

# The Oxy-CaL process: A novel CO<sub>2</sub> capture system by integrating partial oxy-combustion with the Calcium-Looping process

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## Abstract

This paper proposes a novel CO<sub>2</sub> capture technology from the integration of partial oxy-combustion and the Calcium Looping capture process based on the multicycle carbonation/calcination of limestone derived CaO. The concentration of CO<sub>2</sub> in the carbonator reactor is increased by means of partial oxy-combustion, which enhances the multicycle CaO conversion according to Thermogravimetric analysis results carried out in our work, thus improving the CO<sub>2</sub> capture efficiency. On the other hand, energy consumption for partial oxy-combustion is substantially reduced as compared to total oxy-combustion. All in all, process simulations indicate that the integration of both processes has potential advantages mainly regarding power plant flexibility whereas the overall energy penalty is not increased. Thus, the resulting energy consumption per kilogram of CO<sub>2</sub> avoided is kept below 4 MJ/kg CO<sub>2</sub>, which remains below the typical values reported for total oxy-combustion and amine based CO<sub>2</sub> capture systems whereas CO<sub>2</sub> capture efficiency is enhanced in comparison with the Calcium Looping process.

**Keywords:** Calcium-Looping, oxy-combustion, CCS, SPECCA

## 1. Introduction

Post-combustion CO<sub>2</sub> capture and storage (CCS) is considered as one key strategy to mitigate global warming [1,2]. In order to achieve a commercial deployment of CO<sub>2</sub> capture in fossil fuel power plants, several technologies are being analyzed aimed mainly at maximizing the capture efficiency while energy penalty and capital cost are minimized [3,4]. Among diverse possibilities, already commercial amine-based capture systems and the Calcium-Looping (CaL) process, currently under pilot-scale stage, have attracted a great deal of attention in the last years [5,6]. Although CO<sub>2</sub> capture by chemical absorption using MEA (monoethanolamine) is a well-established process in industry, the commercial deployment of this technology for post-combustion CO<sub>2</sub> capture at the large scale is hindered by the high energy penalty (8-12%) mainly due to sorbent regeneration [7–9], amine toxicity [10] and degradation [11].

45

46 The CaL process is based on the carbonation/calcination reaction of solid CaO particles,  
47 which is carried out in two interconnected circulating fluidized bed (CFB) reactors [12]. This  
48 second generation capture technology has several potential advantages when compared with  
49 amine scrubbing such as a higher CO<sub>2</sub> capture efficiency (above 90%), lower energy penalty  
50 over the power plant (4-9%) [6] and the use of low cost, widely available and non-toxic  
51 natural minerals as CaO precursors such as limestone or dolomite [13]. Even though several  
52 pilot scale plants (~ 1-2 MW<sub>th</sub>) are already showing promising results [14,15] the CaL  
53 technology has not reached a demonstration stage yet. The main causes that hinder such step  
54 forward are linked to the excessively large size of the capture system (carbonator reactor  
55 height ~ 40 m; carbonator solids inventory ~ 400 ton; additional coal consumption for CO<sub>2</sub>  
56 capture ~ 45-55%), which increases significantly both capital and operating costs (CAPEX  
57 and OPEX) for power generation [16,17].

58

59 Another interesting possibility to mitigate CO<sub>2</sub> emissions from power plants is the oxy-  
60 combustion technology, which has been successfully demonstrated in large-scale pilot  
61 projects (30 MW<sub>e</sub>) [18–20]. Essentially, oxy-combustion consists of replacing air by pure O<sub>2</sub>  
62 (mixed with CO<sub>2</sub>-rich flue gas recycled) as combustion gas, which yields a highly-  
63 concentrated CO<sub>2</sub> flue gas stream. After water condensation and purification, the CO<sub>2</sub> stream  
64 (~95% vol) is suitable for compression and storage or utilization [21]. The main drawback for  
65 the commercial deployment of oxy-combustion is the high energy consumption for pure O<sub>2</sub>  
66 production in the air separation unit (ASU) via cryogenic distillation, which causes an energy  
67 penalty in the range of 7–13% [22,23] or, equivalently, over 20% additional fuel  
68 consumption for power production.

69

70 In this paper a novel system (Oxy-CaL) for CO<sub>2</sub> capture is investigated based on the  
71 combination of partial oxy-combustion and the CaL process with the goal of exploiting the  
72 synergies between such technologies. Basically, Oxy-CaL consists of carrying out a partial  
73 oxy-combustion process to produce a flue gas with a CO<sub>2</sub> concentration in the range 30-60%  
74 vol, which is then sent to the CaL capture process. In a similar way, other authors have  
75 analyzed the integration of partial oxy-combustion and MEA [24], which is expected to help  
76 mitigating MEA degradation and energy consumption.

77

78 The manuscript starts by showing experimental results from a thermo-gravimetric analysis  
79 (TGA) on the multicycle conversion of limestone derived CaO under realistic calcination  
80 conditions (high temperature and high CO<sub>2</sub> concentration). In these TGA tests, the CO<sub>2</sub>  
81 concentration in the carbonation environment was varied in the range 15-60% vol in order to  
82 address the effect of an excess of CO<sub>2</sub> in the carbonator over the typical vol% in the flue gas  
83 at typical combustion conditions (~15%). Moreover, the carbonation temperature was varied  
84 in the range 625-680°C, which affects critically the carbonation kinetics in the solid-state  
85 diffusion-controlled stage as will be seen. These TGA results are used afterwards in the Oxy-  
86 CaL integration model to calculate the CO<sub>2</sub> capture efficiency from process simulations. The  
87 energy penalty arising from the diverse CO<sub>2</sub> capture technologies considered (total oxy-

88 combustion, CaL and Oxy-CaL) is analyzed. Finally, the oxy-CaL performance is assessed  
89 and compared with those of other CO<sub>2</sub> capture systems.

90

91 Our results show that the Oxy-CaL system is a promising hybrid concept to be applicable in  
92 new power plants, allowing for a substantial reduction of energy penalty as compared to total  
93 oxy-fuel combustion. Moreover, the Oxy-CaL system leads to a high CO<sub>2</sub> capture efficiency  
94 in comparison with the CaL process, which would serve to reduce significantly the carbonator  
95 reactor size.

96

## 97 **2. Thermogravimetric analysis**

### 98 **a. Materials and Methods**

99 The material employed in this work was natural limestone of high purity (99.6% wt CaCO<sub>3</sub>),  
100 received from Segura S.L (Matagallar quarry, Pedrera, Spain). Carbonation/calcination  
101 multicycle tests were carried out using a thermogravimetric analyzer TGA Discovery (TA  
102 Instruments 2011) equipped with an infrared halogen lamp furnace wherein the lamps are  
103 placed symmetrically with respect to a SiC enclosure to minimize undesired heat transfer  
104 phenomena. This setup allows for high heating/cooling rates (300°C/min), which is a  
105 necessary requirement to mimic realistic conditions in the CaL process where the solids are  
106 rapidly circulated between the reactors. The TGA instrument is also equipped with a high  
107 sensitivity balance (<0.1 µg) characterized by a small baseline dynamic drift (<10 µg). A  
108 thermocouple is located close to the sample and underneath it for a reliable measurement and  
109 control of temperature in the sample.

110

111 TGA experiments consisted of 20 carbonation/calcination cycles preceded by a precalcination  
112 of the sample at 950°C for 5 minutes under a 30% air/70% CO<sub>2</sub> vol/vol. atmosphere. Then,  
113 the temperature is decreased at 300°C/min to introduce the carbonation stage at the desired  
114 temperature and under a given CO<sub>2</sub>/air mixture. After that, the sample is calcined for CaO  
115 regeneration by quickly increasing the temperature at 300°C/min to 950°C under a high CO<sub>2</sub>  
116 concentration environment (30% air/70% CO<sub>2</sub> vol/vol.) as representative in the calciner  
117 environment [25,26]. Short residence times of 5 minutes for both calcination and carbonation  
118 stages have been employed as corresponds to realistic conditions. In order to mimic the  
119 integration of the CaL process with oxy-fuel combustion (Oxy-CaL), four different CO<sub>2</sub>/air  
120 mixtures were tested for the carbonation stage: 15% CO<sub>2</sub>/85% air, 30% CO<sub>2</sub>/70% air, 45%  
121 CO<sub>2</sub>/55% air and 60% CO<sub>2</sub>/40% air (vol/vol). Three different carbonation temperatures  
122 (625°C, 650°C and 680°C) were used. Tests are labeled as CaL-T for those in which  
123 carbonation was performed under 15% CO<sub>2</sub>, where T stands for the carbonation temperature,  
124 and Oxy-CaL-vol-T, where vol is the vol% of CO<sub>2</sub> in the CO<sub>2</sub>/air mixture and T is the  
125 carbonation temperature. Samples of small and fixed mass (~10 mg) were employed to avoid  
126 mass transfer resistance within the sample. Intraparticle pore diffusion limitations on the  
127 reaction rate are also avoided by using particles of size below 100 µm [27,28].

