# Effect of H<sub>2</sub>S on the behaviour of an impregnated NiO-based oxygen carrier for Chemical-Looping Combustion (CLC)

Cristina Dueso, María T. Izquierdo, Francisco García-Labiano\*, Luis F. de Diego, Alberto Abad, Pilar Gayán, Juan Adánez.

Instituto de Carboquímica (ICB-CSIC), Department of Energy and Environment Miguel Luesma Castán 4, 50018 Zaragoza, Spain Phone number: +34 976 733 977 Fax number: +34 976 733 318

\*Corresponding Author. Tel.: +34-976-733977; fax: +34-976-733318; *E-mail address*: glabiano@icb.csic.es

#### Abstract

Gaseous fuels for chemical-looping combustion (CLC) process can contain sulphurcompounds which can affect the oxygen-carrier behaviour, especially if NiO is used as active phase. In this work several samples of a NiO-based oxygen-carrier prepared by impregnation, NiO18- $\alpha$ Al, extracted from a CLC unit after continuous operation with CH<sub>4</sub> containing 500 vppm of H<sub>2</sub>S were characterized. Part of the fed sulphur to the system was release as SO<sub>2</sub> in the air-reactor during the CLC experiments while the rest remained in the solid particles. Mainly Ni<sub>3</sub>S<sub>2</sub> was found in the oxygen-carrier extracted from the fuel-reactor, although small amounts of NiSO<sub>4</sub> were also detected. On the contrary, NiSO<sub>4</sub> was the main sulphur compound in the oxygen-carrier from the airreactor and only a low concentration of Ni<sub>3</sub>S<sub>2</sub> was found. Despite the accumulated sulphur and the oxygen transport capacity loss during the operation, the oxygen-carrier was capable of recovering the initial reactivity for the CH<sub>4</sub> combustion after a time without H<sub>2</sub>S feeding to the CLC system. In addition, a study about the possible regeneration of the oxygen-carrier in the air-reactor working at different temperatures and oxygen concentrations was performed. Independently of the operating conditions,

part of the sulphur remained in the solid and total regeneration was not possible. The analysis of the NiO18- $\alpha$ Al oxygen-carrier after the CLC operation using TPR and XPS techniques revealed that sulphur reacted preferably with free NiO instead of NiAl<sub>2</sub>O<sub>4</sub>. Although Ni<sub>3</sub>S<sub>2</sub> was the majority sulphide in the fuel-reactor, minor amounts of other sulphides such as NiS were detected. It also was found that sulphur was preferably concentrated in the outer surface of the particles. Taking into account all the results found, a previous desulphuration process of the fuel would be necessary when using NiO-based oxygen-carriers in the CLC system.

#### Keywords

Chemical-looping combustion (CLC); oxygen-carrier; nickel oxide; sulphur.

#### **1. Introduction**

During last decades, negative effects of global warming have been a constant source of concern to the scientific community. Huge research efforts have been done trying to turn the trend of the increasing greenhouse gas emissions to the atmosphere, primarily  $CO_2$ . Since about a third of the global  $CO_2$  emissions come from the burning of fossil fuels for power generation [1] and a significant change in the dependence on this kind of fuels is not expected in the near future,  $CO_2$  capture and storage in geological formations has emerged as a possible solution to reduce anthropogenic emissions. A large number of processes which accomplish the capture of  $CO_2$  from combustion sources are currently available or under development, i.e. pre-combustion, post-combustion or oxyfuel technologies. Nevertheless, most of these techniques have several drawbacks as the high cost and energy consumption and, therefore, a decrease in the global efficiency of the energy generation process.

Within this framework, Chemical-Looping Combustion (CLC) has been identified as a promising technology for clean fossil fuel combustion since the  $CO_2$  capture is inherent to the process. Hence, any separation step or additional energy is not needed to produce a pure  $CO_2$  stream ready for compression and sequestration. A metal oxide acts as an oxygen-carrier circulating between two fluidized-bed reactors, the fuel- and the airreactor, and transports the oxygen from the combustion air to the fuel, thus avoiding the dilution of flue gases with the N<sub>2</sub> of the air. In the fuel-reactor, the fuel gas ( $C_nH_{2m}$ ) is oxidized to  $CO_2$  and  $H_2O$  by the metal oxide which is reduced at the same time to a metal or the reduced form of the oxide. Almost pure  $CO_2$  can be obtained after  $H_2O$  condensation. The metal or reduced oxide is regenerated with air in the air-reactor for a new cycle. Only N<sub>2</sub> and unreacted  $O_2$  are released from this reactor. Although the reduction can be exothermic or endothermic depending on the type of metallic oxide and the fuel, and the oxidation reaction is always exothermic, the total amount of heat generated in the CLC system is the same as from conventional combustion with the mixture of the air and the fuel.

Several transition state metals, such as Ni, Cu, Mn, Fe and Co, have been proposed as the most suitable materials for CLC [2-4]. The metal oxide is usually supported on an inert material, which increases its mechanical strength and provides a higher surface area for the reaction. Oxygen-carriers prepared over Al<sub>2</sub>O<sub>3</sub>, SiO<sub>2</sub>, TiO<sub>2</sub> and yttrium stabilized zirconia (YSZ) as binders can be found in the literature. NiO-based oxygen-

carriers have attracted most attention for gas combustion due to their high reactivity with methane, main component of natural gas and refinery gas [5-8], and their thermal stability. More than 20 different oxygen-carriers based on nickel oxide have been investigated in different CLC continuous units from 300 W<sub>th</sub> to 140 kW<sub>th</sub> for 2500 hours [2], showing very high combustion efficiencies. Nevertheless, thermodynamic restrictions avoid full conversion of the fuel into CO<sub>2</sub> and H<sub>2</sub>O and small CO and H<sub>2</sub> concentrations can be found in the outlet gas stream from the fuel-reactor of the CLC system. The use of Al<sub>2</sub>O<sub>3</sub> as support for NiO-based oxygen-carriers has been widely studied in the literature due to its good fluidization properties and thermal stability. However, a drawback of this material is NiAl<sub>2</sub>O<sub>4</sub> formation [9]. At high calcination temperatures (> 1073 K), part of NiO can react with the alumina to form nickel aluminate [10], which has lower reactivity than free NiO.

Gaseous fuels for the CLC process, such as natural gas, refinery gas or synthesis gas from coal gasification, may contain different amounts of impurities, like sulphur in the form of H<sub>2</sub>S or COS, which can interact with the metallic oxide, thus affecting the performance of the CLC system. Although the H<sub>2</sub>S concentration in the natural gas is usually quite low ( $\approx$ 20 vppm) [11], the content of this compound may reach values of 800 vppm in a refinery gas [12] and even 8000 vppm in a raw synthesis gas [13]. Sulphur in the fuel could react with the metallic oxide in the oxygen-carrier to form sulphur-containing compounds, such as sulphides and sulphates, that could decrease the oxygen-carrier reactivity and, as a consequence, the energy efficiency of the process. Moreover, the low melting point of some sulphides could produce the agglomeration of the particles and affect the solids circulation pattern between the interconnected fluidized bed reactors [10].

Although the poisoning effect of sulphur for Ni catalysts is especially well-known and a lot of works about this issue can be found in the literature, not many articles about the effect of sulphur on oxygen-carriers for the CLC process have been published up to now. First, several thermodynamic studies appeared. Mattisson et al. [10] determined that  $H_2S$  could be partially oxidized to  $SO_2$  in the fuel-reactor using a NiO-based oxygen-carrier and  $CH_4$  as fuel, improving the conversion degree at high temperatures and low pressures. Both gaseous compounds,  $SO_2$  and  $H_2S$ , could react with nickel in the particles to form Ni<sub>3</sub>S<sub>2</sub>, the most stable sulphide. Nickel sulphate was not present under the reducing fuel-reactor conditions. Later, Jerndal et al. [14] investigated the

thermodynamics of oxygen-carriers based on Ni, Cu, Fe, Mn, Co, W and sulphates of Ba and Sr regarding to carbon deposition and formation of sulphides and sulphates using CH<sub>4</sub>, H<sub>2</sub> and CO as fuels and found the same results for the nickel behaviour in the presence of sulphur. Wang et al. [15] also studied carbon deposition and the sulphur effect on the performance of oxygen-carriers based on five metal oxides (NiO, CuO, Fe<sub>2</sub>O<sub>3</sub>, Mn<sub>3</sub>O<sub>4</sub> and CoO) through thermodynamic simulations using syngas as fuel and they again observed the tendency of nickel to react with the sulphur to form Ni<sub>3</sub>S<sub>2</sub> in the particles.

