Mn-based oxygen carriers prepared by impregnation for Chemical Looping Combustion with diverse fuels

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Abstract

Chemical Looping Combustion (CLC) is considered one of the low cost alternatives for CO\textsubscript{2} capture for fossil fuels combustion and to reach negative emissions through biomass CLC. The cornerstone of the CLC process is the oxygen carrier performance that represents the main additional cost with respect to the conventional combustion. Manganese-based oxygen carriers are subjected to a growing interest because they are low cost, not toxic and environmentally friendly. In this work five impregnated oxygen carriers, with manganese oxide Mn\textsubscript{3}O\textsubscript{4} or Mg\textsubscript{6}MnO\textsubscript{8} as their active phase and three commercial supports based on zirconia and synthetic calcium aluminate, were prepared. Their behaviour for CLC was examined by TGA, batch fluidized bed reactor, TPR, SEM-EDX and XRD. After a preliminary screening two carriers (Mn-ZrM and Mn-ZrSG) were subjected to multiple redox cycles by TGA and batch fluidized bed reactor. Both showed high solids conversion by TGA under the tested conditions, appropriated resistance to fracture, rate indexes relatively high, although Mn-ZrM showed agglomeration and deactivation during batch fluidized bed tests.

Reactivity in batch fluidized bed reactor of the Mn-ZrSG oxygen carrier with methane increases with temperature although suffered from significant deactivation. This was different to the results found during multiple redox cycles by TGA. There was not a clear reason for this decrease in the
reactivity that likely could be due to the uncomplete oxidation in the batch fluidized bed reactor, although further investigations are needed. On the other hand, it presented high and constant reactivity with CO and H₂ in all the range of temperatures tested, being suitable for iG-CLC processes of coal or biomass and syngas combustion. Agglomeration problems were not found and the attrition losses were small. Calculated lifetime was around 11000 hours, much higher than any other Mn-based material developed or tested for CLC.

**Keywords:** CO₂ capture; Chemical Looping Combustion; Oxygen Carrier; Manganese oxide

1. Introduction

Chemical Looping Combustion (CLC) is considered as a low cost process for CO₂ capture during combustion of gaseous and solid fuels due to its inherent CO₂ separation and its low energy penalty. Chemical Looping Combustion process is based on the transfer of the oxygen from the air to the fuel through a solid oxygen carrier, avoiding the direct contact between the fuel and the air. In CLC, as shown in Fig. 1, fuel and air are never mixed and this kind of combustion can be classified as an unmixed combustion [1]. In the first step (reaction 1), the fuel is oxidized to CO₂ and H₂O by a metal oxide (MeₓOᵧ) that is reduced to MeₓOᵧ₋₁. The water generated in the process can be easily separated by condensation, thus obtaining a highly concentrated stream of CO₂, ready for transportation and storage [2]. The reduced metal oxide is further oxidized with air (reaction 2) in a second step and the regenerated material is ready for a new cycle.

\[
(2n+m) \text{Me}_x\text{O}_y + C_n\text{H}_{2m} \rightarrow (2n+m) \text{Me}_x\text{O}_{y-1} + m\text{H}_2\text{O} + n\text{CO}_2 \quad (1)
\]
\[
2 \text{Me}_x\text{O}_{y-1} + \text{O}_2 \rightarrow 2 \text{Me}_x\text{O}_y \quad (2)
\]

CLC concept has been proposed to be carried out in different types or reactors and configurations. The more common configuration of the CLC system corresponds to two interconnected fluidized
bed reactors, which are designated as: air reactor (AR) and fuel reactor (FR), with particles of metal oxide (oxygen carrier) circulating between the two reactors [2].

![Simplified diagram of the CLC system](image)

**Fig. 1.** Simplified diagram of the CLC system.

An overview of the CLC process applied to the combustion of gaseous and solid fuels can be found in a number of review publications [3-7]. In the last few years, significant efforts have been made in the area of oxygen carrier development. [3,4,6]. Most of the oxygen carriers proposed in the literature as suitable for gas combustion (CH₄, natural gas, CO + H₂) are synthetic materials using as active metal oxides (CuO, Fe₂O₃, NiO and Mn oxides) and some mixed oxides with perovskite structure, containing Mn. Moreover, there are some studies showing the suitability of the use of minerals as iron and manganese ores, ilmenite or waste materials coming from steel industry and alumina production, although their performance is usually lower [6].

The cornerstone of CLC process is the performance of the oxygen carrier. This material must display a number of different characteristics in order to be suitable for CLC. Oxygen carriers needs to have sufficient oxygen transport capacity ($R_{oc}$), with high reactivity both for reduction and oxidation reactions and this must be maintained for a large number of redox cycles. Oxygen carriers need to have favourable thermodynamics regarding fuel conversion to CO₂ and H₂O. Moreover, negligible carbon deposition during reduction is needed to avoid C shortcutting to the air reactor, which reduces overall CO₂ capture efficiency. Good fluidization properties and no
agglomeration in the reactors are fundamental for the smooth operation of interconnected fluidized bed reactors. Environmental and health issues must be considered to ensure the process meets future high standards of environmental performance.

Resistance to the attrition is a key point in order to reduce losses of elutriated fines and to reduce the oxygen carrier makeup costs when fluidized bed reactors are used. Thus, the main additional cost for CLC corresponds to the cost of the oxygen carrier replacement. The cost of the makeup stream of solids to replace the loss of fines will depend on: the lifetime of the oxygen carrier particles, the inventory in the CLC system and the cost of the oxygen carrier, which is mainly affected by the oxygen carrier reactivity, metal used and its content in the carrier. The oxygen carrier inventory depends mainly on the oxygen carrier reactivity; the redox pair used; the flow characteristics of gas and solids in the reactors, the solid circulation between reactors and the bed pressure drop [6]. Rate indexes can be used to extrapolate inventories, although give inaccurate values which only can be considered for initial estimations. However rate indexes can be used to compare the reactivity of different oxygen carriers. The rate index corresponds to the % of metal oxide mass that reacts per minute [8] and can be determined at a gas concentration corresponding to the mean value in the reactor.

Rate index (%/min)= 60·100 (dω/dt) (3)

where ω is calculated by ω = m/m_{ox}, m_{ox} being the mass of the sample in the most oxidized state and m the mass of the sample at any reacting time t.

Ni oxygen carriers suffer from environmental and health risks and Cu is a relatively expensive material and temperatures lower than 900 °C are recommended due to the low Tammann temperature of metallic copper, as well as in order to extend the lifetime of most promising materials impregnated on alumina [6]. On the contrary iron is considered a low cost and non-toxic material, although reactivity of Fe-based oxygen carriers is limited. Mn-based oxygen carriers are
also cheap materials, non-toxic and environmentally friendly. Moreover, they have high melting point and oxygen transport capacity when compared with Fe-based oxygen carriers. Mn-based oxygen carriers accomplish most basic requirements to be used as oxygen carrier [6] and when compared with iron carriers, it developed a similar or a superior activity, besides being inexpensive and non-toxic [9]. Another benefit is that some fuels have sulphur compounds in their composition, that react with the oxygen carrier forming sulphides in the CLC process, for example, materials that have in their base Ni and Co. However, manganese-based carriers, are thermodynamically stable in a CLC environment with fuels that have sulphur in their composition [6,10].