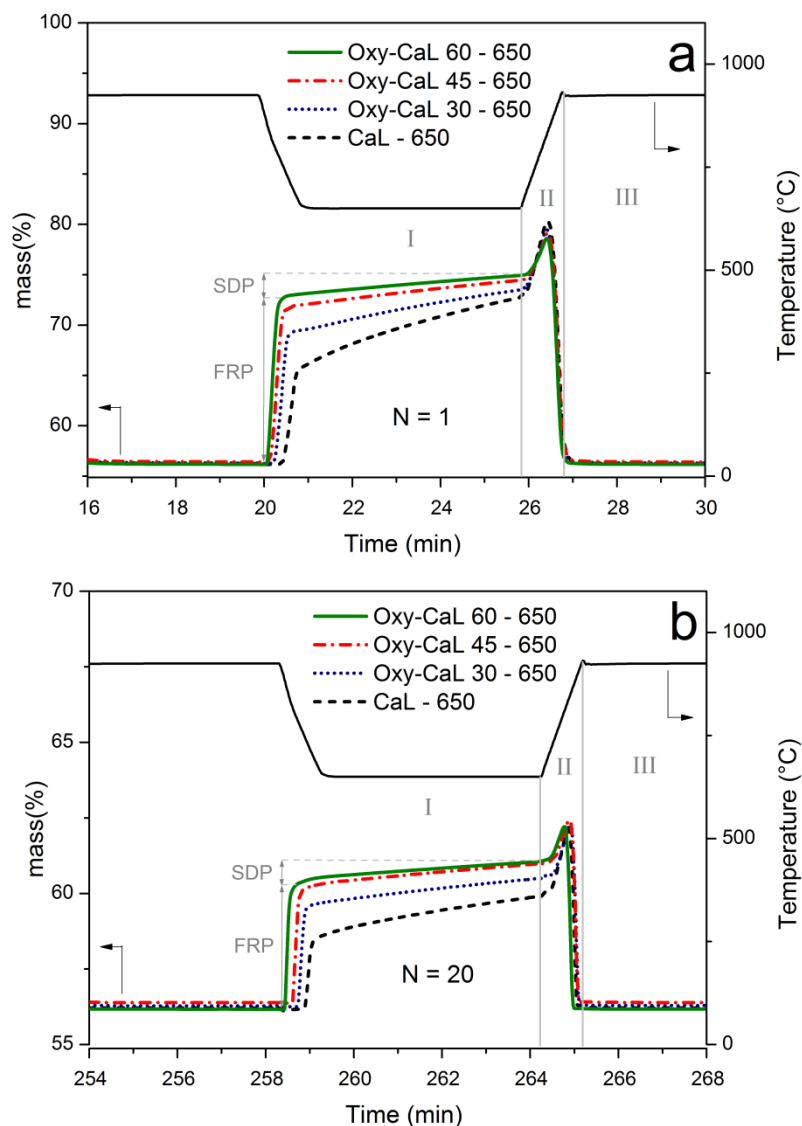
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## **b. Experimental results and discussion**

129 Figure 1 shows examples of thermograms obtained for the 1<sup>st</sup> (Figure 1a) and 20<sup>th</sup> (Figure 1b)  
130 carbonation/calcination cycles under CaL (15% vol CO<sub>2</sub> carbonation) and Oxy-CaL (30%,  
131 45% and 60% vol CO<sub>2</sub> carbonation) conditions. These thermograms show the time evolution  
132 of temperature and sample mass % along the cycles and illustrate already an important effect  
133 of raising the carbonation CO<sub>2</sub> vol concentration on the reaction kinetics. As well-known  
134 from previous studies CaO carbonation is seen to take place along two well-differentiated  
135 stages [29–31]. The first stage consists of a reaction-controlled fast phase on the surface of the  
136 particles that ends up when a 30-50 nm thick carbonate layer is built up on the CaO surface  
137 [29]. This first phase is followed up by a second slower phase limited by the solid-state  
138 diffusion of CO<sub>3</sub><sup>2-</sup> mobile ions and counter-current diffusion of O<sup>2-</sup> anions across the CaCO<sub>3</sub>  
139 product layer [31,32].

140 As may be seen in Figure 1, carbonation in the fast phase is markedly enhanced as the CO<sub>2</sub>  
141 concentration is increased, whereas, on the contrary, diffusion-controlled carbonation is  
142 markedly hindered. Thus, carbonation in the diffusion-controlled stage contributes  
143 significantly to the overall capture capacity under CaL conditions but loses relevancy as the  
144 CO<sub>2</sub> vol% is increased. As will be seen from process simulations this effect on the  
145 carbonation kinetics has remarkable implications on the role of key process operation  
146 parameters such as the solids residence time in the carbonator.

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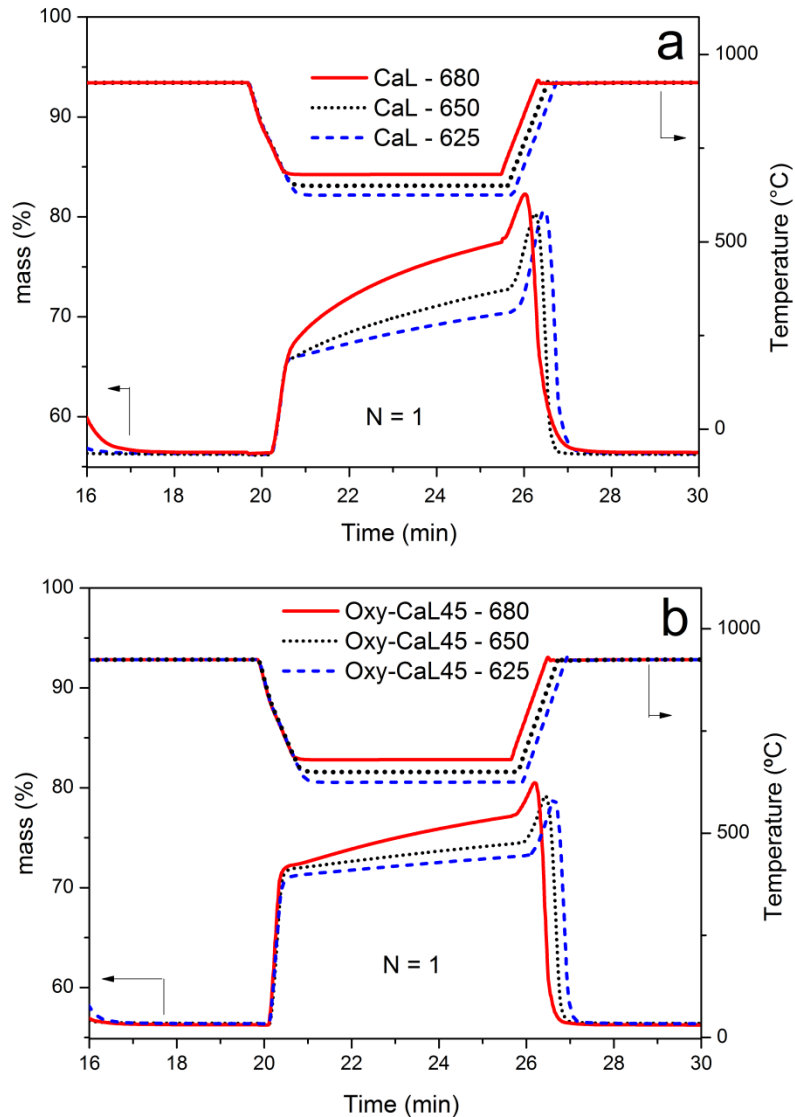


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150 Figure 1. Time evolution of the sorbent mass % during the carbonation and calcination stages in the a) 1<sup>st</sup>  
151 cycle (N=1) and b) 20<sup>th</sup> cycle (N=20) for limestone tested under CaL (15% vol CO<sub>2</sub> carbonation) and Oxy-  
152 CaL (30%, 45%, 60% vol CO<sub>2</sub> carbonation) conditions. Carbonation temperature is fixed to 650°C. I  
153 indicates the carbonation stage, II the transition stage and III the calcination stage. Mass gain in the two  
154 phases of carbonation (fast reaction-controlled phase FRP and solid-state diffusion controlled phase SDP)  
155 are indicated.