More recently other authors presented studies performed in thermobalance (TGA) and CLC pilot plants [16-20]. They found that, although the presence of H<sub>2</sub>S affected the reaction rates and reactivities of the oxygen-carriers, the formation of sulphides was completely reversible and SO<sub>2</sub> was released in the outlet stream from the air-reactor. Forero et al. [21] analysed the behaviour of a Cu-based oxygen-carrier prepared by impregnation in a 500 W<sub>th</sub> CLC continuous unit. Sulphur was mainly released as SO<sub>2</sub> in the fuel-reactor ( $\approx$ 95%). Formation of Cu<sub>2</sub>S took place in this reactor but it was oxidized in the air-reactor forming SO<sub>2</sub>. No deactivation of the oxygen-carrier was observed even working with high H<sub>2</sub>S concentrations (1300 vppm). Mayer [22] investigated the performance of a mixture of two nickel-based oxygen-carriers (N-VITO and N-VITO Mg) prepared by spray drying in a 120 kW<sub>th</sub> CLC unit using natural gas containing 48 vppm of H<sub>2</sub>S. A decrease in the combustion efficiency was observed during the sulphur treatment, but the oxygen-carrier was regenerated when the H<sub>2</sub>S feeding stopped.

In previous works carried out at ICB-CSIC, NiO18- $\alpha$ Al oxygen-carrier had been successfully used in a CLC continuous plant with CH<sub>4</sub> [23] and syngas [24] as fuels, and it showed suitable properties to be considered as oxygen-carrier for CLC process. The behaviour of this solid in the presence of impurities in the fuel, such as hydrocarbons [25] and sulphur [26], was also studied. Several tests with different H<sub>2</sub>S concentrations (100-1000 vppm) in the fuel (CH<sub>4</sub>) and temperatures (1103-1153 K) were carried out in a 500 W<sub>th</sub> continuous plant. During these experiments, a loss of reactivity was always detected, although it had low relevance for H<sub>2</sub>S concentrations below 100 vppm. In addition, a compilation of possible reactions involving sulphur in CLC was made according to the gaseous sulphur species obtained at the outlet stream from the air and the fuel-reactor and the assumed solid compounds formed on the

oxygen-carrier.  $H_2S$  and  $SO_2$  were found at very low concentrations (<10 vppm) in the fuel-reactor outlet stream but sulphur reacted with NiO to form Ni<sub>3</sub>S<sub>2</sub>. This compound was transferred to the air-reactor where it reacted with the air to produce  $SO_2$  or NiSO<sub>4</sub> depending on the temperature. A mass balance to the system also indicated a gradual accumulation of sulphur in the oxygen-carrier particles.

In the work mentioned above [26], a compilation of possible reactions involving sulphur in CLC was made. Therefore, one of the aims of this work was to establish which sulphur compounds were present in the NiO18- $\alpha$ Al oxygen-carrier used in the 500 W<sub>th</sub> continuous CLC unit after CH<sub>4</sub> combustion tests in the presence of H<sub>2</sub>S and hence to show if the proposed reactions took place in the system.

In addition, the most suitable situation for the industrial CLC operation with Ni-based oxygen-carriers would be the total release of the sulphur introduced in the fuel as  $SO_2$  in the air-reactor, in order to keep a high reactivity of the oxygen-carrier during the operation time. For this reason, another objective of this work was to analyse the possible oxygen-carrier regeneration, determining the effect of temperature and oxygen concentration in the air-reactor.

Different characterization techniques such as temperature programmed reduction and oxidation (TPR/TPO) and X-ray photoelectron spectroscopy (XPS) were used in order to determine the sulphur compounds in the oxygen-carrier particles and their degree of interaction with the support.

#### 2. Experimental

#### 2.1. Oxygen-carrier

A nickel-based oxygen-carrier, so-called NiO18- $\alpha$ Al, was used in this work. The desired NiO content in the solid, 18 wt.%, was obtained by applying two successive steps of hot incipient wet impregnation on an  $\alpha$ -Al<sub>2</sub>O<sub>3</sub> support, as described by Gayán et al. [27]. The  $\alpha$ -Al<sub>2</sub>O<sub>3</sub> support particles were prepared by calcination of commercial  $\gamma$ -Al<sub>2</sub>O<sub>3</sub> (Puralox NWa-155, Sasol Germany GmbH) of 100-300 µm at 1423 K for 2 h. Table 1 shows the main properties of the fresh oxygen-carrier. The oxygen transport capacity was defined as the mass fraction of oxygen that can be used in the oxygen

transfer, calculated as  $R_{OC} = (m_{ox} - m_{red})/m_{ox}$ , where  $m_{ox}$  and  $m_{red}$  are the masses of the oxidized and reduced forms of the oxygen-carrier, respectively.

In the experiments carried out in this work, two different types of sulphur-containing samples were used. The first one corresponds to samples obtained after operation in a continuous CLC plant burning  $CH_4$  in presence of  $H_2S$ , while the second one was made up of sulphided particles prepared in a batch fluidized bed. More details about these samples can be found below.

#### **2.2.** Characterization

The reducibility of the NiO18- $\alpha$ Al oxygen-carrier was determined by temperatureprogrammed reduction (TPR) experiments in a flow apparatus Micromeritics AUTOCHEM II. The sample was placed in a quartz tube and flushed with Ar during 1 hour. After this pre-treatment, the oxygen-carrier particles were heated at 7 K/min from room temperature to 1323 K feeding a 20 ml/min flow of a mixture 10 vol.% H<sub>2</sub>-90 vol.% Ar. During these tests, the accuracy in the measurements of the TPR peak was  $\pm$ 1 K. The amount of H<sub>2</sub> consumed was monitored continuously. Temperature programmed oxidation (TPO) runs were performed after cooling down the samples using 20 ml/min of a mixture 5 vol.% O<sub>2</sub>-95 vol.% He and heating up to 1273 K at 15 K/min.

The X-ray photoelectron spectroscopy (XPS) analyses were carried out with an ESCAPlus OMICROM system equipped with a hemispherical electron energy analyser. The spectrometer was operated at 10 kV and 15 mA, using a non-monochromatized MgK<sub> $\alpha$ </sub> X-ray source (hv = 1253.6 eV) and under vacuum (< 5 x 10<sup>-9</sup> Torr). Analyser pass energy of 50 eV was used for survey scans and 20 eV for detailed scans. Binding energies (BE) are referenced to the C 1s peak (284.5 eV) from adventitious carbonaceous contamination. A survey scan (1 sweep/200 ms dwell) was acquired between 1100 and 0 eV. Current region sweeps for Ni 2p, O 1s, C 1s, Al 2p and S 2p were obtained. The CASA XPS data processing software allowed smoothing, Shirley type background subtraction, peak fitting and quantification.

#### 3. Results and discussion

#### 3.1. Sulphur distribution in a 500 W<sub>th</sub> CLC continuous unit.

In a previous work [26], tests carried out in a 500  $W_{th}$  CLC continuous pilot plant allowed studying the behaviour of the NiO18- $\alpha$ Al oxygen-carrier during the combustion of CH<sub>4</sub> (30 vol.%) in the presence of H<sub>2</sub>S. Sulphur content in the oxygencarrier particles extracted from the unit was too low to determine the sulphur compounds (sulphides and/or sulphates) present in the samples by usual characterization techniques. For these reason, new experiments were carried out in the CLC plant.

#### **3.1.1. Preparation of the sulphur-containing materials**

In this work, new combustion tests were performed in the 500  $W_{th}$  CLC continuous unit with 30 vol.% of CH<sub>4</sub> as fuel for three consecutive days. The operating temperatures were 1143 and 1223 K in the fuel and the air-reactor, respectively. The first day experiment corresponded to the reference test without H<sub>2</sub>S. 500 vppm of H<sub>2</sub>S were fed to the system together with the CH<sub>4</sub> for 11 hours during the last two days. The gas velocity at the inlet of the fuel-reactor and air-reactor were 0.1 m/s and 0.45 m/s, respectively. The solids circulation rate was fixed at 12 kg/h.