Several oxidation states may be involved in redox reactions of manganese. MnO₂, compound decompose in air at 500°C, while the Mn₂O₃ is thermodynamically stable in air at temperatures below 900°C [6,11,12]. Thus, the redox transformation among Mn₃O₄ to MnO is considered the most feasible for CLC [6,13]. The Mn₃O₄/MnO system is capable to convert completely CH₄ to CO₂ and H₂O [10]. Despite the promising properties of manganese to be used as oxygen carrier, few materials based on Mn were tested in CLC, especially when compared to nickel, copper and iron. Synthetic manganese-based oxygen carriers reported in the literature are combinations of active phase and support, where in most of them solid state reactions between Mn and the support occurred, decreasing the reactivity of these materials. Manganese-based carriers deposited/impregnated with alumina or silica, usually presented aluminates and silicates formation, respectively. The formation of these mixed oxides, reduces the reactivity of the material, presenting low fuel conversion [8,11,14-16]. Titanium oxide (TiO₂) was also used as a support for Mn₃O₄, showing low reactivity due to the formation of titanates during the sintering process of the material [15]. Some comprehensive screening studies [8,15-21] selected Mn-based oxygen carriers supported on ZrO₂ or ZrO₂ stabilized with MgO as suitable for CLC process.

One oxygen carrier selected in [8] containing 40 % Mn₃O₄ and 60 % MgO stabilized ZrO₂ was tested in a 300 W continuous CLC unit [9] at temperatures from 850 to 950 °C burning syngas and
natural gas during 70 h with encouraging results. This carrier was best suited for syngas combustion reaching combustion efficiencies higher than 99%. For natural gas combustion efficiencies of 87.8% at 850 °C and 99% at 959 °C were found. Although crushing strength was 1.1 N, the attrition losses found were low, 0.038%/h, with a calculated lifetime of 2630 h. The reaction kinetics was determined for this carrier and reaction orders of 1 for reduction with CH₄ and 0.65 with O₂ were found. The activation energies were 119 kJ/mol and 19 kJ/mol for the reduction and the oxidation respectively [12].

Manganese ores have a great potential to be considered as oxygen carriers in CLC processes using solid fuels or syngas and its reactivity has been measured in different investigations. Depending of the origin of the ore the reactivity and the attrition behaviour show significant differences [22-28]. Mn-Fe mixed oxides depending on the composition and calcination conditions are capable of generate gaseous oxygen and also to use lattice oxygen for CH₄, CO and H₂ combustion [28-32]. Some investigations on Mn-Fe mixed oxides have been also carried out showing the capability of this kind of materials to burn fuels in CLC units [32-34]. Rydén et al. [33,34] used Fe-Mn and Mn-Fe-Ti [34] materials in a 300Wₘₜ unit to burn methane. Good conversion of methane was found although the lifetime of the particles was extremely short. (Mn₇₉Fe)₂O₃ was investigated in a 500 Wₘₜ unit burning gaseous fuels and also different solid fuels [32] showing good reactivity for CO, H₂ and solid fuels with high lifetimes (6120 h).

The aim of this work is the development of impregnated Mn based oxygen carriers using different supports and their characterization for Chemical Looping Combustion. Mn-based oxygen carriers prepared by incipient wetness impregnation on CaAl₂O₄ and ZrO₂ of different sources were developed and examined for CLC by TGA and batch fluidized bed reactor testing. In order to select promising materials for CLC in continuous units, oxygen carriers were evaluated regarding the mechanical properties, reactivity with fuel gases and O₂ in order to select the more promising materials. Selected materials were further tested in a batch fluidized bed reactor during multiple redox cycles with fuel gases to know the fluidization behaviour, agglomeration behaviour and the
attrition resistance. Fresh and after being used carriers were also tested by several characterization techniques to analyse the evolution of their chemical and structural properties during CLC redox cycles.

2. Experimental

2.1. Oxygen Carrier preparation

Five oxygen carriers were prepared by incipient wetness impregnation using as supports calcium aluminate or zirconia, with manganese oxides as the active phase. The active metal oxide was obtained from manganese nitrate solutions (5 or 7 M of Mn (NH₃)₂.4 H₂O, Merck) impregnated on the supports. Three zirconia-based supports were used: MELChemicals, YanYu Industrial and Saint Gobain (Zirpro). The CaAl₂O₄ support was prepared by a mechanical mixing followed by pelletizing under pressure and calcination [35]. Table 1 presents the supports used, the adopted nomenclature, the methodology of preparation used, the metal oxide fraction and the oxygen transport capacity of the developed materials [36]. The oxygen transport capacity of the materials (ROC) was determined using the temperature programed reduction results obtained in the CI Electronics thermobalance used in the reactivity experiments.

**Table 1.** Characteristics of the developed oxygen carriers.

<table>
<thead>
<tr>
<th>Support</th>
<th>Nomenclature</th>
<th>Prep. method</th>
<th>Steps</th>
<th>Mn₃O₄ (%)</th>
<th>ROC</th>
</tr>
</thead>
<tbody>
<tr>
<td>CaAl₂O₄ (Syntetic)</td>
<td>Mn-CaAl</td>
<td>IHI</td>
<td>2</td>
<td>15.9</td>
<td>1.1</td>
</tr>
<tr>
<td>ZrO₂ (Yanyu)</td>
<td>Mn-ZrY</td>
<td>IHI</td>
<td>2</td>
<td>18.8</td>
<td>1.3</td>
</tr>
<tr>
<td>ZrO₂ (Saint Gobain)</td>
<td>Mn-ZrSG</td>
<td>IHI</td>
<td>2</td>
<td>16.3</td>
<td>1.1</td>
</tr>
<tr>
<td>ZrO₂ (MELChemicals)</td>
<td>Mn-ZrMH</td>
<td>IHI</td>
<td>1</td>
<td>21.4</td>
<td>2.7</td>
</tr>
<tr>
<td>ZrO₂ (MELChemicals)</td>
<td>Mn-ZrM</td>
<td>II</td>
<td>1</td>
<td>15.3</td>
<td>2.0</td>
</tr>
</tbody>
</table>

Example of nomenclature of the oxygen-carrier: Mn-ZrSG, the Mn matches the active phase (manganese oxide); the Zr is the inert support (ZrO₂-based), SG the support source (in this case Saint Gobain). In preparation method “H” indicates that the oxygen carrier was prepared by incipient hot impregnation.
2.2. Oxygen Carrier Characterization

The oxygen carriers were characterized by X-ray diffraction (XRD), in a Bruker D8 Advance Series 2 equipment, using CuKα as the radiation source, with a voltage of 40 KV, a current of 40 mA, a graphite diffracted beam monochromator, with a step of 0.05 degrees. The sample holder used had a zero background, manufactured in Silicon Crystal and is indicated for small amounts of sample. The sample holder was kept in horizontal position with rotation, to minimize the effect of preferential orientation of the crystals. Data were collected in the range of 2θ (two theta) from 3 to 80 degrees. The identification of crystalline structures and phases presented were obtained by reference sheets of the JCPDS (Joint Committee on Powder Diffraction Standards) contained in the database of the ICDD (International Center for Diffraction Date, 2003 version).

The resistance to fracture for both supports and oxygen carriers were measured using a Shimpo FGN-5X equipment. The crushing strength is the average of 20 measurements performed for each material. Temperature programmed reduction (TPR) experiments were obtained with an AUTOCHEM II equipment from Micromeritics. This analysis used about 180 mg of dried sample. The sample was heated to a temperature up to 1000 °C, with a heating rate of 7 °C/min, using 10% H2-90%Ar as reducer gas, with a flow rate of 0.02 L/min. The reduction temperatures and H2 consumption data were obtained by a thermal conductivity detector (TCD). The morphology of the oxygen carriers was evaluated before and after the reactor experiments, by Scanning Electron Microscopy (SEM) and Energy Dispersive Spectroscopy (EDS) using an equipment of Shimadzu, model SSX 550 SuperScan.