156 Figure 2a shows the thermograms corresponding to the 1<sup>st</sup> cycle obtained from TGA tests  
157 performed at different carbonation temperatures (625, 650 and 680°C) under CaL conditions  
158 (15% vol CO<sub>2</sub> carbonation). As may be observed, a variation of just about 25°C around the typical  
159 carbonation temperature used in pilot-scale plants (~650°C) has a significant effect on the CO<sub>2</sub>  
160 uptake in the diffusion-controlled stage, which notably affects the overall capture capacity. Thus,  
161 carbonation in this phase is enhanced with temperature while a decrease of the carbonation  
162 temperature yields a rapid decay of the carbonation rate in this solid-state diffusion-controlled  
163 stage. This result is consistent with the strong dependence on temperature measured elsewhere for  
164 C<sup>14</sup> isotope diffusivity in CaO and for the effective product layer diffusivity of CO<sub>3</sub><sup>2-</sup> mobile ions

165 in the range of carbonation temperatures used in our work [30,33]. A similar behavior has been  
 166 observed for the samples tested under Oxy-CaL conditions. Nonetheless, the variation with the  
 167 temperature of the CO<sub>2</sub> capture capacity in the diffusion-controlled stage plays a relatively minor  
 168 role on the overall capture capacity under Oxy-CaL conditions as compared to CaL conditions.  
 169 This may be seen in Figure 2b, which shows the 1<sup>st</sup> cycle of the thermograms obtained under  
 170 different carbonation temperatures for the sample tested under Oxy-CaL 45 conditions.  
 171



172  
 173 Figure 2. Time evolution of the sorbent mass % during the 1<sup>st</sup> cycle (N=1) for limestone under (a) CaL  
 174 (15% vol CO<sub>2</sub> carbonation) and (b) Oxy-CaL 45 (45% vol CO<sub>2</sub> carbonation) conditions for different  
 175 carbonation temperatures (625°C, 650°C and 680°C) as indicated.

176  
 177 The parameter used to compare the multicycle capture performance of limestone under CaL  
 178 and Oxy-CaL conditions is CaO conversion, defined as the ratio of CaO mass converted to  
 179 CaCO<sub>3</sub> in the carbonation stage of each cycle to the sorbent mass before carbonation.  
 180 Multicycle CaO conversion data can be generally well fitted by the following semi-empirical  
 181 equation [6,34,35]:

182

$$X_N = X_r + \frac{X_1}{k(N-1) + \left(1 - \frac{X_r}{X_1}\right)^{-1}}; \quad (N = 1, 2, \dots) \quad (1)$$

183

184 where  $N$  is the cycle number,  $X_1$  is CaO conversion at the first cycle,  $k$  is the deactivation rate  
 185 constant and  $X_r$  is the residual CaO conversion, which is approached asymptotically after a  
 186 very large number of cycles. Figure 3a shows multicycle CaO conversion data and best fit  
 187 curves from equation (1) for the samples tested under CaL and Oxy-CaL conditions for a  
 188 carbonation temperature of 650°C. Best fitting parameters are summarized in Table 1. As well  
 189 known, CaO conversion decreases progressively with the cycle number due to enhanced grain  
 190 sintering in the calcination stage at high temperature and under high CO<sub>2</sub> partial pressure  
 191 [6,36,37], which reduces the CaO surface area available for fast carbonation in each cycle.  
 192 Note however, that the deactivation rate is decreased as the CO<sub>2</sub> concentration in the  
 193 carbonation stage is increased. Thus, the residual conversion  $X_r$  takes values of 0.062, 0.070,  
 194 0.076 and 0.081 for the samples tested under CaL-650, Oxy-CaL 30-650, Oxy-CaL 45-650  
 195 and Oxy-CaL 60-650 conditions, respectively.

196

197 The relative contributions to the overall CaO conversion of the fast reaction controlled phase  
 198 (FRP) and solid-state diffusion controlled phase (SDP) have been analyzed by extracting from  
 199 the thermograms the values of CaO conversion in each one of these phases ( $X_{FRP}$  and  $X_{SDP}$ ,  
 200 respectively, see Figure 1a). Data on  $X_{FRP}$  and  $X_{SDP}$  are shown in Figure 3b and Figure 3c. As  
 201 was inferred from Figure 2, it is seen that  $X_{FRP}$  becomes increasingly relevant while  $X_{SDP}$  is  
 202 decreased as the CO<sub>2</sub> concentration in the carbonation environment is increased.

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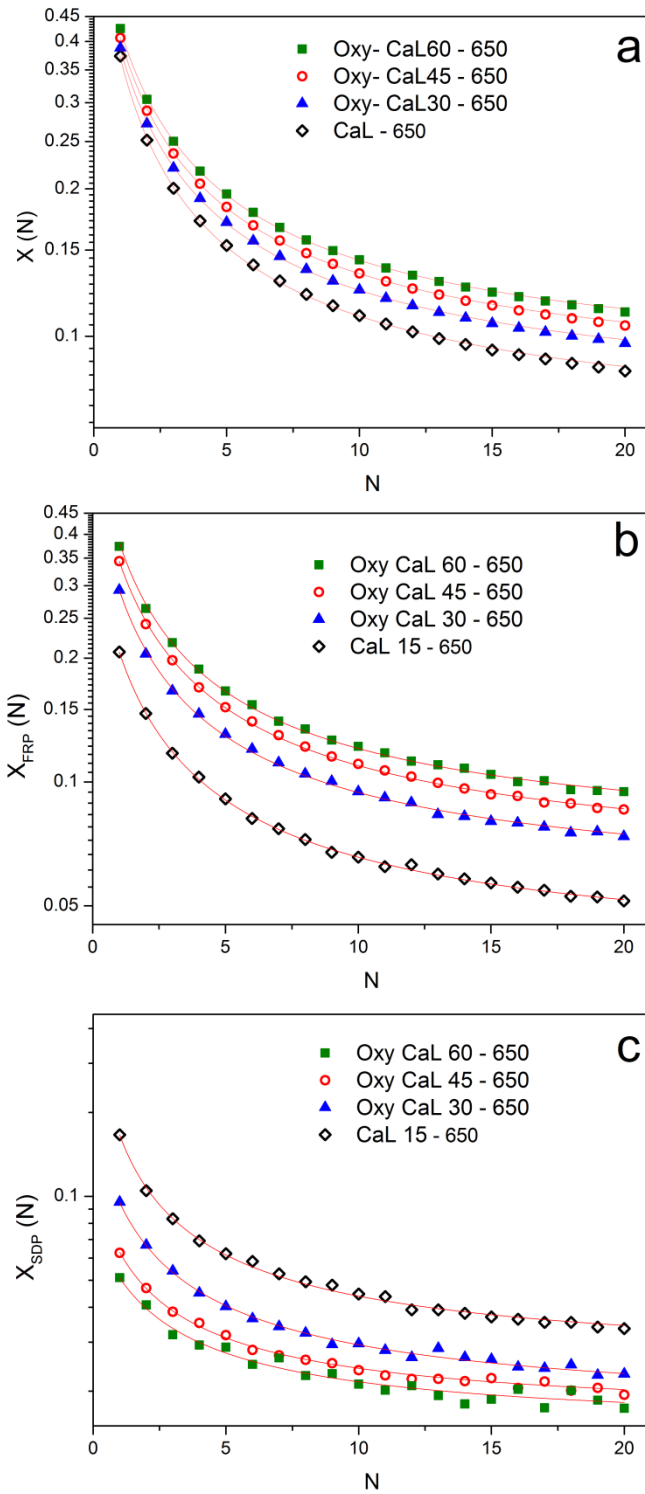
204 Table 1: Values of the deactivation rate constant  $\kappa$  and residual conversion  $X_r$  obtained from the best fits of  
 205 Eq. (1) to TGA experimental data for carbonation at different CO<sub>2</sub> concentrations (15% vol in the CaL  
 206 tests; 30%, 45%, and 60% vol in the Oxy-CaL tests).