Figure 1 shows the temperature profiles in both reactors and the riser and the gas concentration at the outlet stream of the fuel and air-reactors during the third day of experimentation. CH<sub>4</sub> was fed to the system when the temperature reached the desired value. At the beginning, some CH<sub>4</sub> was detected due to the formation of nickel sulphides or sulphates in the oxygen-carrier during the previous test. Nevertheless, this concentration decreased with time while CO<sub>2</sub> increased until reaching the steady state conditions. The oxygen-carrier seemed to recover its initial combustion capacity. After H<sub>2</sub>S feeding, CO<sub>2</sub> concentration decreased and some unburnt CH<sub>4</sub> appeared, while the CO and H<sub>2</sub> concentrations increased slowly up to 1.5 and 3 vol.%, respectively. H<sub>2</sub>S and SO<sub>2</sub> were found at very low concentrations (< 10 vppm) in the fuel-reactor outlet stream. SO<sub>2</sub> was mainly released from the air-reactor and its concentration increased with time. A sulphur mass balance in the system and the reactivity loss of the oxygen-carrier indicated that part of the sulphur was being accumulated in the oxygen-carrier.

These results were completely reproducible compared with the values obtained by García-Labiano et al. [26] working with the same oxygen-carrier, NiO18- $\alpha$ Al. After the operation, the solid was carefully extracted from each part of the CLC unit

(air-reactor, fuel-reactor and riser) without mixing and was reserved for later tests in the batch fluidized-bed reactor.

#### **3.1.2.** Sulphur distribution in the oxygen-carrier particles after the operation

The loss of reactivity of the oxygen-carrier observed during the tests in the 500  $W_{th}$  CLC unit was attributed to the formation in the solid particles of some sulphur compounds like Ni<sub>3</sub>S<sub>2</sub> and NiSO<sub>4</sub>. To determine and to quantify the presence of these compounds in the solid after operation, the material extracted from the fuel-reactor, airreactor and riser was divided into fractions of 150 g. These batches were heated in a batch fluidized bed reactor whose description in detail can be found elsewhere [28] from room temperature to 1223 K in air or inert atmosphere (N<sub>2</sub>). After that, the gas atmosphere was changed to nitrogen or air, respectively. SO<sub>2</sub> released in the outlet stream was continuously analysed in a non-dispersive infrared analyser (NDIR) Siemens ULTRAMAT U22.

Figure 2 shows the  $SO_2$  concentration emitted from the samples extracted from the fuelreactor, the air-reactor and the riser of the CLC unit during the different steps with nitrogen and air. Each peak was attributed to the reactions of the sulphur-containing compounds, sulphides and sulphates, thermodynamically possible at each temperature. According to thermodynamic calculations using HSC Chemistry software [29], the possible reactions involving sulphur compounds are the following:

Air atmosphere:

$$Ni_3S_2 + 7/2O_2 \rightarrow 3NiO + 2SO_2 \tag{1}$$

$$Ni_3S_2 + 9/2 O_2 \rightarrow 2 NiSO_4 + NiO$$
<sup>(2)</sup>

Nitrogen atmosphere:

$$NiSO_4 \rightarrow NiO + SO_2 + \frac{1}{2}O_2 \tag{3}$$

 $Ni_3S_2 + 7 NiSO_4 \rightarrow 10 NiO + 9 SO_2 \tag{4}$ 

The results obtained during the heating in air were first considered. Nickel sulphide present in the samples could be oxidized directly to NiO and SO<sub>2</sub> at high temperatures and cause the release of SO<sub>2</sub> according to reaction 1. The only SO<sub>2</sub> peak detected at 1170 K during the heating of the samples from the fuel-reactor, air-reactor and riser could be assigned to this reaction. In addition, nickel sulphide could be also oxidized to NiSO<sub>4</sub> through reaction 2, without any SO<sub>2</sub> release. To demonstrate this hypothesis, air was switched to N<sub>2</sub> and SO<sub>2</sub> was released again from the reactor. This peak would correspond to the decomposition of the nickel sulphate (reaction 3), either already present in the initial sample or formed in the previous oxidation step.

When the samples were heated up in N<sub>2</sub>, two SO<sub>2</sub> peaks were observed at  $\approx$ 600 K and  $\approx$ 1200 K. Thermodynamic calculations using HSC Chemistry software [29] showed that Ni<sub>3</sub>S<sub>2</sub> could not decompose in this inert atmosphere, so the gaseous SO<sub>2</sub> only could come from reactions involving NiSO<sub>4</sub>. The peak at high temperature could be attributed to the decomposition of this compound to give NiO and SO<sub>2</sub> by reaction 3. At low temperatures, both sulphur-containing compounds can react according reaction 4 to release SO<sub>2</sub>. Further changes of gas atmosphere produced similar results to the ones explained above during the tests heating in air.

Assuming that the proposed reactions 1-4 took place in the system, the integration of the different peaks showed in Figure 2 gave us the distribution of the sulphur between  $Ni_3S_2$  and  $NiSO_4$  in the samples, as well as an approximation of the total sulphur released at those conditions. These data are summarized in Table 2.  $Ni_3S_2$  was the major sulphur compound in the sample extracted from the fuel-reactor although some small amounts of  $NiSO_4$  remained in the particles. On the contrary,  $NiSO_4$  was the major sulphur compound in the samples extracted from the air-reactor with small contribution of the  $Ni_3S_2$ . Part of the sulphur retained in the solid as nickel sulphide in the fuel-reactor was released as  $SO_2$  in the air-reactor during the operation in the CLC continuous plant. For this reason, the sulphur amount in the air-reactor sample was lower. This agreed with the results shown in the previous work by García-Labiano et al [26].

The  $SO_2$  concentration curves of oxygen-carrier from the riser should be the same as the ones obtained with the solid from the air-reactor. Nevertheless, as can be seen in Table 2 and Figure 2, small differences were observed during the heating in N<sub>2</sub> as well as in air. The  $SO_2$  peaks from NiSO<sub>4</sub> decomposition increased their intensity. This seemed to indicate that the lower temperature in the riser with respect to the air-reactor favoured

the formation of NiSO<sub>4</sub> from the reaction between gaseous SO<sub>2</sub> and NiO in the oxidized oxygen-carrier through the reverse reaction 3. This fact confirmed the formation of sulphur compounds in the riser proposed by García-Labiano et al. [26]. Lower temperatures in the riser were due to the specific design of this CLC plant. Nevertheless, formation of NiSO<sub>4</sub> could be avoided in an industrial scale CLC plant working at higher temperatures in the riser.

In any case, the above tests showed that a great part of the sulphur present in the samples remained inside the particles despite of the successive regeneration steps with nitrogen or air. It must be considered that the desired situation during the operation of an industrial scale CLC plant would be the emission of all the sulphur contained in the fuel as  $SO_2$  in the air reactor. This will allow having a fully regenerated and highly reactive oxygen carrier at the fuel reactor inlet. To analyse this possibility, new experiments were planned in the batch fluidized bed reactor to study the influence of different parameters (temperature and oxygen concentration) in the oxygen carrier regeneration, as explained below.

#### 3.2. Studies about the regeneration of the oxygen-carrier

During previous combustion tests in the CLC continuous unit feeding mixtures  $CH_4$ - $H_2S$  with the NiO18- $\alpha$ Al oxygen-carrier, it was observed that the SO<sub>2</sub> concentration in the outlet stream of the air-reactor varied depending on the air-reactor and riser temperatures [26]. To analyse the possible regeneration of the sulphur-containing oxygen-carrier, new sulphided material with high S content was prepared and tested at different conditions of temperature and oxygen concentration in the batch fluidized bed reactor.

#### 3.2.1. Preparation of the sulphided sample in a BFB.

About 400 g of NiO18- $\alpha$ Al particles were reduced with 25 vol.% H<sub>2</sub> (N<sub>2</sub> balance) at 1223 K for more than one hour in the batch fluidized-bed reactor. When the H<sub>2</sub> concentration in the outlet stream reached the initial value, indicating that the sample was fully reduced, the system was cooled down to 873 K and 1300 vppm of H<sub>2</sub>S in N<sub>2</sub> (268 l/h, 10 cm/s) were introduced into the reactor for 45 minutes. 5 vol.% of H<sub>2</sub> was fed together with H<sub>2</sub>S to avoid the decomposition of this gas inside the pipes. These

conditions favoured the formation of nickel sulphide, considering the results from previous thermodynamic analyses [26]. Outlet stream gases were analysed in a gas chromatograph (Varian 3400-CX GC) equipped with a PORAPAK-Q packed column and a sulphur-specific flame photometric detector (FPD), calibrated in the range 0-300 vppm for both H<sub>2</sub>S and SO<sub>2</sub>. These gases were not detected during the sulphidation process, so it could be concluded that sulphur was totally accumulated in the oxygencarrier. The final sulphur content in the solid determined by a total sulphur analysis was 0.095 wt.%. This meant that the conversion of Ni to Ni<sub>3</sub>S<sub>2</sub> was 1.85 %. Although this percentage was low, it represented a higher amount of sulphur than in the case of the samples extracted after the operation in the CLC continuous unit. The preparation process of sulphided particles (BFB-S sample) was carried out twice, so  $\approx$  800 g of NiO18- $\alpha$ Al oxygen-carrier were available for subsequent experiments.