2.3. Oxygen Carrier reactivity testing

2.3.1. Thermogravimetric Analyser

The reactivity of oxygen carriers was measured using a thermogravimetric analyser, CI Eletronics. The setup has a system to receive the data, and a gas supplying system with mass flow controllers to prepare the reacting gas mixture. The methodology used the variations of mass in the oxygen
carrier, when it was subjected to temperatures and different reacting gases during redox cycles. The conversion of the samples was obtained using Eqs. (4) and (5) 1 and 2 [12].

\[
\text{Reduction: } X_{\text{red}} = \frac{m_{\text{ox}} - m}{m_{\text{ox}} - m_{\text{red}}} \\
\text{Oxidation: } X_{\text{ox}} = 1 - \frac{m_{\text{ox}} - m}{m_{\text{ox}} - m_{\text{red}}}
\]

where \(X_{\text{red}}\) and \(X_{\text{ox}}\) are the reduction and oxidation conversions, respectively; \(m_{\text{ox}}\) is the mass of the sample in the most oxidized state under the tested conditions; \(m_{\text{red}}\) is the mass of the sample in the most reduced state; and \(m\) the mass of the sample. For data evaluation it was considered that the mass of 100% would correspond to the sample in its higher oxidation state analysed.

A sample of 50 mg was heated up by synthetic air until the reacting temperature (850 or 950 °C). When the desired temperature was reached and the stability of the system was checked, the experiments began and the oxygen carrier was subjected to several reduction/oxidation cycles. To prevent the mixture of the reduction and oxidation gases, nitrogen was introduced between each reduction and oxidation stage.

During the experiments, the gas flow was maintained at 25 NL/h, seeking to reduce the oscillations occurred when there was a change of gases and to minimize the resistance to gas film diffusion. A gas composition of 15% of fuel (H₂ or CH₄) and 20% of water vapour was used during reduction, and synthetic air during oxidation. The steam was used to prevent the deposition of carbon when carbon-containing gases were used. This experiment was able to determine, together reactivity several other parameters, which ensure that the carrier is feasible for CLC, such as the effective amount of metal oxide, the rate index and the oxygen transport capacity of the oxygen carrier.

2.3.2. Fluidized bed facility

A batch fluidized bed reactor, was used since it resemble the conditions of a CLC process, and it allows to evaluate the distribution of the products formed in the reduction and oxidation steps; fluidization/agglomeration problems; to determine the attrition rates; and to analyse the carbon
deposition. Fig. 2 shows a diagram of the batch fluidized bed setup used. The setup consists of a gas supply system, a fluidized bed batch reactor, two solid recovery filters from the reactor and a system that analyses the outlet gases of the reactor [37]. The gas supply and steam system were composed by different mass flow controllers that are connected to the bottom of the reactor. Always between a period of reduction or oxidation, pure N₂ was flowed to avoid contact between the reducer and oxidant gases. This system allowed the use of different fuel gas mixtures, as well as different oxidizing gases.

![Diagram of the discontinuous fluidized bed reactor.](image)

**Fig. 2:** Schematic diagram of the discontinuous fluidized bed reactor.

For the experiments, around 300 grams of oxygen carrier were used, with a particle size from 100 to 300 µm. The superficial gas velocity was fixed at least three times the minimum fluidization velocity. The gas velocity was chosen to have a good fluidization without particle entrainment. Table 2 presents the composition of the reactive gases and temperatures used during the experiments performed in the reactor. Reduction reaction times were selected to match the stoichiometric fuel gas consumption. For the oxidation the time needed for complete oxidation was used.
Table 2. Experimental conditions used during batch fluidized bed reactor experiments.

<table>
<thead>
<tr>
<th>Temperature (°C)</th>
<th>Reduction Stage</th>
<th>Oxidation Stage</th>
</tr>
</thead>
<tbody>
<tr>
<td>850, 900 and 950 °C</td>
<td>25%CH₄+10%H₂O + 65%N₂</td>
<td>5%O₂ + 95%N₂</td>
</tr>
<tr>
<td></td>
<td>50%H₂ + 20%H₂O + 30%N₂</td>
<td>50%CO + 20%CO₂ + 30%N₂</td>
</tr>
</tbody>
</table>

The composition of the gases at the outlet of the fluidized bed batch reactor was determined using different continuous gas analysers. The concentrations of CO, CO₂ and CH₄ was determined through a non-dispersive infrared analyser (NDIR) from Maihak S710/UNOR. The H₂ concentration was determined by a thermal conductivity detector (TCD) from MAIHAK S710 model THERMOR, and the concentration of O₂ with a paramagnetic analyser from Siemens OXYMAT 5E model. The concentrations of the gases are recorded continuously after the reactor outlet. From these data the solid conversion as a function of time was calculated. Time corrections were made, due to the delay to reach the gas analysers. The Eqs. (6) and (7) were determined based on the mass balance and were used to calculate the solid conversion in the reduction and oxidation steps, respectively [37].

Reduction stage:

\[
X_{\text{red}} = \frac{1}{n_0} \int_{t_0}^{t_{\text{out}}} \left( F_{\text{out}} (2y_{\text{CO}_2,\text{out}} + y_{\text{H}_2\text{O},\text{out}}) + F_{\text{in}} (2y_{\text{CO}_2,\text{in}} + y_{\text{H}_2\text{O},\text{in}}) \right) \, dt
\]

(6)

Oxidation stage:

\[
X_{\text{ox}} = \frac{2}{n_0} \int_{t_0}^{t_{\text{out}}} F_{\text{out}} \left( \frac{F_{\text{in}}}{F_{\text{out}}} y_{\text{O}_2,\text{in}} - y_{\text{O}_2,\text{out}} - \frac{1}{2} y_{\text{CO}_2,\text{out}} - y_{\text{CO}_2,\text{in}} \right) \, dt
\]

(7)
where $X_{\text{red}}$ and $X_{\text{ox}}$ is the conversion of the oxygen carrier during the reduction and oxidation stages, respectively; $F_{\text{in}}$ and $F_{\text{out}}$ are the molar gas flow at the inlet and outlet of the reactor, respectively; $y_{i,\text{in}}$ is the molar fraction of the $i$ gas at the inlet of the reactor; $y_{i,\text{out}}$ is the molar fraction of the $i$ gas in the reactor outlet; $n_0$ is the number of moles of oxygen that the oxygen carrier can provide in its most oxidized state; and $t$ is the time of the reaction.

To determine the attrition rate, the solids recovered in the filters were weighted and subsequently its particle size distribution was determined, whereas the particles with a size under than 40 $\mu$m were considered as generated by attrition. The attrition rate was calculated through Eq. (8).

$$n_{\text{attrition}} = \frac{m_{\text{elut}}}{m_{\text{OC}}} * 100 * 3600 \quad \text{(8)}$$

where $n_{\text{attrition}}$ is the attrition rate ($%/h$); $m_{\text{elut}}$ is the mass of oxygen carrier with a particle size smaller than 40 $\mu$m; $m_{\text{OC}}$ is the total mass of the oxygen carrier in the bed, and $t$ the time of operation in which the sample was collected from filters. From the attrition rate, it is also possible to calculate the average lifetime of the oxygen carriers.

3. Results

3.1. Oxygen Carrier characterization

X-ray diffraction (XRD) of the supporting porous particles was done to know the main constituents of these materials. The synthetic CaAl$_2$O$_4$ support showed CaAl$_2$O$_4$ and CaAl$_4$O$_7$ phases in the monoclinic structure, evidencing the efficiency of the methodology used in the synthesis of the material [35]. The supports from YanYu Industrial and Saint Gobain were composed by zirconia (ZrO$_2$) and were used without any prior treatment to obtain the manganese-based oxygen carrier.