T carb =650°C		CaL-650	Oxy-CaL30-650	Oxy-CaL45-650	Oxy-CaL60-650
$X_{overall}$	$X_1$	0.373	0.388	0.407	0.425
	$\kappa$	0.731	0.667	0.651	0.633
	$X_r$	0.061	0.070	0.076	0.081
	$R_{sqr}$	0.999	0.999	0.999	0.999
$X_{FRP}$	$X_1$	0.207	0.293	0.344	0.374
	$\kappa$	0.660	0.661	0.641	0.674
	$X_r$	0.037	0.053	0.060	0.068
	$R_{sqr}$	0.999	0.999	0.999	0.998
$X_{SDP}$	$X_1$	0.166	0.095	0.063	0.051
	$\kappa$	0.828	0.686	0.711	0.633
	$X_r$	0.025	0.016	0.016	0.014
	$R_{sqr}$	0.998	0.998	0.993	0.980

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211

212 Figure 3. (a) CaO conversion versus cycle number for carbonation/calcination tests carried out under CaL  
 213 and Oxy-CaL conditions. (b) Conversion in the fast reaction controlled phase. (c) Conversion in the solid-  
 214 state diffusion phase. Carbonation is carried out at 650°C under 15% vol CO<sub>2</sub> (CaL) and 30% vol, 45% CO<sub>2</sub>  
 215 and 60% vol CO<sub>2</sub> for the Oxy-CaL tests as indicated. Calcination in all the tests is performed at 950°C  
 216 (70% CO<sub>2</sub>/30% air vol/vol). Solids lines are the best fits of equation 1 to data.

217



218 The effect of varying the carbonation temperature around 650°C on the multicycle CaO  
219 conversion performance for the different CaL and Oxy-CaL conditions was also investigated  
220 in our work. Data on the multicycle CaO conversion ( $X_N$ ,  $X_{FRP}$  and  $X_{SDP}$ ) for the tests carried  
221 out under CaL and Oxy-CaL-45 conditions at 625, 650 and 680°C are shown in Appendix A.  
222 The results show that the overall conversion increases with the carbonation temperature while  
223 conversion in the fast reaction controlled phase is essentially independent of the carbonation  
224 temperature in the range of temperatures tested. The main effect of varying the carbonation  
225 temperature is therefore observed on the conversion in the solid-state diffusion controlled  
226 phase, which is significantly enhanced with the carbonation temperature (see Figure 15c and  
227 Figure 15f in Appendix A). A similar behavior was observed for the samples tested under  
228 Oxy-CaL-30 and Oxy-CaL-60 conditions.

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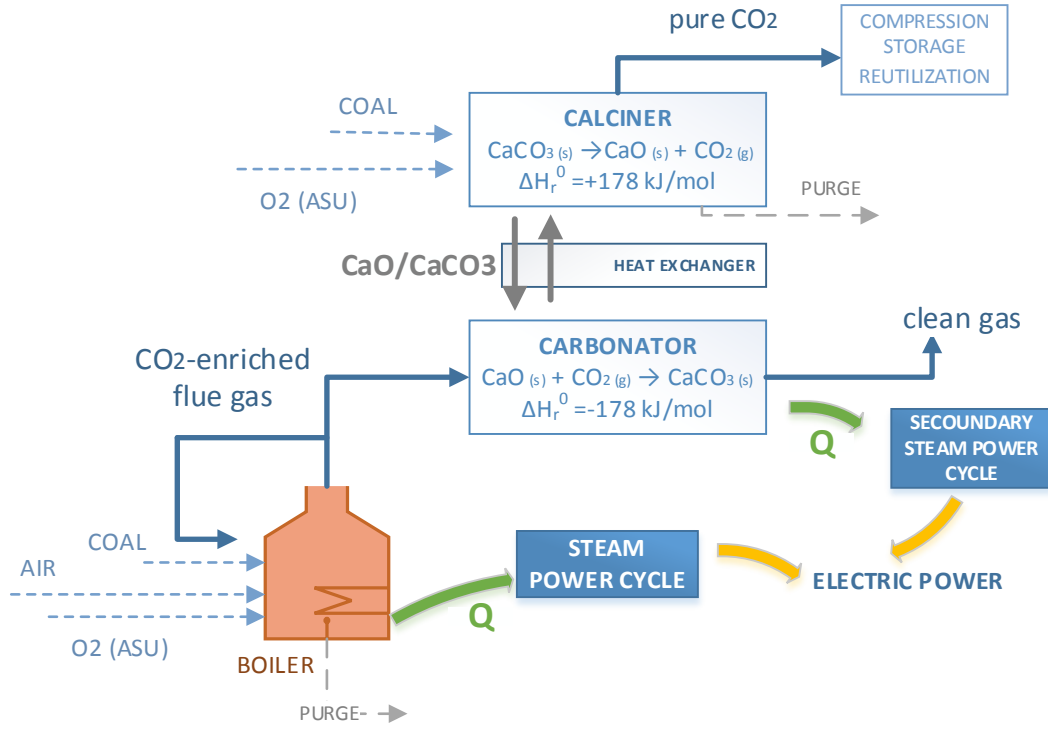
### 231 3. The Oxy-CaL process

#### 232 a. Description

233 The Oxy-CaL process newly proposed in the present manuscript is a CO<sub>2</sub> capture hybrid  
234 system based on the combined use of partial oxy-combustion and the CaL capture process.  
235 The basic idea behind Oxy-CaL is to exploit the enhancement of CO<sub>2</sub> capture capacity in the  
236 CaL process as the CO<sub>2</sub> concentration in the carbonation environment is increased (as seen  
237 above from the TGA tests) whereas the energy penalty for partial oxy-combustion to increase  
238 the CO<sub>2</sub> concentration in the flue gas is notably reduced as compared to total oxy-combustion.  
239 Figure 5 shows a schematic representation of this integration as applied to CO<sub>2</sub> capture in a  
240 coal fired power plant (CFPP).

241  
242 As can be seen in Figure 5, the Oxy-CaL process is initiated by partial oxy-combustion of  
243 coal using to this end a mixture of air, nearly pure oxygen (purity  $\geq 95\%$ ) and CO<sub>2</sub>-enriched  
244 recycled flue gas at combustor temperatures between 850°C and 950°C. As a result, the flue  
245 gas stream exiting the boiler reaches a CO<sub>2</sub> vol concentration in the range 30–60% (depending  
246 on the air/O<sub>2</sub>/CO<sub>2</sub> mixture composition) instead of the typical 15% vol concentration obtained  
247 from combustion with just air. The heat released by combustion is used for electric power  
248 production by means of a steam power cycle. Once partial oxy-combustion is carried out, the  
249 CO<sub>2</sub>-enriched flue gas is sent to the CaL process. The CO<sub>2</sub> present in the flue gas reacts in the  
250 carbonator with a fluidized bed of CaO particles at temperatures around 650°C. The  
251 carbonated particles are then circulated to the calciner reactor in which fast decomposition of  
252 CaCO<sub>3</sub> occurs to regenerate the CaO solids and produce a rich CO<sub>2</sub> stream ready to be  
253 compressed and transported for storage or other uses.

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255  
256 Figure 4: Oxy-CaL process schematics  
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## 260 b. CO<sub>2</sub> capture efficiency (carbonator model)

261 The CO<sub>2</sub> capture efficiency in the CaL process will be assessed by means of the carbonator  
262 model described in detail elsewhere [38]. Accordingly, the CO<sub>2</sub> capture efficiency can be  
263 expressed as a function of the total solids inventory in the carbonator ( $W_s$ ), or the moles  
264 number of Ca-based solids ( $N_{Ca}$ ), the solids residence time in the carbonator ( $\tau$ ) and the flow  
265 rate of fresh limestone makeup fed into the system ( $F_0$ ). In this model an average CaO  
266 conversion ( $X_{ave}$ ) is defined by the sum of the average conversion in the fast reaction-  
267 controlled phase ( $X_{ave,FRP}$ ) and conversion in the solid-state diffusion controlled phase  
268 ( $X_{ave,SDP}$ ), where both  $X_{ave,FRP}$  and  $X_{ave,SDP}$  are calculated by assuming that the gas passes  
269 in plug flow across a bed of perfectly mixed solids in the carbonator. The interested reader  
270 may see ref. [38] for further details on the carbonator model.