#### **3.2.1.** Effect of temperature on the oxygen-carrier regeneration

The sulphided samples previously prepared were regenerated at different temperatures ranging from 1073 to 1263 K, typical temperatures of the air-reactor in a CLC plant. A bed of alumina (300-500  $\mu$ m) was heated to the desired temperature and fluidized with air (190 l/h) in the batch fluidized bed reactor. To introduce the sulphided oxygen-carrier into the reactor, a small lock-hopper located in the upper part of the reactor was used. Then, 20 g of the sulphided particles were quickly introduced into the bed and the released SO<sub>2</sub> was measured continuously using a non-dispersive infrared analyser (Siemens ULTRAMAT U22). When the SO<sub>2</sub> concentration fell down to zero, air was replaced by N<sub>2</sub> trying to regenerate the oxygen-carrier particles completely.

Figure 3 shows the sulphur emitted as  $SO_2$  with respect to the total sulphur present in the sample using air and subsequently nitrogen as fluidization agents at different temperatures. The amount of  $SO_2$  released at temperatures below 1173 K, corresponding to reaction 1, was very low and most of the nickel sulphide was converted into nickel sulphate (reaction 2). This fact can be explained attending to the thermodynamic equilibrium for NiSO<sub>4</sub> decomposition, shown in Figure 4. Independently of oxygen concentration and  $SO_2$  partial pressure, decomposition started being noticeable at 900 K. Nevertheless, the reaction rate might be very low at that temperature since about 200 degrees more were necessary to observe  $SO_2$  formation experimentally. The change from air to  $N_2$  produced SO<sub>2</sub> emissions again (reactions 3 and 4), indicating the presence of NiSO<sub>4</sub> in the sample formed during the previous air feeding step.

It must be pointed out that the oxygen-carrier was never fully regenerated at any of the temperatures tested. A maximum of 12 wt.% of the sulphur could be emitted as  $SO_2$  at the oxidation conditions in air atmospheres. Even if the solid was later treated with nitrogen, only 56 wt.% of the sulphur would be released at 1263 K. These results indicated that the regeneration of the sulphided particles is rather difficult even at high temperatures and part of the sulphur always remains in the solid.

#### 3.2.2. Effect of oxygen concentration on the oxygen-carrier regeneration

The effect of the oxygen concentration on the oxygen-carrier regeneration was also analysed. The experimental procedure was similar to the one described above. In this case, different  $O_2$ - $N_2$  mixtures were used as fluidizing agent instead air at 1223 K.

Figure 5 shows the percentage of sulphur emitted as  $SO_2$  when the sulphided particles were fed into the fluidized bed reactor with different  $O_2$  concentrations. As can be observed, the  $SO_2$  release decreased as the  $O_2$  content in the gas fed to the reactor raised. Therefore, an important effect of the  $O_2$  was inferred since the nickel sulphide oxidation to form nickel sulphate was favoured when the oxygen concentration was high. This fact was confirmed when the  $O_2$ - $N_2$  mixture was replaced by pure  $N_2$  and the  $SO_2$  release as a consequence of the NiSO<sub>4</sub> decomposition took place. In any case, it must be highlighted that only about 40 wt.% of the total sulphur present in the particles reacted with the  $O_2$ , either to release  $SO_2$  or to form sulphates, and the rest remain inside the particle without regeneration.

Nevertheless, regeneration tests carried out during the experiments in the CLC pilot plant with fuel containing  $H_2S$  showed that the oxygen-carrier was capable of recovering its initial reactivity after several hours without  $H_2S$  feeding [26]. Figure 6 shows the TGA reactivity data for the fresh and the regenerated oxygen-carrier after experiments with 300-1000 ppm of  $H_2S$ . As can be seen, the reactivity was very high with almost complete conversion before one minute in both the reduction and the oxidation. At these conditions, the oxygen-carrier could be used with good performance despite the remaining sulphur in the particles. Obviously, although the reactivity could

be recovered, the sulphur remaining in the particles would produce a decrease in the oxygen transport capacity of the material along its life-time. In any case, to corroborate the above results a deeper analysis of the solid particles by means of different characterization techniques was performed.

# **3.3.** Characterization of the NiO18-αAl oxygen-carrier particles after their use in the presence of sulphur

According to the above results, full regeneration of the NiO18- $\alpha$ Al oxygen-carrier would not be possible at the selected conditions. The interaction of sulphur with the complex structure of the oxygen carrier containing different Ni and Al compounds (NiO, NiAl<sub>2</sub>O<sub>4</sub>,  $\alpha$ -Al<sub>2</sub>O<sub>3</sub>, etc.) could affect the regeneration process. For this reason, TPR-TPO and XPS analysis were carried out to analyse the reducibility and to characterize the surface of the oxygen-carrier particles, identifying the main elements, their oxidation state and the type of chemical structure.

Several samples extracted from the CLC continuous unit, before and after the operation with  $H_2S$ , and from the BFB reactor, sulphided and regenerated, were considered for the characterization. The study of fresh oxygen-carrier (so-called Fresh) and particles extracted from the CLC unit after 7 hours of combustion with  $CH_4$  but without  $H_2S$  (sample CLC) were used as reference. Two sulphur-containing samples extracted from the fuel-reactor -CLC-S(FR)- and the air-reactor - CLC-S(AR)- after the operation with  $H_2S$  feeding in the CLC unit were used for further analysis.

For the investigation of the solid from the batch fluidized bed reactor, three samples were selected: the original sulphided oxygen-carrier prepared as explained above (BFB-S), the sulphided oxygen-carrier after a regeneration step with air -BFB-R(A)-, and the material regenerated with air and subsequently with N<sub>2</sub> at 1223 K -BFB-R(AN)-.

#### **3.3.1.** Temperature-programmed reduction (TPR)

Bulk NiO (Sigma-Aldrich Chemie GmbH, Germany, Nickel (II) oxide, black, <10 micron) and  $\alpha$ -Al<sub>2</sub>O<sub>3</sub>, prepared by calcination of commercial  $\gamma$ -Al<sub>2</sub>O<sub>3</sub> (Puralox NWa 155, Sasol Germany GmbH, 100-300  $\mu$ m) were used as reference materials for TPR tests. The TPR curve of bulk NiO powder showed a single peak of hydrogen

consumption with a maximum at 604 K. As expected, an almost negligible peak of hydrogen consumption was observed at 601 K for  $\alpha$ -Al<sub>2</sub>O<sub>3</sub>.

Firstly, a TPR analysis of the fresh NiO18- $\alpha$ Al oxygen-carrier was performed. As can be seen in Figure 7, the fresh particles exhibited three peaks of hydrogen consumption whose positions and bands are reported in Table 3. This could be due to the presence of different NiO species on the alumina and agreed with several studies in literature [30-33]. First peak ( $\approx$ 721 K) could be attributed to isolated NiO units on the surface of the support. The interaction with the alumina was weak but enough to need a higher temperature to be reduced than the one corresponding to bulk NiO. Second peak ( $\approx$ 873 K) was related to the presence of a bi-dimensional NiO monolayer on the top of the support with moderate interaction with  $\alpha$ -Al<sub>2</sub>O<sub>3</sub>. Finally, the third peak ( $\approx$ 1139 K) corresponded to a strong interaction with the support. Either nickel species incorporated in the surface of the support having nickel aluminate character or bulk-NiAl<sub>2</sub>O<sub>4</sub> consisting of nickel cations which migrated deeply under the surface of the alumina support appeared.

After a TPO experiment, a new TPR profile was obtained and some differences with respect to the first analysis were observed, such as a shoulder around 622 K. This could reflect the reduction of Ni<sup>3+</sup> (oxygen excess after TPO) according to the above mentioned literature [31-34]. Moreover, the second hydrogen consumption peak, corresponding to isolated units of NiO, shifted to lower temperatures, indicating that, after the oxidation, the interaction with the support decreased slightly and therefore the reduction of the sample was easier. Peak at 873 K almost disappeared and shifted to around 900 K. This was related to the successive reduction of nickel ions of octahedral and tetrahedral symmetry from non-stoichiometric surface nickel aluminate. The peak corresponding to nickel aluminate appeared at the same temperature but the amount of hydrogen consumption increased significantly. The successive reduction of nickel ions between the two symmetries could explain this fact [34].