In both diffractograms the baddeleyite phase was identified, which has the chemical formula ZrO$_2$ and a monoclinic crystalline structure. The support from MELChemicals (ZrM) was composed by hydrotalcite and zirconia. This support ZrM presented peaks for MgO and ZrO$_2$, both with cubic
structure, although it was not possible to identify any structure for the aluminum present in hydrotalcite. Mn-CaAl, Mn-ZrY and Mn-ZrSG oxygen carriers only have Mn$_3$O$_4$ as their active phase, with the name of hausmanite with tetragonal structure. Equally, the XRD results not showed crystalline structures that could be assigned to interactions of active phase with the support or a change in their structures, indicating that neither CaAl$_2$O$_4$ nor ZrO$_2$ react with manganese oxide in the synthesis conditions used [35]. In the oxygen carriers prepared with MELChemicals (ZrM) (Mn-ZrMH and Mn-ZrM) the same phases were identified: MgAl$_2$O$_4$, MgO, ZrO$_2$, Mg$_6$MnO$_8$ and Mn$_3$O$_4$. The formation of the MgAl$_2$O$_4$ spinel occurred due to reaction between aluminium and magnesium oxides, both from the hydrotalcite when subjected to high temperatures. The active phases identified in these oxygen carriers were Mn$_3$O$_4$ and Mg$_6$MnO$_8$, with the manganese oxidation states of $+\frac{8}{3}$ and +4, respectively. The Mg$_6$MnO$_8$ material has an oxidation state similar of MnO$_2$, which has high oxygen transfer capacity. It is believed that both the excess magnesium oxide in support as well as the calcination conditions promoted the formation of a mixed oxide, composed by magnesium and manganese (Mg$_6$MnO$_8$) [38,39]. Table 3 shows the references letters, chemical formulas, crystalline structures, crushing strength and reduction and oxidation rate indexes found for the prepared oxygen carriers. Although it was not established a relationship between the mechanical resistance, through the crushing strength, and the half-life of an oxygen carrier, it was accepted that particles with the crushing strength lower than 1 N were overly fragile to be used in CLC systems with continuous circulation of solids [8]. Prepared oxygen carriers, presented a crushing strength higher than 1 N, except the material Mn-CaAl. With the impregnation of the support ZrY, a significant increase in the crushing strength from 0.7 to 7.3 was observed.
Table 3. Properties of supports and developed oxygen carriers.

<table>
<thead>
<tr>
<th>Sample</th>
<th>References of Chemical formulas</th>
<th>Structures</th>
<th>Crushing Strength (N)</th>
<th>Rate Index (%/min)</th>
</tr>
</thead>
<tbody>
<tr>
<td>CaAl</td>
<td>01-070-0134 CaAl₂O₄</td>
<td>Monoclinic</td>
<td>0.7</td>
<td>Reduction 4.1</td>
</tr>
<tr>
<td></td>
<td>01-074-1467 CaAl₄O₇</td>
<td>Monoclinic</td>
<td></td>
<td>Oxidation 9.1</td>
</tr>
<tr>
<td>Mn-CaAl</td>
<td>01-080-0382 Mn₃O₄</td>
<td>Tetragonal</td>
<td>0.8</td>
<td>Reduction 4.1</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>Oxidation 9.1</td>
</tr>
<tr>
<td>ZrY</td>
<td>01-074-1200 ZrO₂</td>
<td>Monoclinic</td>
<td>0.8</td>
<td>Reduction 4.0</td>
</tr>
<tr>
<td>Mn-ZrY</td>
<td>01-080-0382 Mn₃O₂</td>
<td>Tetragonal</td>
<td>7.3</td>
<td>Reduction 3.0</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>Oxidation 2.1</td>
</tr>
<tr>
<td>ZrSG</td>
<td>01-074-1200 ZrO₂</td>
<td>Monoclinic</td>
<td>1.1</td>
<td>Reduction 13.1</td>
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<tr>
<td>Mn-ZrSG</td>
<td>01-080-0382 Mn₃O₄</td>
<td>Tetragonal</td>
<td>2.5</td>
<td>Reduction 13.1</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>Oxidation 8.3</td>
</tr>
<tr>
<td>ZrM</td>
<td>01-087-0653 MgO</td>
<td>Cubic</td>
<td>1.0</td>
<td>Reduction 4.0</td>
</tr>
<tr>
<td></td>
<td>00-027-0997 ZrO₂</td>
<td>Cubic</td>
<td></td>
<td>Reduction 4.0</td>
</tr>
<tr>
<td>Mn-ZrM</td>
<td>00-011-0031 Mg₆MnO₈</td>
<td>Cubic</td>
<td>3.4</td>
<td>Reduction 6.8</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>Oxidation 4.1</td>
</tr>
<tr>
<td></td>
<td>01-080-0382 Mn₃O₄</td>
<td>Tetragonal</td>
<td></td>
<td>Reduction 6.8</td>
</tr>
<tr>
<td></td>
<td>01-088-1007 ZrO₂</td>
<td>Tetragonal</td>
<td></td>
<td>Reduction 6.8</td>
</tr>
<tr>
<td></td>
<td>00-030-0794 MgO</td>
<td>Cubic</td>
<td></td>
<td>Reduction 6.8</td>
</tr>
<tr>
<td></td>
<td>01-077-0437 MgAl₂O₄</td>
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<td>Reduction 6.8</td>
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<tr>
<td>Mn-ZrMH</td>
<td>00-011-0031 Mg₆MnO₈</td>
<td>Cubic</td>
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<td>Reduction 5.9</td>
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<tr>
<td></td>
<td>01-080-0382 Mn₃O₄</td>
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<td></td>
<td>Reduction 5.9</td>
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<tr>
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<td></td>
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<tr>
<td></td>
<td>00-030-0794 MgO</td>
<td>Cubic</td>
<td></td>
<td>Reduction 5.9</td>
</tr>
<tr>
<td></td>
<td>01-077-0437 MgAl₂O₄</td>
<td>Cubic</td>
<td></td>
<td>Reduction 5.9</td>
</tr>
</tbody>
</table>

The results for temperature programmed reduction proved the formation of manganese oxides identified by XRD. The reduction of Mn₃O₄ to MnO was demonstrated by the reduction peaks in TPR ranging from 350-550 °C, and the position of the peak depends on the size and morphology of the particles of Mn₃O₄; see Fig. 3a [40,41]. It is noticed that the Mn-ZrSG material has a greater
trend to be reduced than other materials, since its reduction occurs in a lower temperature range. 

The reduction profile of the material Mn-ZrSG reveals there was not interaction between Mn$_3$O$_4$ and support. The material Mn-ZrY has a reduction peak that is intermediate when compared with the previous one, although it presents a region of difficult reduction at temperatures above 500 °C. The carrier Mn-ZrY presented one less intense reduction peak, possibly due to the formation of Mn$_3$O$_4$ particles with a reduction more difficult. This fact could be proven by the region with temperature of 515°C where the reduction occurs slowly. This difficulty to reduce the Mn-ZrY could affect the reduction rate of the material during the reactivity tests, reducing its effective use for oxygen transport. The material Mn-CaAl presented a maximum reduction around 500 °C and it was also possible to check a shoulder at 528 °C, that can be attributed to the low intensity interactions among the particles of the Mn$_3$O$_4$ and the support.

**Fig. 3.** Profiles of the temperature programmed reduction of the oxygen carriers prepared: a) Mn-CaAl, Mn-ZrY and Mn-ZrSG; b) Mn-ZrM and Mn-ZrMH.