271 According to this carbonator model, the average reaction rates in the FRP and SDP phases are  
272 expressed as:

$$r_{ave,FRP} = \frac{X_{ave,FRP}}{t_{FRP}} \quad \text{for } t \leq t_{FRP} \quad (2)$$

$$r_{ave,SDP} = \frac{X_{ave,SDP}}{t_0 - t_{FRP}} \quad \text{for } t_{FRP} < t \leq t_{max} \quad (3)$$

273 where  $r_{ave,i}$  is the average reaction rate in the i-phase (either FRP or SDP),  $t_{FRP}$  is the time  
 274 lag of the FRP phase,  $t_{max}$  is total carbonation time,  $X_{ave,i}$  is the average capture capacity in  
 275 the i-phase, and  $t_0$  is the overall carbonation time lag in the carbonation TGA test (section 2).

276 The average rate of CaO conversion in the kinetically controlled fast phase ( $r_{aveFRP}$ ) at  
 277 atmospheric pressure can be approximated by a first-order kinetic law [30]:

$$r_{aveFRP} = k_s S_{ave} (1 - X)^{\frac{2}{3}} ([CO_2] - [CO_2]_{eq}) \quad (4)$$

278 where  $[CO_2]$  and  $[CO_2]_{eq}$  are the actual and equilibrium  $CO_2$  concentrations, respectively,  $k_s$   
 279 is the kinetic constant and  $S_{ave}$  is the average CaO specific surface area available for reaction  
 280 after N cycles.

281 On the other hand, the average rate of CaO conversion in the diffusion controlled phase can  
 282 be expressed by means of an effective diffusion constant ( $D^*_{eff}$ ) [38].

$$r_{aveSDP} \approx D^*_{eff} ([CO_2] - [CO_2]_{eq}) \quad (5)$$

283 Both Eq. 4 and Eq. 5 can be well fitted to the experimental TGA data shown above, which  
 284 allows us obtaining the values of  $k_s$  and  $D^*_{eff}$  to be used in the kinetic model for each one of  
 285 the CaL and oxy-CaL systems considered. Values of best fitting parameters are shown in  
 286 Table 2.

287 Table 2: Kinetic model parameters obtained from the best fits of Eqs. 4 and 5 to experimental TGA data.

288

	$k_s (x 10^{10}) \left[ \frac{m^4}{mol \cdot s} \right]$	$D^*_{eff} (x 10^5) \left[ \frac{m^3}{mol \cdot s} \right]$
CaL-625	8.87	2.70
CaL-650	10.00	4.63
CaL-680	12.27	8.09
Oxy-CaL30-625	4.429	1.07
Oxy-CaL30-650	6.08	1.61
Oxy-CaL30-680	4.51	2.75
Oxy-CaL45-625	3.21	0.61
Oxy-CaL45-650	3.60	1.02
Oxy-CaL45-680	2.75	1.68

Oxy-CaL60-625	2.00	0.61
Oxy-CaL60-650	2.38	0.71
Oxy-CaL60-680	1.98	1.10

289

290 Once the average CaO conversion is calculated, the capture efficiency in the carbonator  
 291 ( $E_{CO_2}$ ) can be obtained as:

$$E_{CO_2} = \frac{F_R}{F_{CO_2}} X_{ave} \quad (6)$$

292 where  $F_R$  is the solids recirculation flow rate between the calciner and carbonator reactors,  
 293 which is given by  $F_R = W_s/N_{Ca}$ .

294

295 Figure 6 shows the evolution of the CO<sub>2</sub> capture efficiency ( $E_{CO_2}$ ) as the solids recirculation  
 296 flow rate between reactors is decreased or, equivalently, the solids residence time in the  
 297 carbonator ( $\tau = N_{Ca}/F_R$ ) is increased. As may be seen, a significantly higher capture  
 298 efficiency is achieved by increasing the CO<sub>2</sub> concentration in the carbonator in the Oxy-CaL  
 299 systems. Note the differences on the maximum capture efficiency ( $E_{max} = (y_{CO_2,in} - y_{eq}) /$   
 300  $y_{CO_2,in}$ ) for the different systems (obtained for very short residence times) as a consequence  
 301 of the variation of the CO<sub>2</sub> vol % in the carbonator. Even though the use of short residence  
 302 times leads to relatively high capture efficiencies it must be kept in mind that short residence  
 303 times would rise the cost for solids transportation. The sensible heat needed to increase the  
 304 temperature of the solids stream entering the calciner would be also raised as the solids  
 305 recirculation rate is increased to achieve short residence times.

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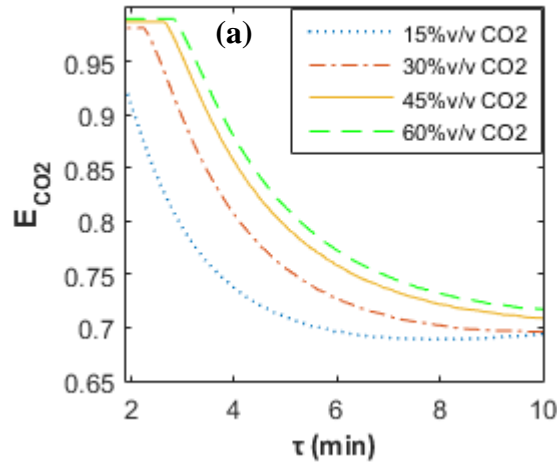
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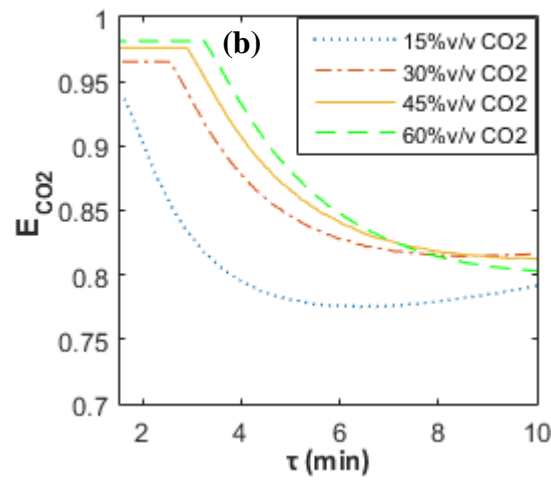
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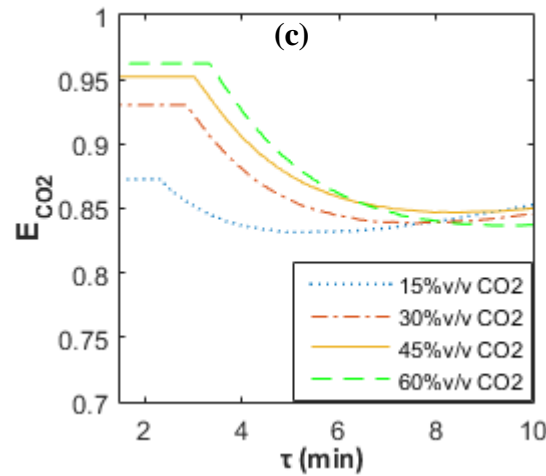
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320



321 Figure 5: CO<sub>2</sub> capture efficiency as a function of the residence time in the carbonator, which is varied by  
322 changing the  $F_R/F_{CO_2}$  ratio. Calculations are made for fixed values of the solids inventory  $W_S =$   
323 400 ton and  $F_0/F_{CO_2} = 0.05$  and different carbonation temperatures. a)  $T_{carb} = 625^\circ C$ ; b)  $T_{carb} =$   
324  $650^\circ C$  and c)  $T_{carb} = 680^\circ C$ .

325

326 As shown in Figure 6, the capture efficiency is decreased as the solids residence time in the  
327 carbonator is prolonged albeit at a minor rate for the CaL process in comparison with the  
328 Oxy-CaL systems. This is due to the relatively higher conversion in the solid-state diffusion

329 controlled phase for the CaL process (as seen above from the TGA tests). This result  
 330 becomes more marked as the carbonator temperature is increased (compare Figure 6a and  
 331 Figure 6c) as a consequence of the enhancement of solid-state diffusivity with temperature.  
 332 The results obtained up to this point show already some hints concerning the CFPP-CO<sub>2</sub>  
 333 capture integration. An increase in the solids residence time in the carbonator would arguably  
 334 allow for a reduction of the energy penalty although the capture efficiency would be  
 335 hampered depending on the capture system. Thus, a comparative assessment of the diverse  
 336 capture systems and their integration into CFPP must necessarily include an evaluation of the  
 337 energy penalty. This will be the subject of the next section.