A TPR analysis of the samples extracted from the CLC continuous pilot plant was also carried out (TPR-1), as can be seen in Figure 8. The position of peaks and bands are summarized in Table 3. Significant differences were found regarding the TPR of the fresh NiO18- $\alpha$ Al oxygen-carrier. It was observed that most of the nickel was present as nickel aluminate in the samples from both the fuel- and the air-reactor, in contrast with the fresh oxygen-carrier that had a higher content of free NiO. At the same time, the

NiO peak was much lower and its asymmetry indicated a contribution of bulk and isolated NiO species showing a weak interaction with the  $\alpha$ -Al<sub>2</sub>O<sub>3</sub> support. These results agreed with a previous work carried out by Dueso et al. [35] where the ratio NiO/NiAl<sub>2</sub>O<sub>4</sub> in the particles was not a constant but a function of the reduction conversion during the previous redox cycles.

After the in-situ oxidation (TPO) of the samples, a second TPR was performed (TPR-2). At these conditions, almost all the nickel aluminate was converted to NiO during the oxidation for the samples extracted from the air-reactor (CLC and CLC-S(AR)). These results were directly related to those observed in the study of the NiO18- $\alpha$ Al oxygen-carrier reactivity [35]. In this case, sulphur accumulated in the solid particles did not seem to affect the reducibility behaviour of the CLC-S(AR) sample. Nevertheless, the sample extracted from the fuel-reactor (CLC-S(FR)) exhibited a different pattern than the above solid. In this case, an important amount of nickel aluminate was maintained. These results could be due to the higher content of Ni<sub>3</sub>S<sub>2</sub> and NiSO<sub>4</sub> in this sample than in the one obtained from the air-reactor, where the oxygen-carrier was partially regenerated. Ni could have migrated to deeper layers of bulk-NiAl<sub>2</sub>O<sub>3</sub> to form the nickel sulphide and sulphate, being more difficult to convert into NiO. Further XPS analysis shown below could explain this fact.

The TPR profiles from the samples coming from the batch experiments -BFB-S, BFB-R(AN) and BFB-R(A)- are depicted in Figure 9. These samples showed TPR profiles quite different from those exhibited by the samples extracted from the CLC plant (Figure 8). TPR profiles of BFB-R(A) and BFB-R(AN) were very similar to that corresponding to the fresh sample, indicating that the regeneration with air or air/N<sub>2</sub> of the sulphided samples did not alter significantly the Ni species distribution. However, the TPR profile of these two samples obtained after a TPO run (TPR-2) changed. The sample oxidized with air, BFB-R(A), suffered a high re-dispersion of the active phase and the peak corresponding to nickel aluminate almost disappeared. Results were similar to the ones found for samples CLC and CLC-S(AR). Nevertheless, the peak of nickel aluminate for the sample exposed to consecutive stages of oxidation with air and N<sub>2</sub> (CLC-R(AN)) was not altered after reoxidation. The only redistribution of the active phase observed took place between NiO species with different interaction strength with the support. This could be also related with the different amounts of sulphur present in both regenerated samples.

The sulphided NiO18- $\alpha$ Al particles (BFB-S) had a different behaviour regarding the rest of the samples from both the CLC pilot plant and the batch fluidized bed reactor. A peak with a branch starting at  $\approx$ 633 K appeared at 555 K. A similar form of the TPR curve was reported by Kuhn et al. [36] for Ni-YSZ treated with H<sub>2</sub>S. However, after the in-situ oxidation, TPR profile was quite similar to the observed with the sample BFB-R(AN), indicating some reversibility of the sulphidation process.

In order to explain the TPR behaviour of these particles (BFB-S) and for comparison purposes, a TPR experiment was carried out with a mixture of 1 wt.% commercial nickel sulphide (Ni<sub>3</sub>S<sub>2</sub> Aldrich, 99.7% trace metals basis, -150 mesh) with the same  $\alpha$ -Al<sub>2</sub>O<sub>3</sub> used to prepare the NiO18- $\alpha$ Al oxygen-carrier. As can be seen in Figure 10, no peaks were detected for the Ni<sub>3</sub>S<sub>2</sub>/ $\alpha$ -Al<sub>2</sub>O<sub>3</sub> mixture and only a branch starting around 628 K, similar to that exhibited by the sample BFB-S, was observed. Thus, the behaviour of BFB-S sample could be explained in the same terms. In addition, the small peak at 555 K could be attributed to the presence of some NiS<sub>2</sub> in the sample, the less stable sulphide, as proposed by Afanasiev [37]. It must be considered in any case that the preparation of sample BFB-S is quite different to the samples existing in the CLC unit where a full reduction of the materials is never reached in the fuel-reactor. However, the lack of conclusive evidences for this hypothesis by TPR analysis could be clarified by the following XPS analysis.

#### **3.3.2.** X-ray photoelectron spectroscopy (XPS)

Binding energy regimes containing the Ni 2p and S 2p emission lines of the samples extracted from the CLC continuous plant are shown in Figure 11. The position of the Ni  $2p_{3/2}$  line varied from 855.4 to 855.6 eV. For these samples the position of the Ni  $2p_{1/2}$  line was found at 17.9 eV higher binding energy in comparison to Ni  $2p_{3/2}$  line. For both emission lines a shake-up satellite was found with a binding energy about 6.2-6.0 eV higher. The position of the Ni  $2p_{3/2}$  line around 855.5 eV with the associated satellite indicated the presence of one or more nickel-oxygen species. The fitting of the peaks showed the presence of several contributions whose BEs are given in Table 4. These BEs could be assigned to different Ni or S compounds according to data available in literature [36, 38-40].

Regarding to nickel peaks, sample CLC exhibited two contributions. The first one was attributed to the presence of Ni<sup>2+</sup> in octahedral sites in the form of NiO showing a weak interaction with the support whereas the second peak could be ascribed to Ni<sup>2+</sup> in the tetrahedral position of a spinel-like structure as NiAl<sub>2</sub>O<sub>4</sub>. This analysis confirmed the distribution of the impregnated nickel in the NiO18- $\alpha$ Al oxygen-carrier between bulk nickel oxide, easy to reduce owing to a low interaction with the  $\alpha$ -Al<sub>2</sub>O<sub>3</sub> support, and NiAl<sub>2</sub>O<sub>4</sub>, formed from the reaction between free NiO and the alumina support at high temperatures, as found in previous works [23, 27, 35] and in the preceding TPR analysis (Figure 8).

For the sulphur-containing samples CLC-S(FR) and CLC-S(AR), the first peak in the Ni 2p region shifted to a lower BE (Table 4), near to the one corresponding to Ni<sup>0</sup>. Nevertheless, the satellite peak is almost unaltered. Therefore, the presence of NiS instead of Ni<sup>0</sup> was considered. Magnus et al. [41] found that NiS is the preferred sulphide when the content of nickel over an alumina support is high. It is known that strong metal-support interactions give rise to the appearance of NiS whilst Ni<sub>3</sub>S<sub>2</sub> is preferentially found in systems with weak metal-support interactions [42]. This could be the case of the NiO18- $\alpha$ Al oxygen-carrier, with a very high content of nickel in comparison with the amount of sulphur in the solid particles and a strong interaction of the metal with the support to form NiAl<sub>2</sub>O<sub>4</sub>. However, curve fitting of Ni 2p<sub>3/2</sub> emission line gave a very small contribution of the case of the CLC sample without sulphur. It seems to indicate that sulphur mainly reacts with nickel in the form of NiO.

However, the most relevant result obtained with these sulphur-containing samples is that a new peak was found in the deconvolution of the Ni  $2p_{3/2}$  region. This peak at about 857 eV could be attributed to either Ni<sub>3</sub>S<sub>2</sub> phase, according to the Ni<sub>3</sub>S<sub>2</sub> standard in Kuhn et al. [36], or to sulphur-oxygen species. Arnoldy et al. [43] proposed a sulphidation mechanism of oxidic nickel catalysts by H<sub>2</sub>S through O-S exchange.

To get an insight into the species found on the different samples, XPS of the S 2p region was obtained and analysed. These results should be consistent with the Ni  $2p_{3/2}$  XPS results showed above. The position of the fitted peaks is presented in Table 4.