XRD on oxygen carriers impregnated on the MELChemicals support showed Mg$_6$MnO$_8$ and Mn$_3$O$_4$ as active phases. The Mg$_6$MnO$_8$ reduction likely occurs in a similar way to pure MnO$_2$, being noticed three peaks of reduction with hydrogen; see Figure 3b. The first one is at low temperature with a peak in 395 °C. The rest were overlapping peaks located at temperatures above 500 °C. It
was found that the peak intensity at low temperature was substantially lower than those that occur at high temperatures. It was observed a reduction shoulder around 400 °C, which can be attributed to the reduction of Mn$^{4+}$ to Mn$^{3+}$ of the superficial Mg$_6$MnO$_8$ [40]. The high intensity in the reduction temperature region from 450-800 °C may be the result of several species of manganese that have close reduction temperature, that contributed to this peak. In this region, the reduction of Mn$^{3+}$ to Mn$^{2+}$ can occur in two steps (Eqs. 9 and 10), as well as the reduction of free Mn$_3$O$_4$ to the MnO (Mn$_3$O$_4$ free was identified by XRD beyond the reductions of Mn$^{4+}$ remaining.

\[
3\text{Mn}_2\text{O}_3 + \text{H}_2 \rightarrow 2\text{Mn}_3\text{O}_4 + \text{H}_2\text{O} \quad (9)
\]

\[
\text{Mn}_3\text{O}_4 + \text{H}_2 \rightarrow 3\text{MnO} + \text{H}_2\text{O} \quad (10)
\]

Considering the extreme reduction conditions, in which the samples were subjected during TPR analysis, the mixed oxide was reduced, according with XRD results as follows:

\[
\text{Mg}_6\text{MnO}_8 + \text{H}_2 \rightarrow 6\text{MgO} + \text{MnO} + \text{H}_2\text{O} \quad (11)
\]

and that the manganese changed from Mn$^{4+}$ to Mn$^{2+}$ after the end of the experiment [40,42].

### 3.2. Reactivity of oxygen carriers

Initially, the five prepared oxygen carriers were subjected to the reduction/oxidation cycles in order to assess their reactivity. The evaluated temperatures were 850 and 950 °C. Variations in reactivity were not observed during multiple redox isothermal tests. Fig. 4 shows the results found for fifth cycle with the five manganese-based oxygen carriers, comparing the reactivity with different fuels (H$_2$ or CH$_4$) and at different temperatures. The following reactions were considered for the mass variations found in the thermobalance for the active phases Mn$_3$O$_4$ and Mg$_6$MnO$_8$:

\[
\text{CH}_4 + 4\text{Mn}_3\text{O}_4 \rightarrow \text{CO}_2 + 2\text{H}_2\text{O} + 12\text{MnO} \quad (12)
\]

\[
\alpha\text{CH}_4 + 4\text{Mg}_6\text{MnO}_8 \rightarrow \alpha\text{CO}_2 + 2\alpha\text{H}_2\text{O} + 4\text{Mg}_6\text{MnO}_{8-\alpha} \quad (13)
\]

\[
\text{H}_2 + \text{Mn}_3\text{O}_4 \rightarrow \text{H}_2\text{O} + 3\text{MnO} \quad (14)
\]
\[ \alpha \text{H}_2 + \text{Mg}_6\text{MnO}_8 \rightarrow \alpha \text{H}_2\text{O} + \text{Mg}_6\text{MnO}_8-\alpha \]  \hspace{1cm} (15)  

\[ 6\text{MnO} + \text{O}_2 \rightarrow 2\text{Mn}_3\text{O}_4 \] \hspace{1cm} (16)  

\[ \text{Mg}_6\text{MnO}_{8-\alpha} + (\alpha/2)\text{O}_2 \rightarrow \text{Mg}_6\text{MnO}_8 \] \hspace{1cm} (17)  

According to the results presented in Fig. 4 the better results for the Mn-CaAl material corresponded to H\textsubscript{2} 950 °C, in which about 80% of solid was converted. The results showed that the rate of oxidation was not influenced by the examined temperatures, and in all tests the carrier was regenerated at virtually the same time. For the oxygen carrier Mn-ZrY the reactivity with hydrogen gas was higher when compared with methane, at both temperatures. Similarly, the conversion achieved in 850 °C was lower than one from 950 °C, needing in both cases around 60 s to achieve the maximum reduction. The reactivity of Mn-ZrSG with H\textsubscript{2} and CH\textsubscript{4} was nearly identical, reaching conversions around 90 %, in both reduction and oxidation, in a time lower than 30 seconds in all cases analyzed. For the carriers Mn-ZrMH and Mn-ZrM, Mn\textsubscript{3}O\textsubscript{4} and Mg\textsubscript{6}MnO\textsubscript{8} were identified as active phases. For Mn\textsubscript{3}O\textsubscript{4} the reactions happen according to the Eq. 12, 14 and 16. While, Eqs. 13, 15 and 17 represent the reactions that occurred with the active phase Mg\textsubscript{6}MnO\textsubscript{8} (where \( \alpha \) is < 1).  

Mn-ZrMH oxygen carrier at 950 °C reached a conversion around 90% in less than 180s with fuel gas (H\textsubscript{2} and CH\textsubscript{4}) and the oxidation happens quickly up to 100 s, and then very slowly, indicating that the active phase presents some difficulty to fully regenerate to its initial state. The experiments performed at 850 °C, with hydrogen the carrier conversion was 100% while with methane was 82%. Oxidations at 850 °C have similar behaviour to the oxidations at 950 °C, where neither the temperature nor the gas reducer gives to step changes in the oxidation of this material. For Mn-ZrM the conversion with hydrogen was the same at both analysed temperatures (100%), while for methane it ranged around 85% and 74% at 850 °C and 950 °C, respectively. All oxidations had similar behaviour, where almost all the oxidation occurs very quickly, up to 50 s, and later a small fraction of the oxidation occurs slowly indicating a change in the controlling step.
Fig. 4. Conversion of oxygen carriers by thermogravimetry for the first cycle with fresh oxygen carriers: a) $T=950 \, ^\circ\text{C}$, using $\text{H}_2$ as the reducer; b) $T=950 \, ^\circ\text{C}$, using $\text{CH}_4$ as the reducer; c) $T=850 \, ^\circ\text{C}$, using $\text{H}_2$ as the reducer; and d) $T=850 \, ^\circ\text{C}$, using $\text{CH}_4$ as the reducer. All the oxidations were with synthetic air.

3.3. Screening of oxygen carriers

A preliminary screening was made considering that the oxygen carrier must have sufficient crushing strength (>1 N), high reactivity and to reach the maximum possible conversion. From the analysed results, it was found that from the five carriers studied only two of them could be considered satisfactory (Mn-ZrSG and Mn-ZrM). These oxygen carriers were further tested by TGA and batch fluidized bed reactor for 100 redox cycles to know if reactivity was maintained.
3.3.1. Reactivity by 100 cycles of reduction/oxidation

The selected Mn-ZrM and Mn-ZrSG oxygen carriers were subjected to 100 cycles of reduction/oxidation in a thermobalance, in order to analyse the reactivity of these materials throughout multiple redox cycles; see Fig. 5. The reducing gas was composed of 15% CH₄ + 20% H₂O + 65% N₂. Synthetic air was used as oxidizing agent and pure nitrogen to purge the system between oxidation and reduction steps. Reduction and oxidation were carried out at the temperature of 950 °C. Each reduction/oxidation step was of 60 s, followed by 30 s of purge.

![Graphs showing solid conversion over time for Mn-ZrM and Mn-ZrSG](image)

Fig. 5. Conversion achieved with Mn-ZrM and Mn-ZrSG materials during 100 redox cycles. Reduction: 15% CH₄ + 20% H₂O; Oxidation: air; Reacting temperature: 950 °C.

During 100 cycles, the Mn-ZrM material initially presented a high conversion (~ 80 %), with a gradual decrease of the solid conversion, decreasing until 30 % in the 50th cycle, indicating one important deactivation. On the other hand, the oxygen carrier Mn-ZrSG presented very fast
reaction rate in both the reduction and oxidation steps, reaching about 95% of its total oxygen transport capacity in less than 10 s. These results showed that the Mn-ZrSG particles are quite reactive with methane and maintained the reactivity over the 100 cycles of reduction/oxidation.

3.3.2. Reactivity in fluidized bed batch reactor

Mn-ZrM material also was evaluated in fluidized bed batch reactor, although it presented serious agglomeration problems during fluidization in the initial experiments conducted. By this reason, the material was considered unfeasible to proceed with the scheduled tests and it was rejected for use in CLC.