#### 338 4. CFPP-CO<sub>2</sub> capture integration models

339 This section is devoted to a comparative assessment of several CO<sub>2</sub> capture technologies,  
 340 namely oxy-combustion, CaL and Oxy-CaL, and their integration into a Coal Fired Power  
 341 Plant (CFPP). Regarding energy penalty, the parameter usually employed in the literature is  
 342 the specific energy consumption for CO<sub>2</sub> avoided (SPECCA) [7], which quantifies the  
 343 additional fuel consumption (in MJ) needed to avoid the emission of 1 kg of CO<sub>2</sub> into the  
 344 atmosphere (Eq. 6)

$$345 \text{SPECCA [MJ/kg}_{CO_2}] = 3600 \frac{\frac{1}{\eta_{plant}} - \frac{1}{\eta_{ref}}}{E_{ref} - E} \quad (6)$$

346 where  $\eta_{ref}, \eta_{plant}$  are the CFPP efficiency, and  $E_{ref}, E$  are the emissions ratio (in kg  
 347 CO<sub>2</sub>/kWh<sub>e</sub>) without and with the capture system integrated, respectively.

348  
 349 A 490 MW<sub>e</sub> CFPP has been chosen in our work as reference plant, which is modelled using  
 350 the commercial software ASPEN PLUS<sup>TM</sup>. In order to simplify the model a number of  
 351 assumptions were made: (i) the system operates at steady conditions; (ii) minimum  
 352 temperature difference is 20°C for all heat exchangers; (iii) ideal behavior of cyclones; (iv)  
 353 solid–solid heat exchange is simulated as a transfer of heat between solids; (v) 89% isentropic  
 354 efficiencies are assumed as constant for all turbomachinery. In this CFPP, air-combustion of  
 355 42.2 kg/s of coal Pittsburgh No. 8 (see [39] for coal type details) takes place in the steam  
 356 boiler (operating at an average temperature of 900°C) to generate 1297 MW<sub>th</sub>, which releases  
 357 to the atmosphere 513.4 kg/s of flue gas with a CO<sub>2</sub> vol concentration of 15% at atmospheric  
 358 pressure. Electric power is produced by means of a reheat supercritical steam cycle ( $P_{vv} = 290$   
 359 bar,  $T_{vv} = 600/620^\circ\text{C}$ ), wherein the steam regenerative process is carried out from four feed-  
 360 water heaters, one of which is a total mixer exchanger type (degasifier). Selected conditions  
 361 lead to a 44% thermal to electric net efficiency. Taking into account parasitic electricity, the  
 362 overall net efficiency drops to 37.81%. This value will be used as a reference to calculate the  
 363 penalty arising from the integration in this plant of the diverse CO<sub>2</sub> capture systems. Main  
 364 inputs and results from the reference plant model are summarized in Table 3. An extended  
 365 table showing additional parameters is shown in Appendix B.

366  
 367

368 Table 3: Main inputs and results for the base case of diverse CO<sub>2</sub> capture systems

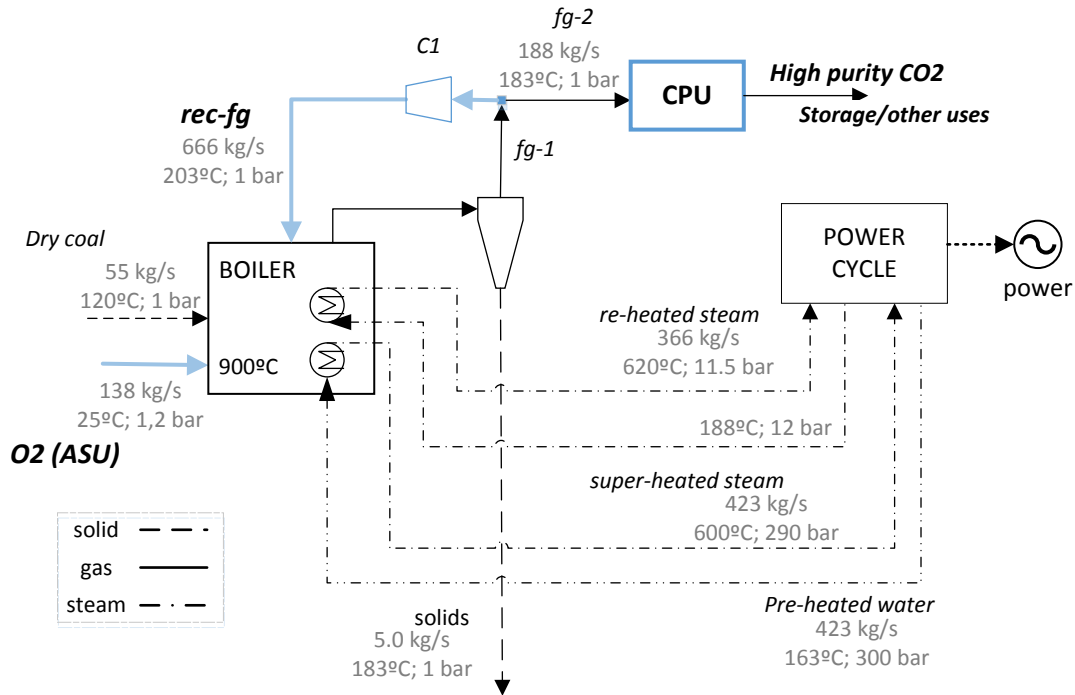
	parameter	Reference CFFP (air combustion)	oxy- combustion	CaL	Oxy- CaL 30	Oxy- CaL 45	Oxy- CaL 60
CFPP	$\dot{m}_{coal}$ (kg/s)	42.20	55.05	42.20	46.10	47.50	48.15
	$\dot{m}_{air}$ (kg/s)	475	-	475	208.90	100.20	43.54
	$\dot{m}_{O_2}$ (kg/s)	-	136.91	-	68.85	96.35	110.503
	$v_{CO_2}$	0.15	0.89	0.15	0.30	0.45	0.60
	$\eta_{CFPP}$	0.377	0.287	0.377	0.351	0.344	0.337
	net work (MW)	490.47	488.80	490.47	498.30	502.30	499.75
	Penalty	-	<b>9.02%</b>	-	<b>2.64%</b>	<b>3.47%</b>	<b>4.03%</b>
	SPECCA (MJ/kgCO <sub>2</sub> )	-	<b>4.06</b>	-	<b>0.94</b>	<b>1.25</b>	<b>1.50</b>
CaL	$T_{calc}$ (°C)	-	-	950	950	950	950
	$T_{carb}$ (°C)	-	-	650	650	650	650
	$E_{CO_2}$	-	-	0.827	0.950	0.976	0.981
	$\dot{m}_{coal}$ (kg/s)	-	-	18.48	22.34	23.40	23.84
	$\dot{m}_{O_2}$ (kg/s)	-	-	48.00	58.41	61.00	62.49
	$\dot{W}_{sec}$ (MW)	-	-	75.80	113.90	126.46	130.88
	$\eta_{int}$	-	-	0.303	0.291	0.288	0.285
	Penalty	-	-	<b>7.43%</b>	<b>6.04%</b>	<b>5.48%</b>	<b>5.27%</b>
	SPECCA (MJ/kgCO <sub>2</sub> )	-	-	<b>3.28</b>	<b>2.63</b>	<b>2.37</b>	<b>2.29</b>
Total	$\dot{m}_{coal,total}$	-	-	60.68	68.44	70.90	71.99
	Penalty <sub>total</sub>	-	<b>9.02%</b>	<b>7.43%</b>	<b>8.68%</b>	<b>8.95%</b>	<b>9.30%</b>
	SPECCA <sub>total</sub> (MJ/kgCO <sub>2</sub> )	-	<b>4.06</b>	<b>3.28</b>	<b>3.56</b>	<b>3.62</b>	<b>3.79</b>

369

370

### a. Total Oxy-combustion

371 Firstly, an oxy-fuel combustion process will be analyzed in order to assess the CO<sub>2</sub> capture  
372 efficiency and the energy consumption in comparison with the above-mentioned reference  
373 plant. The CFFP oxy-combustion model has been developed using ASPEN PLUS<sup>TM</sup> and the  
374 same coal (Pittsburgh No. 8). Figure 7 shows a flow diagram of the oxy-combustion model  
375 that highlights the differences with respect to the reference air-combustion plant. For  
376 simulating coal oxy-combustion a model based on chemical and phase equilibrium through  
377 Gibbs' free energy minimization method is used.