A peak and two broad bands of lower intensity were observed for the sample CLC-S(AR). The peak in the range 166-169 eV indicates the presence of Ni-O-S species according to previously cited literature [36, 40]. According to Buckley and Woods [40], the two bands could correspond to the doublet exhibited by the XPS pattern of Ni<sub>3</sub>S<sub>2</sub> standard. However, after curve fitting of the S 2p emission line, the separation of the BEs found for these samples was higher than that found for the standard sulphide. Moreover, the intensity and the area of the high BE contribution of the doublet found for the standard was lower than those for the low BE contribution. This could be due to the presence of other Ni-S species, such as NiS. These findings agreed with the conclusions made from the analysis of the Ni  $2p_{3/2}$  region, which suggested the presence of this sulphur compound in small amounts together with the major Ni<sub>3</sub>S<sub>2</sub>.

The sample CLC-S(FR) corresponding to the fuel-reactor of the 500  $W_{th}$  CLC unit plant showed a slightly different spectrum than that of the sample extracted from the airreactor. In this case, the intensity of the bands corresponding to nickel sulphides was higher while the intensity of the peak attributed to Ni-O-S species was lower. This result confirmed the proposed formation of nickel sulphide in the oxygen-carrier particles during the combustion of CH<sub>4</sub> in the presence of H<sub>2</sub>S inside the fuel-reactor [26]. It must be also remarked the presence of Ni-O-S species in the samples from the fuelreactor, which could be related with sulphate compounds.

The XPS spectra of the BFB reactor samples are shown in Figure 12. Sulphided sample (BFB-S) showed a Ni 2p XPS pattern similar to those of the samples extracted from the CLC continuous plant (see Figure 11), although the TPR profiles were very different. An XRD analysis of these samples confirmed the presence of metallic nickel, coming from the almost complete reduction of the sample during the sulphidation stage in the BFB reactor, and  $\alpha$ -Al<sub>2</sub>O<sub>3</sub>. However, the Ni 2p<sub>3/2</sub> line surprisingly appeared at 854.3 eV and the shake-up satellite indicated the presence of divalent species. On the other hand, line S 2p showed two wide peaks. First peak was found at 162.2 eV and corresponded to Ni<sub>3</sub>S<sub>2</sub> phase. The curve fitting of this peak gave the expected doublet for Ni<sub>3</sub>S<sub>2</sub> (161.6 and 162.8 eV). The selected operating conditions for the preparation (873 K, 1300 vppm of H<sub>2</sub>S) favoured the formation of a stable nickel sulphide as expected. The second band had the maximum at 169.1 eV, typical for Ni-O-S species. This finding was also quite surprising because the conditions in the BFB reactor were reducing instead oxidizing so this kind of compounds should not be observed. In this case, a

possible oxidation during sample manipulation could be inferred. To confirm that, an analysis of the composition of the inner part of the particles was carried out. The BFB-S sample was sputtered in situ at the same conditions previously reported. The Ni 2p and S 2p XPS lines can be seen in Figure 12. The Ni  $2p_{3/2}$  line shifted to a binding energy corresponding to metallic nickel and the satellite peak shrunk almost completely, indicating that divalent Ni species had almost disappeared. S 2p line showed only one peak related to Ni<sub>3</sub>S<sub>2</sub> phase. However, the concentration of this compound was very low because the intensity of the peak decreased with respect to the analysis of the surface. The peak at higher BE assigned to Ni-O-S species disappeared. Therefore, only the surface of the particles was oxidized. At this high BE only a shoulder corresponding to Ni<sub>3</sub>S<sub>2</sub> phase was found [36]. These results demonstrated that the nickel sulphide is formed preferably in the surface of the particles and that this could be easily oxidized. Finally, the sample regenerated in two stages with air and N<sub>2</sub> (BFB-R(AN)) was analysed. The XPS spectrum of Ni  $2p_{3/2}$  region showed one peak with two contributions: the first one could be attributed to Ni<sup>2+</sup> in octahedral sites as NiO with weak interaction with the support and the second peak could be assigned to  $Ni^{2+}$  in tetrahedral position of a spinel-like structure as NiAl<sub>2</sub>O<sub>4</sub>. As it has been previously mentioned, nickel inside NiO18-aAl particles was distributed between free NiO and NiAl<sub>2</sub>O<sub>4</sub>. So these results agreed with the previous TPR analysis. Despite the regeneration treatment, the S 2p region also showed the presence of sulphur compounds on the surface of the carrier, although twice less intense than the one corresponding to the BFB-S sample. The main contributions were assigned to Ni<sub>3</sub>S<sub>2</sub> phase and Ni-O-S species. The corresponding species could not be followed in Ni  $2p_{3/2}$  region due to the low intensity of the contribution of nickel sulphide to Ni  $2p_{3/2}$  peak compared to the intensity of the contribution of NiO and NiAl<sub>2</sub>O<sub>4</sub> species. In order to determine the evolution of sulphur inside the particles, this sample was also sputtered with an Ar ion gun. Ni  $2p_{3/2}$  peak after sputtering shifted to lower binding energies and the presence of different contributions was evident from the pattern. The satellite shrunk near 20%, indicating the presence of metallic nickel in some extent. This result could not be followed by TPR technique, because of the reasons mentioned above. The intensity of the S 2p peak after sputtering was near twice lower than that corresponding to the received sample and Ni-O-S species almost disappeared, being these results similar than those found for the BFB-S sample.

Although is difficult to quantify, the intensity of the peaks in both samples (as received and after sputtering) indicated that about 1/3 of the sulphur present in the sample remain inside the particle, mostly in the form of nickel sulphide. This result agrees with the found in the regeneration tests carried out in the BFB at different temperatures and oxygen concentrations, where the presence of sulphur inside the samples was also inferred.

Therefore, it can be concluded that the complete regeneration of the oxygen-carriers during oxidation in the air-reactor was not possible, and sulphur was being accumulating inside the particles. A previous desulphurization step of the fuel would be necessary before using it in a CLC plant if Ni-based oxygen-carriers are going to be used. In any case, it must be mentioned that regenerated particles, even containing some sulphur inside, were able to maintain the reactivity regarding CH<sub>4</sub> combustion reaction.

#### 4. Conclusions

Gaseous fuels for chemical-looping combustion (CLC) process can contain sulphurcompounds which affect the oxygen-carrier behaviour, especially if NiO is used as active phase. The loss of reactivity when NiO18- $\alpha$ Al oxygen-carrier was used in a CLC continuous unit with CH<sub>4</sub> containing H<sub>2</sub>S as fuel was attributed to the formation of nickel sulphides and sulphates. In this work the distribution of the sulphur in the solid particles extracted from each part of the plant after the operation ( $CH_4 + 500$  vppm  $H_2S$ for 11 h) was determined from a heating up process in a fluidized bed reactor with nitrogen or air atmospheres. 86% of the sulphur in the particles from the fuel-reactor was as Ni<sub>3</sub>S<sub>2</sub> while 14% corresponded to NiSO<sub>4</sub>. The amount of sulphur in the airreactor was reduced to almost a half and the ratio Ni<sub>3</sub>S<sub>2</sub>-NiSO<sub>4</sub> was reversed (16-84). Although the oxygen-carrier recovered from the riser kept the proportions between both sulphur compounds with respect to the sample of the air-reactor, the amount of sulphur increased almost to the levels in the fuel-reactor. Lower temperatures in this part of the unit could favour the reaction of the gaseous SO2 formed in the air-reactor with the NiO in the solid particles although this particularity of our installation will not be produced at industrial scale.

A regeneration study of the oxygen-carrier was also performed in order to determine the optimal situation to emit as much sulphur as possible in the air-reactor avoiding sulphur accumulation in the particles. Additional tests with a sulphided NiO18- $\alpha$ Al oxygen-

carrier (0.095 wt.%) previously prepared were carried out in a batch fluidized bed reactor at different temperatures (1073-1263 K) and oxygen concentrations (0-21 vol.%). Independently of the operating conditions, part of the sulphur remained in the solid and total regeneration was not possible. Nevertheless, the oxygen-carrier seemed to reach its initial reactivity after a regeneration step, so the remaining sulphur did not affect the behaviour of the oxygen carrier regarding  $CH_4$  combustion, although the oxygen transport capacity was decreased through the operation time.

The analysis of the NiO18- $\alpha$ Al oxygen-carrier samples using TPR and XPS techniques revealed that the sulphur reacted preferably with free NiO instead of NiAl<sub>2</sub>O<sub>4</sub>.