Mn-ZrSG oxygen carrier was initially tested with methane at temperatures from 850 to 950 °C. Fig. 6 shows the product distribution during redox cycles (no 1 and 20). At all the temperatures tested the CH₄ conversion was not complete. The CH₄ slip decreased with temperature, although it was important even at 950 °C. The gas product distribution showed significant concentrations of CO and H₂ at the reactor outlet. CO and CO₂ appeared also during oxidation indicating the C deposition during the reduction step. The maximum conversion of CH₄ at 950 °C was found during the first cycle. In this Figure, a deactivation of the oxygen carrier with the number of redox cycles is also observed. Moreover, it can be observed that O₂ concentration does not return quickly to its initial value during oxidation. This slow increase indicates that oxidation was slow and not complete. Unconverted CH₄ was always present at high concentrations together a decrease of the CO₂ concentration at the outlet of the reactor and one increase in the concentrations of CO and H₂. This was different to the results obtained during 100 redox cycles TGA, although it could be due to the insufficient oxygen carrier inventory in the batch reactor. It was not clear the possible reasons for this behaviour because, although the fluidization was not smoot, there was not observed any agglomeration in the bed. Difficulties in oxidation could contribute to the deactivation of the oxygen carrier. Samples taken from the bed maintained its oxygen transport capacity although its reactivity had an important decrease with a change in the reduction rate index from 13.1 to 5.
Fig. 6. Gas product distribution during redox cycles in discontinuous fluidized bed reactor working with the Mn-ZrSG using CH₄ as fuel: a) 1st cycle at 950 °C; b) cycle 20 at 950 °C.

Moreover the oxygen carrier Mn-ZrSG was examined for the combustion of H₂ and CO. The reduction reaction of manganese oxide when using CO or H₂ as fuel is slightly exothermic [10], and an increase of 15 °C inside the reactor was observed during the reductions stage. Fig. 7 shows the gas product distribution obtained at temperatures from 800 to 950 °C using CO (50 % CO + 20 %CO₂ + 30 % N₂) as fuel. At the three analysed temperatures, similar behaviour was found (Figs. 7a; 7b; 7c), where initially it was seen basically CO₂ concentrations, indicating that in this interval all fuel was burned. After 25 s, the unburned CO concentrations gradually increased reaching a maximum of 10%, due to the decrease in the amount of oxygen available in the oxygen carrier. During the oxidation, in the first step all the oxygen feed in was consumed in the oxidation of the oxygen carrier. The O₂ concentration increased rapidly until it reached the input gas concentration. Also in the oxidation step, compounds such as CO and CO₂ were not detected, what indicates that there was not carbon formation during the reduction stage. Multiple redox cycles were carried out to follow the possible deactivation of the oxygen carrier. Fig. 7d shows the gas product distribution in the redox cycle no. 20. It was possible to observe that the particles of Mn-ZrSG oxygen carrier kept its reactivity and solid conversion after 20 redox cycles in the same
conditions. Considering all cycles analysed for Mn-ZrSG oxygen carrier with carbon monoxide as fuel, the conversion of the solid \( \chi_{\text{red}} \) in the reduction step was 0.83.

**Fig. 7.** Gas product distribution in the batch fluidized bed reactor working with Mn-ZrSG using CO as fuel: a) \( T = 850 \, ^\circ\text{C} \); b) \( T = 900 \, ^\circ\text{C} \); c) \( 950 \, ^\circ\text{C} \) (first cycle); d) after 20 cycles at \( T = 950 \, ^\circ\text{C} \).

The hydrogen gas was another fuel examined in the fluidized bed batch reactor with Mn-ZrSG oxygen carrier. Fig. 8 shows the gas product distribution at the outlet of the reactor during one reduction/oxidation cycle of Mn-ZrSG, using hydrogen in different conditions. It turns out that the redox cycles behaviour were virtually equal at the three temperatures, where nearly the whole reduction period the conversion of \( H_2 \) to generate \( H_2O \) occurred, since no hydrogen gas concentration was observed until the last steps of the reduction. The increasing concentrations of \( H_2 \) at the end of the reduction (after 60 s) were attributed to low concentrations of the active phase, \( \text{Mn}_3\text{O}_4 \), when the oxygen carrier conversion was higher than 50 \%. This does not occur in a continuous circulating fluidized bed CLC system where the supply of the oxidized oxygen carrier
(Mn₃O₄) is constant. During most part of the oxidation, the oxygen concentration was zero, because it was completely consumed in the regeneration of the oxygen carrier, while its velocity was limited by the oxygen supply to the reactor.

Moreover the reactivity of Mn-ZrSG was evaluated during 20 reduction/oxidation cycles at 950 °C with hydrogen. Fig. 8d shows the gas product distribution during the redox cycle no. 20. In all redox cycles using H₂ as fuel, it was found that this carrier kept its reactivity with hydrogen gas. The solid conversion during the reduction step was 0.68.

The formation of carbon on the oxygen carrier particles can be a problem, since it can deactivate the material, to produce some agglomeration problems and mainly it decrease the efficiency of CO₂ capture because in the CLC unit C is transferred together the oxygen carrier to the air reactor in which it is oxidized to CO₂ and emitted to the atmosphere. In all the experimental work with
methane gas at 950 °C with Mn-ZrSG, it was found in all the 20 cycles the generation of CO and CO₂ during the oxidation step. Similar behaviour was found at lower temperatures. In all the oxidation cycles, the concentration of carbon monoxide and carbon dioxide were lower than 2%. However, in any experiment carried out with CO as fuel it was not noticed the formation of carbon during the reduction period.

3.3.3. Fluidization behaviour

As it was previously indicated, Mn-ZrM material presented serious problems of agglomeration and fluidization during batch fluidized bed tests. However during all the testing time with Mn-ZrSG oxygen carrier, about 74 hours, the pressure drop in the bed was kept almost constant indicating that agglomeration problems were not present, although fluidization was not smooth with CH₄. Moreover, Mn-ZrSG particles extracted at the end of the tests did not showed any evidence of agglomeration. To determine the attrition rate of the oxygen carrier during redox cycles, 60 reduction/oxidation cycles were performed under fluidization conditions, during 39 hours of fluidization. In all redox cycles methane gas was used as reducer, oxygen as oxidant gas (5% O₂) and pure nitrogen to purge the system. Fig. 9 shows the attrition rate as a function of time, for the oxygen carrier Mn-ZrSG.

Fig. 9. Attrition rate of the Mn-ZrSG oxygen carrier during multi-cycle reaction in the fluidized bed during 39 hours of operation (60 redox cycles).
A greater attrition rate occurred during the first two cycles, which can be attributed to the rounding effect of particle irregularities. After the 20th cycle, a small increase in the attrition loss occurred, that can be associated to the fragmentation and/or attrition on the surface of the particles. It is important to notice that the Mn-ZrSG oxygen carrier presented a very low attrition rate. Considering all the testing done the attrition rate was 0.009 weight %/hour, which corresponds to an average operation lifetime for the Mn-ZrSG particles of more than 11000 hours, what is a high value when compared to other oxygen carriers reported in the literature [8,43].

3.4. Characterization after redox cycles

3.4.1. X-ray Diffraction (XRD)

In Fig. 10a the diffractograms obtained for the Mn-ZrM oxygen carrier fresh and after the experiments in batch fluidized bed reactor are presented. Not agglomerated particles and other ones taken from agglomerates were analysed. It was possible to notice significant differences in the obtained diffractograms, highlighting the decrease of peaks that were related with MgAl2O4 (2 theta in 16.3; 30.7; 36.3; 44.1; 58.4 and 64.3°) by comparing the samples fresh and after the experiments at the reactor. Comparing the particles tested in the reactor in a similar way, it was possible to observe a greater amount of peaks characteristic related with the Mg6MnO8 structure on the agglomerated particle with values of 2 theta in 18.4; 35.5 and 57.3°. After redox cycles, significant structural changes in Mn-ZrM material occurred, as proved by the differences in the peaks and their corresponding intensities, which indicate a greater concentration of manganese oxides in the agglomerated particles. This is in accordance with the TPR and SEM results, which will be presented later.
Fig. 10. X-ray diffraction for oxygen carriers fresh and after the experiments in the reactor: a) Mn-ZrM and b) Mn-ZrSG.