378

379 Figure 6: Schematics of the total oxy-combustion CFPP. Differences with air-combustion CFPP are  
 380 highlighted in blue color (for interpretation of the references to color in this figure, the reader is referred to  
 381 the web version of this article.)

382

383 In this CFPP oxy-combustion case, combustion of 55.3 kg/s of coal Pittsburgh No. 8 with  
 384 137.6 kg/s of a high purity (95%) O<sub>2</sub> stream from ASU releases 187.8 kg/s of flue gas with a  
 385 CO<sub>2</sub> vol concentration (dry-basis) of 89% at atmospheric pressure, which, after purification  
 386 and compression, is ready to be stored [40] or used in other industrial processes [41,42]. A  
 387 flue gas recirculation with a recycle ratio of 0.78 is carried out in order to control the flame  
 388 temperature in the boiler [43]. A compressor (C1 in the figure) is used to overcome the  
 389 pressure drop in the reactor. ASU energy consumption has been estimated as 200 kWh per kg  
 390 of pure O<sub>2</sub> [44,45]. CO<sub>2</sub> purification unit (CPU) specific energy consumption has been fixed  
 391 to 143 kWh/tCO<sub>2</sub> [22] in order to simplify the model. Power is produced using the same  
 392 reheat supercritical steam cycle ( $P_{vv} = 290$  bar,  $T_{vv} = 600/620^\circ\text{C}$ ) as in the reference air-  
 393 combustion CFPP case. Main inputs and results from the model are summarized in Table 3.  
 394 From the simulations, the resulting specific CO<sub>2</sub> emissions are 86.2 g/kWh corresponding to  
 395 90% CO<sub>2</sub> capture efficiency. Remaining CO<sub>2</sub> emissions result from the nearly 8% of CO<sub>2</sub> lost  
 396 in the purification process, which is consistent with results from previous works [40]. A net  
 397 thermal to electric efficiency of 28.8% is achieved yielding an energy penalty of 9.1% points,  
 398 which is within the range of previously reported results (8-12%) [18,46].  
 399

400

## b. Calcium-Looping

401 In this section, the CaL integration model for post-combustion CO<sub>2</sub> capture used in the  
 402 present work is summarized. The reader interested in further details may see ref. [47] where a  
 403 similar model is thoroughly described. The CaL process is accomplished by using two



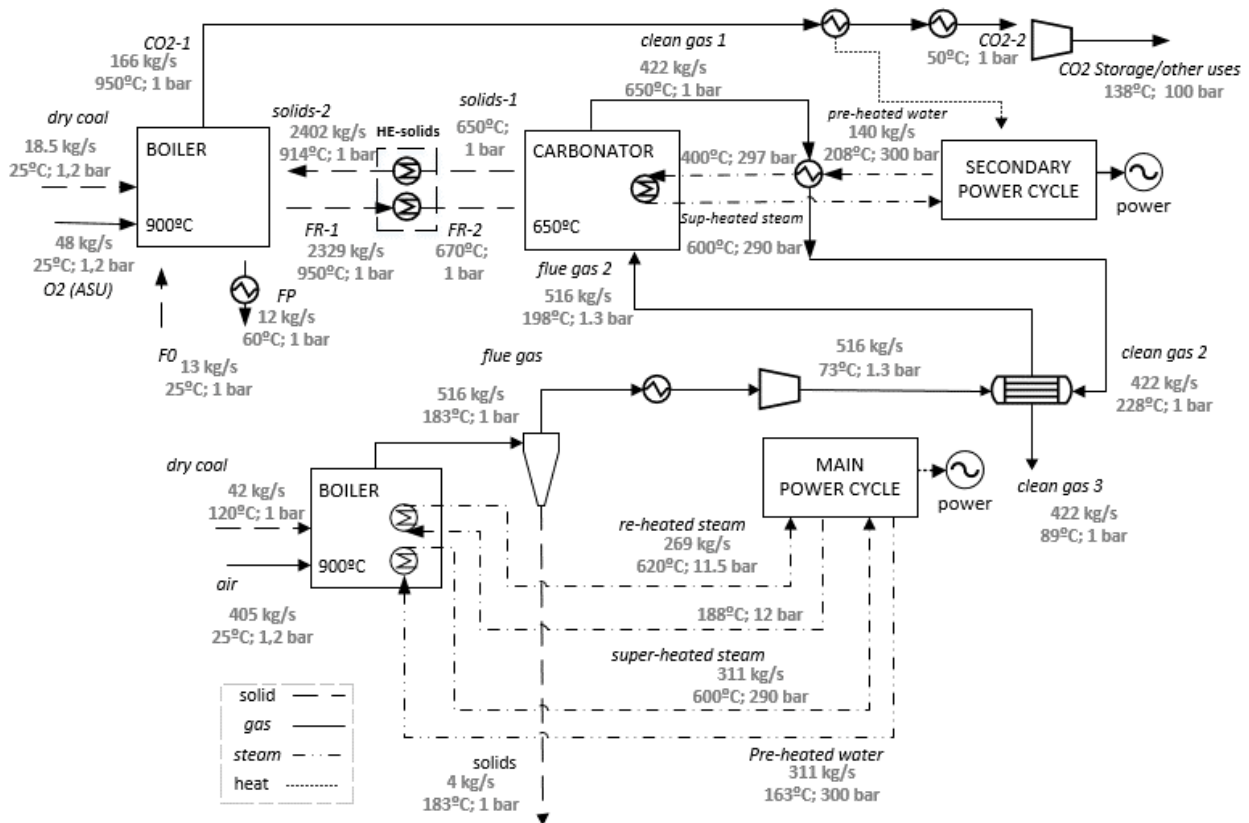
404 interconnected circulating- fluidized-bed (CFB) reactors, both operated under atmospheric  
405 pressure at gas velocities of approximately 5 m/s [48,49]. CaO particles react in the  
406 carbonator according to the carbonation reaction (Eq.7) at temperatures between 625-680°C  
407 with the CO<sub>2</sub> present in the flue gas stream coming from the CFPP plant. The partially  
408 carbonated particles are then circulated into the calciner reactor in which fast decomposition  
409 of CaCO<sub>3</sub> occurs at 950°C [15,25,50,51] to regenerate the sorbent and produce a rich CO<sub>2</sub>  
410 stream to be compressed and transported for storage or other uses. The regenerated CaO  
411 particles are recovered at the calciner exit by a cyclone and sent back to the carbonator for a  
412 new cycle. CO<sub>2</sub> capture in this reactor is modeled according to an equilibrium reactor  
413 following the model described in section 3.b.



414 Figure 8 shows a schematic representation of the CaL process. The heat produced in the  
415 exothermic carbonation reaction as well as the sensible heat recovered from the streams  
416 exiting the calciner is used in a secondary steam cycle for electricity generation. Given the  
417 large flow rate of solids recirculated between reactors in the CaL cycle, a heat exchanger  
418 (simplified as a heat transfer between solids with a temperature approach of 20°C) is  
419 incorporated for transferring sensible heat between the solids leaving the calciner (FR in  
420 Figure 8, with a temperature of about 950°C) and the solids entering into it to be heated up to  
421 the calcination temperature.

422  
423 In order to attain full calcination in short residence times of the limestone makeup fed into the  
424 calciner, the temperature in the calciner reactor must be 930° o even higher [14,52]. This  
425 makes necessary to supply a large amount of heat to the calciner, which is accomplished by  
426 in-situ oxy-fuel combustion in order not to dilute CO<sub>2</sub> in this reactor. CO<sub>2</sub> compression is  
427 modelled as a multi-stage compression to 100 bar refrigerated with water at ambient  
428 temperature. Pressure drop of the flue gas across the carbonator is calculated from the Kunii–  
429 Levenspiel (K–L) fluid dynamics model [53,54]. A compressor is used to counteract this  
430 pressure drop. Energy consumption derived from solids transportation has been set at 20 MJ  
431 per ton of solids [55]. Main inputs and results from the model are summarized in Table 3. The  
432 base case for CFPP-CaL integration leads to a CO<sub>2</sub> capture efficiency of 82.7% for an overall  
433 plant efficiency of 30.3%, which implies an energy penalty of 7.4% points in the range of  
434 previous values reported in literature [6,56,57]. As regards specific energy consumption (Eq.  
435 6), a SPECCA value of 3.3 MJ/kg CO<sub>2</sub> is achieved.