Ni3S2 was found to be the major sulphide in the samples obtained after the operation in the 500  $W_{th}$  CLC unit although minor amounts of NiS were also detected. It was also found that sulphur was preferably concentrated in the outer surface of the particles. TPR analysis showed that sulphur accumulated in the oxygen-carrier did not affect the reducibility behaviour. The sulphidation and later stages of oxidation with air and N<sub>2</sub> or only air did not alter significantly the Ni species distribution between NiO and NiAl<sub>2</sub>O<sub>4</sub>. Nevertheless, the presence of a low S 2p peak in the XPS analysis of the regenerated samples of sulphided NiO18- $\alpha$ Al oxygen-carrier indicated that total regeneration of the particles after their use with H<sub>2</sub>S did not take place. Taking into account all the results found, a previous desulphuration process of the fuel would be necessary when using NiO-based oxygen-carriers in the CLC system.

#### Acknowledgements

This research was conducted with financial support from the European Commission, under the Sixth Framework Program, Contract no. 019800-CLC GAS POWER, by the CCP2 (CO<sub>2</sub> Capture Project), a partnership of BP, Chevron, Conoco-Phillips, Eni Technology, Norsk Hydro, Shell, Suncor and Petrobras. The authors thank I. Fernández, N. Fernández and E. Ayllón their help with the XPS and TPR analyses of the samples.

#### References

[1] IPCC, Special report on carbon dioxide capture and storage, Prepared by Working Group III of the Intergovernmental Panel on Climate Change [Metz, B., O. Davidson, H. C. de Coninck, M. Loos, and L. A. Meyer (eds.)], Cambridge University Press, Cambridge, United Kingdom and New York, NY, USA, 2005.

[2] J. Adánez, A. Abad, F. García-Labiano, P. Gayán, L.F.d. Diego, Prog. Energ. Combust. 38 (2012) 215-282.

[3] M.M. Hossain, H.I. de Lasa, Chem. Eng. Sci. 63 (2008) 4433-4451.

[4] A. Lyngfelt, M. Johansson, T. Mattisson, in: 9th International Conference on Circulating Fluidized Beds (CFB-9), Hamburg, Germany, 2008.

[5] J. Adánez, L.F. de Diego, F. García-Labiano, P. Gayán, A. Abad, J.M. Palacios, Energ. Fuel 18 (2004) 371-377.

[6] J. Adánez, F. García-Labiano, L.F. De Diego, P. Gayán, J. Celaya, A. Abad, Ind.Eng. Chem. Res. 45 (2006) 2617-2625.

[7] M. Ishida, M. Yamamoto, T. Ohba, Energ. Convers. Manage. 43 (2002) 1469-1478.
[8] E. Jerndal, T. Mattisson, I. Thijs, F. Snijkers, A. Lyngfelt, Int. J. Greenh. Gas Control 4 (2010) 23-35.

[9] R. Villa, C. Cristiani, G. Groppi, L. Lietti, P. Forzatti, U. Cornaro, S. Rossini, J Mol. Catal. A-Chem. 204-205 (2003) 637-646.

[10] T. Mattisson, M. Johansson, A. Lyngfelt, Fuel 85 (2006) 736-747.

[11] D. Bucker, D. Holmberg, T. Griffin, Technoeconomic evaluation of an oxyfuel power plant using mixed conducting membranes., in: Carbon Dioxide Capture for Storage in Deep Geological Formations-Results from the CO<sub>2</sub> Capture Project, Elsevier, Oxford, 2005.

[12] A. Lyngfelt, B. Kronberger, J. Adánez, J.X. Morin, P. Hurst, E.S. Rubin, D.W. Keith, C.F. Gilboy, M. Wilson, T. Morris, J. Gale, K. Thambimuthu, The grace project: Development of oxygen carrier particles for chemical-looping combustion. Design and operation of a 10 kW chemical-looping combustor, in: Greenhouse Gas Control Technologies 7, Elsevier Science Ltd, Oxford, 2005, pp. 115-123.

[13] D. Hebden, H. Stroud, Coal gasification processes, in: Chemistry of Coal Utilization, John Wiley & Sons, New York, 1981.

[14] E. Jerndal, T. Mattisson, A. Lyngfelt, Chem. Eng. Res. Des. 84 (2006) 795-806.

[15] B. Wang, R. Yan, D.H. Lee, D.T. Liang, Y. Zheng, H. Zhao, C. Zheng, Energ. Fuel 22 (2008) 1012-1020.

[16] R.D. Solunke, G. Veser, Fuel 90 (2011) 608-617.

[17] R.D. Solunke, G. Veser, Energ. Fuel 23 (2009) 4787-4796.

[18] H. Tian, T. Simonyi, J. Poston, R. Siriwardane, Ind. Eng. Chem. Res. 48 (2009) 8418-8430.

[19] L. Shen, Z. Gao, J. Wu, J. Xiao, Comb. Flame 157 (2010) 853-863.

[20] E. Ksepko, R.V. Siriwardane, H. Tian, T. Simonyi, J.A.P. Jr., A. Zinn, M. Sciazko,

in: Les Rencontres Scientifiques de l'IFP - 1<sup>st</sup> International Conference on Chemical Looping, Lyon, Francia, 2010.

[21] C.R. Forero, P. Gayán, F. García-Labiano, L.F. de Diego, A. Abad, J. Adánez, Int.

J. Greenh. Gas Control 4 (2010) 762-770.

[22] K. Mayer, in: 3rd High Temperature Solid Looping Network Meeting, 30th August-1st September, Vienna (Austria), 2011.

[23] J. Adanez, C. Dueso, L.F. de Diego, F. Garcia-Labiano, P. Gayan, A. Abad, Energ.Fuel 23 (2009) 130-142.

[24] C. Dueso, F. García-Labiano, J. Adánez, L.F. de Diego, P. Gayán, A. Abad, Fuel88 (2009) 2357-2364.

[25] J. Adánez, C. Dueso, L.F. de Diego, F. García-Labiano, P. Gayán, A. Abad, Ind.Eng. Chem. Res., 48 (2009) 2509-2518.

[26] F. Garcia-Labiano, L.F. de Diego, P. Gayan, J. Adanez, A. Abad, C. Dueso, Ind. Eng. Chem. Res. 48 (2009) 2499-2508.

[27] P. Gayán, C. Dueso, A. Abad, J. Adánez, L.F. de Diego, F. García-Labiano, Fuel88 (2009) 1016-1023.

[28] L.F. de Diego, P. Gayán, F. García-Labiano, J. Celaya, A. Abad, J. Adánez, Energ. Fuel 19 (2005) 1850-1856.

[29] HSC Chemistry 6.1, Chemical reaction and equilibrium software with thermochemical database and simulation module, Oututec Research, Pori, Finland, 2008.

[30] C. Li, Y.-W. Chen, Thermochim. Acta 256 (1995) 457-465.

[31] B. Mile, D. Stirling, M.A. Zammitt, A. Lovell, M. Webb, J. Catal. 114 (1988) 12.

[32] A. Kadkhodayan, A. Brenner, J.Catal. 117 (1989) 10.

[33] F. Patcas, D. Honicke, Catal. Commun. 6 (2005) 4.

[34] J. Wang, L. Dong, Y. HU, G. Zheng, Z. Hu, Y. Cheng, J. Solid State Chem. 157(2001) 274-282.

- [35] C. Dueso, A. Abad, F. García-Labiano, L.F. de Diego, P. Gayán, J. Adánez, A. Lyngfelt, Fuel 89 (2010) 3399-3409.
- [36] J.N. Kuhn, N. Lakshminarayanan, U.S. Ozkan, J. Mol. Catal.s A-Chem. 282 (2008)12.
- [37] P. Afanasiev, Appl. Catal. A-Gen. 303 (2006) 110-115.
- [38] C.D. Wagner, A.V. Naumkin, A. Kraut-Vass, J.W. Allison, C.J. Powell, J.R.
- Rumble, in, NIST (National Institute of Standards and Technology), 2003.
- [39] C.D. Wagner, W.M. Riggs, L.E. Davis, J.F. Moulder, Handbook of X-ray
- Photoelectron Spectroscopy, Perkin-Elmer Corp, 1979.
- [40] A.N. Buckley, R. Woods, J. Mol. Catal. A-Chem. 21 (1991) 575-582.
- [41] P.J. Mangnus, E.K. Poels, A.D.V. Langeveld, J.A. Moulijn, J. Catal. 137 (1992)92-101.
- [42] S.P.A. Louwers, R. prins, J. Catal. 133 (133) 94-111.
- [43] P. Arnoldy, J.A.M.V.D. Heijkant, D.D.d. Bok, J.A. Moulijn, J. Catal. 92 (1985) 35-55.

Table 1. Properties of the fresh NiO18- $\alpha$ Al oxygen-carrier.