For Mn-ZrSG oxygen carrier, fresh and after all the experiments in the reactor, the same crystalline structures were identified, which are: ZrO$_2$ with monoclinic structure, and the Mn$_3$O$_4$ with tetragonal structure; see Fig. 10b. The intensities of the diffraction peaks were virtually the same. This highlights the excellent ability of Mn-ZrSG oxygen carrier in maintaining its crystalline structure after several reduction/oxidation cycles and hours of operation.

### 3.4.2. Temperature programmed of reduction

Fig. 11 shows temperature programmed reduction profiles of Mn-ZrM and Mn-ZrSG, both fresh and after the experiments in the reactor. For Mn-ZrM material (Fig. 10a) different behaviour can be observed between the fresh and agglomerated or not agglomerated particles, proving that samples were substantially modified. Both, agglomerated or non-agglomerated samples, presented a significant decrease in the amount of H$_2$ consumption, compared with that consumed by the fresh particles. This decrease in hydrogen consumption changed from 15 to 60%, between agglomerated and non-agglomerated particles.
Fig. 11. Temperature programmed reduction for oxygen carriers fresh and after reactor experiments: a) Mn-ZrM and b) Mn-ZrSG.

In Fig. 11b it was observed that the oxygen carrier Mn-ZrSG after 74 hours of operation in a fluidized bed batch reactor, kept its reducing behaviour, with only a slight variation of the temperature. Furthermore, the hydrogen consumption was the same for particles fresh and after the tests in the reactor, and maintained the same oxygen transport capacity.

3.4.3. Scanning Electron Microscopy (SEM)

Fig. 12 shows micrographs of the Mn-ZrM material fresh and after the tests in the reactor. It can be observed the presence of agglomerated particles in Fig. 12b, which were responsible for the de-fluidization of the system and for the decrease in reactivity of this material. The agglomerated material was composed mainly by smaller material particles stuck. The morphology of the fresh particles and of those that did not agglomerate in the reactor was similar. The chemical microanalysis by energy dispersive spectroscopy (EDX) showed a greater superficial concentration of manganese in the particle that was agglomerated (Fig. 12b), corroborating the assumption that the manganese oxides migrated to the surface and between particles.
Fig. 12. Scanning Electron Microscopy of the Mn-ZrM oxygen carrier, fresh and after the experiments in the reactor: a) fresh; b) agglomerated particles and c) non-agglomerated particles.

Fig. 13 shows the micrographs obtained for Mn-ZrSG oxygen carrier, fresh and after the experiments performed in the batch fluidized bed reactor. According to Fig. 13, the morphology of particles, fresh and after the experiments in the fluidized bed reactor, were similar, with rounded aspect and almost identical sizes. One higher superficial concentration of manganese in particles tested in the reactor was observed by EDX. This migration happened from the inner to the external surface of the particles during operation at high temperature in the reactor.
Fig. 13. Scanning Electron Microscopy and EDX profiles of the Mn-ZrSG oxygen carrier submitted in reactor: a) fresh; b) after the experiments.

4. Discussion

The results of XRD and TPR showed that Mn-CaAl, Mn-ZrY and Mn-ZrSG oxygen carriers formed as active phase Mn$_3$O$_4$, which was in accordance with the reports from the literature and thermodynamic data [6, 9-12]. On the other hand, MELChemicals support which has in its composition MgO, (originated from hydrotalcite), favoured the solid state reaction with manganese oxide, during calcination at 950 °C, forming a mixture of manganese oxides: Mg$_6$MnO$_8$ and Mn$_3$O$_4$ as active phases. The advantage of the mixed oxide Mg$_6$MnO$_8$ was its ability to provide gaseous oxygen, similar to MnO$_2$. It was verified that high Mg/Mn molar ratios, when calcined in air at temperatures above 800 °C, improved the formation of the Mg$_6$MnO$_8$ [38,39] compound.
The material Mn-CaAl showed a crushing strength lower than 1N, although it presented high reaction rates (%/min) and the maximum conversion reached at 850 °C was of 73%. Because of this, the oxygen carrier Mn-CaAl was not further considered.

The carrier Mn-ZrY showed the greatest value of crushing strength (7.3 N). On the other hand, its reactivity with reducing gases was low, especially at 850 °C, which could be attributed to the no complete reduction of Mn$_3$O$_4$ under the examined conditions, as it was verified by the TPR experiments. Moreover the oxidation was slow which makes difficult the oxygen carrier regeneration. As a consequence, the values of the rate indexes for reduction and oxidation were the lower ones found between all the prepared oxygen carriers. By the above reasons this oxygen carrier was not selected for further investigations.

These reactivities were higher than that of the Mn$_3$O$_4$/ZrO$_2$ oxygen carrier obtained by mechanical mixing [15]. The slow re-oxidation found for Mn-ZrMH and Mn-ZrM oxygen carriers, was associated to the active phase Mg$_6$MnO$_8$. This was attributed to the partial destruction of the spinel structure of the mixed oxide when it was reduced under the conditions examined, resulting in MgO and MnO formation (Eq. 11). However the regeneration to Mg$_6$MnO$_8$ in air at high temperatures occurs slowly and it was not complete. It was reported that some mixed oxides, such as calcium manganese perovskites, with Mn (IV) in the structure should be reduced up to +3, to preserve the perovskite structure of the active phase [44]. Similarly, it is believed that the Mg$_6$MnO$_8$ should be reduced only to a certain level to keep its structure and facilitate the regeneration, although this fact that needs to be confirmed.

Oxygen carriers Mn-ZrMH and Mn-ZrM presented crushing strength values higher than 3 N and the largest oxygen transport capacity among the prepared materials. The maximum conversion reached by these two oxygen carriers was quite similar and the rate index for reduction was slightly higher for Mn-ZrM. During multiple redox cycles by TGA with Mn-ZrM, it was verified that the oxidation step controlled the reactivity of the oxygen carrier because when the Mg$_6$MnO$_8$ phase is reduced it cannot return completely to its initial conversion during the oxidation; and/or
the residence time in the oxidation was not high enough to recover the active phase. These reasons drive to a decrease in the amount of oxygen available to each subsequent cycle, leading to a decreased conversion of the oxygen carrier until the stabilization. This fact can be assigned to the early reaction rate to be controlled by chemical reaction, and later, when most of the particles had reacted, the reaction can be controlled by the solid diffusion [12]. Moreover during batch fluidized bed reactor tests, defluidization of the bed was observed which was originated by the agglomeration of the oxygen carrier particles. By this reason Mn-ZrM was not considered as suitable for CLC.

Mn-ZrSG oxygen carrier had a sufficient crushing strength (2.5 N) well above 1 N. Mn-ZrSG oxygen carrier presented high reduction and oxidation reactivities reaching the maximum conversion in less than 30 seconds. During all tests the maximum conversion reached during oxidation and reduction steps were around 90 % and the rate indexes found for reduction and oxidation were high, 13.1 and 8.5 respectively at 950 °C. During multiple redox cycles (100 cycles) by TGA, the reactivity, the maximum redox conversion and the physic-chemical properties were maintained, as it was verified in the XRD, TPR and SEM. The above results make this oxygen carrier as a promising material for CLC processes and it was qualified for more advanced studies.

However a different behaviour was found for Mn-ZrSG oxygen carrier when it was tested with methane in the batch fluidized bed reactor. Results indicated the low reactivity of the carrier with a non-complete CH\textsubscript{4} conversion and the formation of carbon in all the conditions tested. Reactivity after reaction was reduced showing a rate index of 5.2 but the oxygen transport capacity was maintained. This indicates that higher bed inventories were needed to reach complete CH\textsubscript{4} conversion. Product distribution was not adequate for Chemical Looping Combustion of CH\textsubscript{4} because always CO and H\textsubscript{2} appeared after short periods of reduction. Moreover, both CO and H\textsubscript{2} concentrations increased and the initial time for appearance was shortened with the number of redox cycles, indicating some deactivation of the oxygen carrier. These differences with TGA results cannot be attributed to in-bed agglomeration problems and
could be likely due to the uncomplete oxidation, although a further investigation is needed to clarify the reasons for this behaviour. Experiments performed with Mn-ZrSG and carbon monoxide as fuel, presented good performance. CO burning efficiency increased with temperature and one solid conversion as high as 0.83 in all cycles was found. During the main part of the reduction step all CO was converted to CO₂. After this period, CO₂ concentrations decreased while the CO increased. This change in the reduction was associated with the conversion degree of the oxygen carrier and the lattice oxygen availability. However in a continuous reactor it is not expected unburned CO at the outlet, since the supply of oxidized Mn-ZrSG will be constant, and the appearance of the CO is produced by reduction of active phase (Mn₃O₄) in the last reduction stages. Moreover, in these experiments it was not possible to verify carbon deposition in any condition evaluated, indicating that this carrier did not favour solid carbon formation reactions from the CO nor by inverse Boudouard reaction (2CO → C + CO₂).

Using hydrogen gas as fuel, during the first 60 s, the combustion of H₂ was 100% and in all analysed cycles the conversion of the solid was constant, 0.68, not being influenced by the temperature nor the cycle number. At the end of reduction, the concentration of hydrogen increases quickly. The above results indicated the suitability of Mn-ZrSG oxygen for combustion of mixtures (CO+H₂) and for solid fuel CLC. Thus, this oxygen carrier is considered suitable for gas combustion, especially for syngas combustion. For CH₄, although the material was highly reactive, some further investigations are needed to explain results found in the batch fluidized bed reactor that mainly could be attributed to a loss of reactivity originated by uncomplete oxidation. Likely, difficulties in the bed fluidization that did not reach bed agglomeration could partially explain this behaviour.

To put in context the reactivity of the selected oxygen carrier, it was made a comparison with different Mn based oxygen carriers tested in the literature showing rate indexes. Table 4 shows the rate indexes with CH₄, CO and H₂ together the crushing strength (CS) and the lifetime of Mn
based oxygen carriers both synthetic (Mn₃O₄, Mg-ZrO₂, Mn-Fe) and different Mn ores when they are available. Considering only the reactivity through the rate indexes, rhodonite (OXMN010A) was the oxygen carrier with higher reactivity with CH₄, CO and H₂. It is necessary to mention that Mn ores lost their reactivity with the redox cycles with very important decreases in the rate index for (MnSA, MnGBHNE, MnGBMPB, MnBr). By this, only values in brackets in Table 4 were considered. It can be observed that after some redox cycles, rate indexes with CH₄ for Mn ores are quite low. The oxygen carrier developed Mn-ZrSG showed relatively high rate indexes for CH₄, CO and H₂.

Table 4. CLC characteristics of tested natural and synthetic Mn materials.

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<th>Oxygen carrier</th>
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<th>Rate index (%/min)</th>
<th>CS (N)</th>
<th>Lifetime (h)</th>
<th>Ref.</th>
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<tr>
<td>Mn77Fe</td>
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<td>17.9</td>
<td>38.8</td>
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<td>14.2 (1.8)</td>
<td>4.6 (2.0)</td>
</tr>
<tr>
<td>MnGBHNE</td>
<td>Ore, Gabon</td>
<td>9.2 (1.3)</td>
<td>6.4 (1.4)</td>
<td>19.2 (9.0)</td>
<td>1.8 (1.4)</td>
</tr>
<tr>
<td>MnGBMPB</td>
<td>Ore, Gabon</td>
<td>9.3 (2.2)</td>
<td>9.0 (2.5)</td>
<td>26.4 (14.8)</td>
<td>2.4 (0.5)</td>
</tr>
<tr>
<td>MnBR</td>
<td>Ore, Brasil</td>
<td>12.7 (1.7)</td>
<td>8.2 (3.0)</td>
<td>20.5 (12.5)</td>
<td>1.0 (0.4)</td>
</tr>
<tr>
<td>Mn Alheim</td>
<td>Ore, Norway</td>
<td>7.2</td>
<td>---</td>
<td>19.8</td>
<td>---</td>
</tr>
<tr>
<td>Mn Sinai-A</td>
<td>Ore</td>
<td>3.6</td>
<td>---</td>
<td>5.4</td>
<td>---</td>
</tr>
<tr>
<td>Mn Guizhou</td>
<td>Ore</td>
<td>11.3</td>
<td>---</td>
<td>11.3</td>
<td>---</td>
</tr>
</tbody>
</table>

Rate index by TGA at 950 C. Lifetime calculated from batch fluidized bed reactor. Preparation- IMP: Impregnated; SD: spray drying; FG freeze granulation. In brackets properties after several redox cycles.
The main advantage of Mn-ZrSG comes from the presence of ZrO₂ as support for impregnation that drives to a lifetime as high as 11000 h. This lifetime is very high in comparison with the synthetic Mn₃O₄-ZrO₂ [9]. With respect to Mn-Fe that has a relatively high lifetime (6150 h) the developed material has much higher reactivity with CH₄, CO and H₂. Moreover, Mn ores examined in the literature showed significant attrition rates driving to calculated lifetimes ranging from 150 to 1600 h. Only rhodonite (OXMN010A) oxygen carrier had high reactivity and one high lifetime of 2900 h.

Mn-ZrSG showed during TGA tests good reactivities with CH₄, CO and H₂ and maintained the reactivity with the redox cycles. Moreover during batch fluidized bed reactor redox cycles excellent performance was found burning CO and H₂ showing very low attrition rates with calculated lifetimes around 11000 hours. Thus this oxygen carrier can be considered suitable for CLC, specially for syngas combustion and solid fuel chemical looping through in-situ gasification chemical looping (iG-CLC).

5. Conclusions

In this work five impregnated oxygen carriers, with manganese oxide as their active phase and three commercial supports based on zirconia and synthetic calcium aluminate, were prepared and their behaviour for CLC was examined by TGA, batch fluidized bed reactor, TPR, SEM-EDX and XRD.

Mn-CaAl, Mn-ZrY and Mn-ZrSG oxygen carriers, presented the Mn₃O₄ active phase, while for Mn-ZrM and Mn-ZrMH active phases were mixed oxide Mg₆MnO₈ and Mn₃O₄. After preliminary screening two carriers (Mn-ZrM and Mn-ZrSG) were subjected to multiple redox cycles by TGA and batch fluidized bed reactor. Both showed high solids conversion by TG under the tested conditions; appropriated resistance to fracture; rate indexes relatively high although Mn-ZrM showed agglomeration and deactivation during batch fluidized bed tests.
Reactivity of the Mn-ZrSG oxygen carrier with methane increases with temperature operating with CH₄ although suffered from significant deactivation. This was different to the results found during multiple redox cycles by TGA. There was not a clear reason for this decrease in the reactivity that likely could be due to the uncomplete oxidation in the batch fluidized bed reactor, although further investigations are needed. On the other hand, it presented high and constant reactivity with CO and H₂ in all the range of temperatures tested, being suitable for iG-CLC processes of coal or biomass and syngas combustion. Agglomeration problems were not found and the attrition losses were small. Calculated lifetime was around 11000 hours, much higher than any other Mn based material developed or tested.

**Acknowledgements**

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