NiO content (wt.%)	18
Particle size (µm)	100-300
Apparent density (g/cm <sup>3</sup> )	2.5
BET surface area (m <sup>2</sup> /g)	7
Porosity (%)	43
Mechanical strength (N)	4.1
XRD phases	α-Al <sub>2</sub> O <sub>3</sub> , NiO, NiAl <sub>2</sub> O
Oxygen transport capacity	0.04

	Sulphur in the sample	Sulphur distribution (%	
	(mg S/100 g sample)	$Ni_3S_2$	NiSO <sub>4</sub>
Fuel-reactor	11.6	86	14
Air-reactor	5.9	16	84
Riser	11.2	17	83

**Table 2.** Sulphur distribution in the oxygen-carrier samples extracted from the CLCcontinuous unit after  $CH_4$  combustion experiments in the presence of  $H_2S$ .

\* Inferred from SO<sub>2</sub> release in BFB according to reactions 1-4.

Sample	TPR-1	TPR-2 <sup>*</sup>
Fresh	721, 873, 1141	622, 710, 873, 1141
CLC	647, 683, 763, 1217	620, 705, 1078
CLC-S(FR)	583, 658, 833, 1116	626, 700, 875, 1133
CLC-S(AR)	689, 735, 1135	629, 721, 1086
BFB-S	556	556
BFB-R(A)	719, 821, 1162	729, 853, 922, 1123
BFB-R(AN)	711, 803, 1185	722, 856, 1167
* After a TPO ru	n	

**Table 3.** Position of the peaks and bands in TPR analysis. Data in K.

	Ni 2p region BE (eV)			S 2p region		
Sample				BE (eV)		
CLC	853.6	855.6				
CLC-S(FR)	852.6	855.4	856.6	160.0	163.2	168.5
CLC-S(AR)	853.2	855.5	856.7	160.5	163.1	168.5
BFB-S	854.3			161.6	162.8	169.1
BFB-S*	851.9			160.7	162.7	
BFB-R(AN)	854.0	855.7		161.0	162.6	167.6
BFB-R(AN)*	851.8	852.6	855.6	160.5	163.3	167.4

**Table 4.** Surface analysis from XPS spectra.

\* After sputtering

Captions of Figures

**Figure 1.** Temperature and gas concentration profiles during an experiment in the 500  $W_{th}$  CLC continuous pilot plant. CH<sub>4</sub> = 30 vol.%, H<sub>2</sub>S = 500 vppm, T<sub>FR</sub> = 1143 K, T<sub>AR</sub> = 1223 K, G<sub>s</sub> = 12 Kg/h.

**Figure 2.**  $SO_2$  concentration released during the heating of the NiO18- $\alpha$ Al oxygencarrier extracted from the fuel-reactor, air-reactor and riser in air and nitrogen atmosphere.

**Figure 3.** Effect of temperature on the sulphur emitted as  $SO_2$  during successive regeneration stages with air and nitrogen with sulphided samples of NiO18- $\alpha$ Al oxygen-carrier.

Figure 4. Thermodynamic equilibrium for NiSO<sub>4</sub> decomposition.

**Figure 5.** Effect of oxygen concentration on the sulphur emitted as SO<sub>2</sub> during successive regeneration stages with  $O_2$ - $N_2$  mixtures and nitrogen with sulphided samples of NiO18- $\alpha$ Al oxygen-carrier. T= 1223 K.

**Figure 6.** TGA reactivity for the fresh and the regenerated oxygen carrier. T = 1223 K. 15 vol.% CH<sub>4</sub> during reduction; air during oxidation.

**Figure 7.** TPR profile of the fresh NiO18- $\alpha$ Al oxygen-carrier ( \_\_\_\_\_) and after a TPO run(- - -). Theoretical temperature values of the different structures obtained from references [30-33].

**Figure 8.** TPR profiles of the samples extracted from the CLC continuous pilot plant burning a fuel containing sulphur: (\_\_\_\_) Fresh oxygen-carrier, (\_\_\_.) CLC, (\_\_\_.) CLC-S(AR), and (\_\_\_.) CLC-S(FR).

**Figure 9.** TPR profiles of the samples prepared in the BFB reactor: (\_\_\_\_) fresh oxygencarrier, (\_\_\_) sulphided particles BFB-S, (\_\_\_\_) after regeneration with air BFB-R(A), (\_\_\_) and after regeneration with air and  $N_2$ , BFB-R(AN).

**Figure 10.** TPR profiles of the sulphided NiO18- $\alpha$ Al oxygen-carrier (BFB-S) and the pattern sample (1 wt.% Ni<sub>3</sub>S<sub>2</sub>/ $\alpha$ -Al<sub>2</sub>O<sub>3</sub>).

**Figure 11.** XPS surface spectra for Ni 2p and S 2p of the samples extracted from the CLC continuous pilot plant during combustion tests.

**Figure 12.** XPS surface spectra for Ni 2p and S 2p of sulphided sample prepared in the batch fluidized bed reactor, BFB-S.

**Figure 13.** XPS surface spectra for Ni 2p and S 2p of regenerated sample prepared in the batch fluidized bed reactor, BFB-R(AN).



**Figure 1.** Temperature and gas concentration profiles during an experiment in the 500  $W_{th}$  CLC continuous pilot plant. CH<sub>4</sub> = 30 vol.%, H<sub>2</sub>S = 500 vppm, T<sub>FR</sub> = 1143 K, T<sub>AR</sub> = 1223 K, G<sub>s</sub> = 12 Kg/h.



**Figure 2.**  $SO_2$  concentration released during the heating of the NiO18- $\alpha$ Al oxygencarrier extracted from the fuel-reactor, air-reactor and riser in air and nitrogen atmosphere.



Figure 3. Effect of temperature on the sulphur emitted as  $SO_2$  during successive regeneration stages with air and nitrogen with sulphided samples of NiO18- $\alpha$ Al oxygen-carrier.



Figure 4. Thermodynamic equilibrium for NiSO<sub>4</sub> decomposition.



**Figure 5.** Effect of oxygen concentration on the sulphur emitted as  $SO_2$  during successive regeneration stages with  $O_2$ - $N_2$  mixtures and nitrogen with sulphided samples of NiO18- $\alpha$ Al oxygen-carrier. T= 1223 K.



**Figure 6.** TGA reactivity for the fresh and the regenerated oxygen carrier. T = 1223 K. 15 vol.% CH<sub>4</sub> during reduction; air during oxidation.



**Figure 7.** TPR profile of the fresh NiO18-αAl oxygen-carrier ( \_\_\_\_\_) and after a TPO run(- - -). Theoretical temperature values of the different structures obtained from references [30-33].



**Figure 8.** TPR profiles of the samples extracted from the CLC continuous pilot plant burning a fuel containing sulphur: (\_\_\_\_) Fresh oxygen-carrier, (\_\_\_\_) CLC, (\_\_\_\_) CLC-S(AR), and (\_\_\_\_) CLC-S(FR).



**Figure 9.** TPR profiles of the samples prepared in the BFB reactor: (\_\_\_\_) fresh oxygencarrier, (\_\_\_) sulphided particles BFB-S, (\_\_\_\_) after regeneration with air BFB-R(A), (\_\_\_) and after regeneration with air and N<sub>2</sub>, BFB-R(AN).



Figure 10. TPR profiles of the sulphided NiO18- $\alpha$ Al oxygen-carrier (BFB-S) and the pattern sample (1 wt.% Ni<sub>3</sub>S<sub>2</sub>/ $\alpha$ -Al<sub>2</sub>O<sub>3</sub>).



**Figure 11.** XPS surface spectra for Ni 2p and S 2p of the samples extracted from the CLC continuous pilot plant during combustion tests.



**Figure 12.** XPS surface spectra for Ni 2p and S 2p of sulphided sample prepared in the batch fluidized bed reactor, BFB-S.



**Figure 13.** XPS surface spectra for Ni 2p and S 2p of regenerated sample prepared in the batch fluidized bed reactor, BFB-R(AN).

### **Graphical Abstract**



## Highlights

 $H_2S$  in the fuel affects the reactivity of NiO-based oxygen-carriers. > Sulfur reacts preferentially with NiO in the particles instead of NiAl<sub>2</sub>O<sub>4</sub>. > Ni<sub>3</sub>S<sub>2</sub>, NiS and small amounts of NiSO<sub>4</sub> were found in the fuel-reactor particles. > NiSO<sub>4</sub> and small amounts of Ni<sub>3</sub>S<sub>2</sub> were found in the air-reactor particles. > Total regeneration of the accumulated sulfur in the particles was not possible.