436  
437



438  
439 Figure 7: CFPP-CaL integration scheme

440  
441

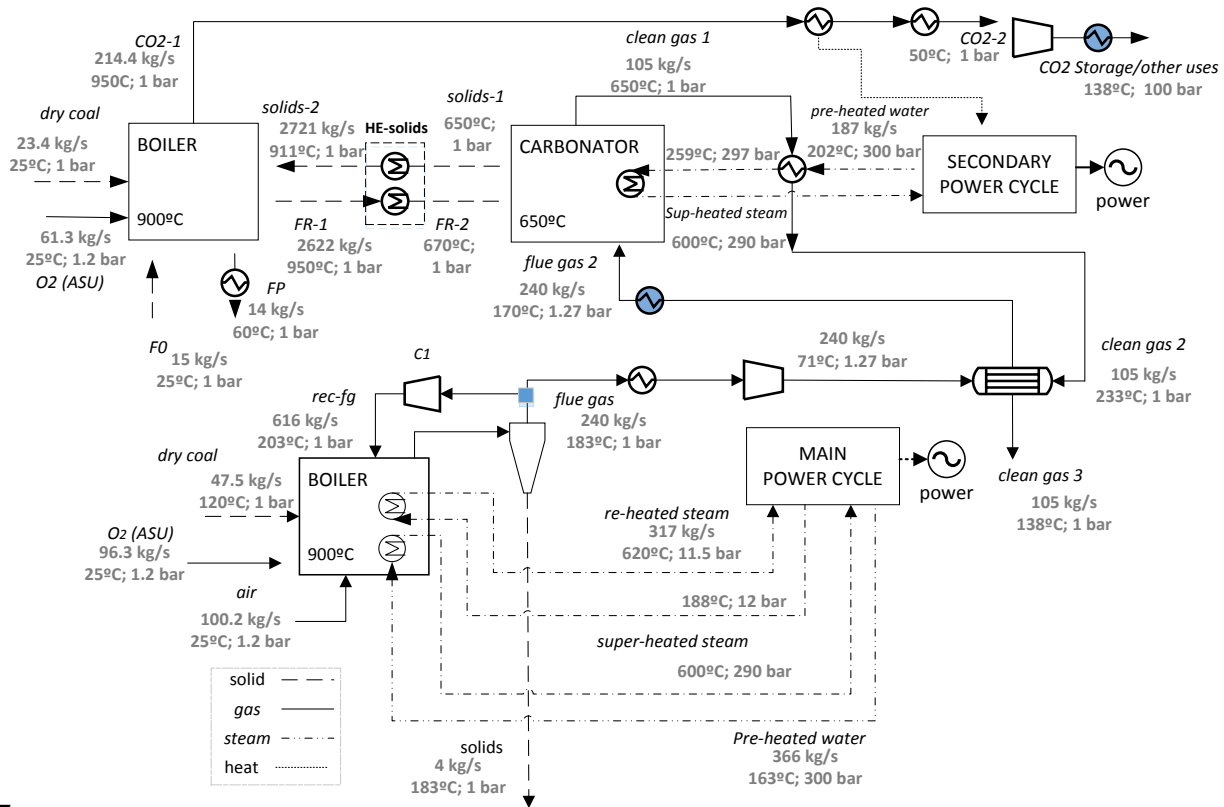
### 442 c. Oxy-CaL

443 This section describes the novel Oxy-CaL hybrid system proposed in the present work. Figure  
444 9 shows a schematic representation of the process, which has been simulated for several  
445 values of the CO<sub>2</sub> vol % in the flue gas effluent from partial oxy-combustion. As can be seen  
446 in Figure 9, the process is initiated by an oxy-fuel combustion similar to that described in  
447 section 4.a. Partial oxy-combustion is carried out to obtain a CO<sub>2</sub> vol% in the range of 30-  
448 60% in the flue gas at the boiler exit. To this end, a mixture of air and O<sub>2</sub> is used in the boiler  
449 for combustion. The air/O<sub>2</sub> ratio is calculated to achieve a given CO<sub>2</sub> vol% in the flue gas  
450 (45% in the case illustrated in Figure 9). As in the case of total oxy-combustion, recirculation  
451 of the flue gas serves to control the flame temperature in the boiler, whose value is kept the  
452 same for all the simulations.

453 Since the amount of pure O<sub>2</sub> for partial oxy-combustion is substantially decreased (99.3 kg/s  
454 to achieve a 45% vol CO<sub>2</sub> concentration instead of 138 kg/s for total oxy-combustion), power  
455 consumption in the ASU is notably reduced. Furthermore, the CPU unit for CO<sub>2</sub> purification  
456 is not needed since this step is carried out after the CaL process. Altogether, the energy  
457 penalty for partial oxy-combustion is significantly reduced. Thus, energy penalty is 3.47% in  
458 the Oxy-CaL 45 system as compared to 9.02% for total oxy-combustion.

459

460 After partial oxy-combustion, the CO<sub>2</sub> rich flue gas is sent to the carbonator reactor to follow  
 461 up with the CaL process, being before slightly compressed (to overcome the pressure drop in  
 462 the carbonator) and preheated with the hot gas streams exiting the CaL cycle. In contrast with  
 463 the CaL scheme (Figure 8), flue gas preheating is carried out in the Oxy-CaL system by using  
 464 firstly the compressed CO<sub>2</sub> stream since the thermal capacity of the flue gas at the carbonator  
 465 exit is lower than in the case of the CaL. The CO<sub>2</sub> entering into the carbonator reacts with the  
 466 CaO solids coming from the calciner as in the conventional CaL model (section 4.b)  
 467 according to the carbonation reaction (Eq. 7). As discussed in section 3, the capture efficiency  
 468 is significantly enhanced when the CO<sub>2</sub> concentration in the flue gas is increased (Figure 6).  
 469 Thus, the Oxy-CaL-45 system has a CO<sub>2</sub> capture efficiency of 97.6% in the base case (see  
 470 Figure 6a) as compared to 82.7% in the base case of the CaL system. Such increase in the  
 471 capture capacity implies also the handling of a larger amount of CaCO<sub>3</sub>, which leads to higher  
 472 heat needs in the calciner for CaO regeneration, and therefore to a higher consumption of coal  
 473 and O<sub>2</sub> as can be seen by comparison of Figure 8-9. Nonetheless, the amount of CO<sub>2</sub> captured  
 474 in this Oxy-CaL-45 system is 29% over that captured by means of CaL (section 4.b) and 14%  
 475 above the CO<sub>2</sub> captured by total oxy-combustion (section 4.a).  
 476



477  
 478 Figure 8: General Oxy-CaL-45 (45% vol CO<sub>2</sub> concentration in the flue gas by partial oxy-combustion)  
 479 integration scheme

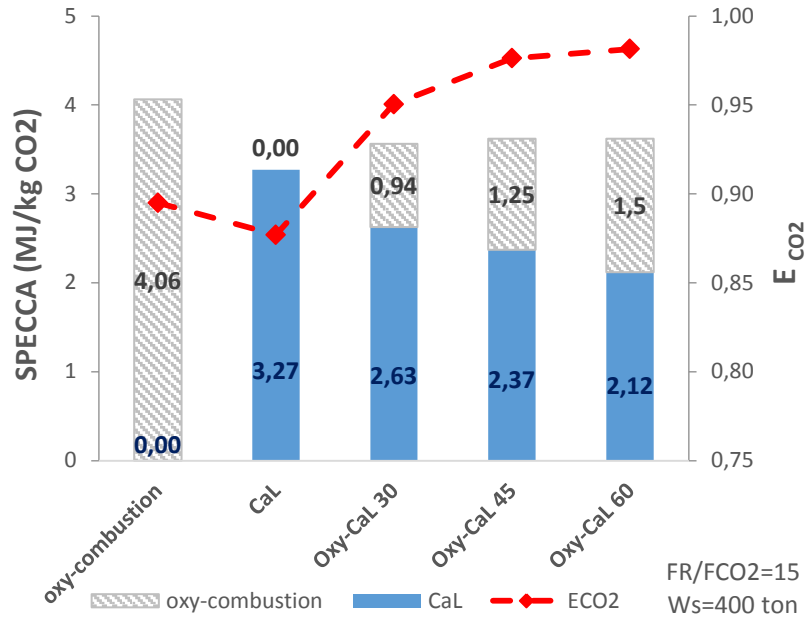
480  
 481 Despite the need of additional coal and O<sub>2</sub> for oxy-combustion in the calciner, the increase in  
 482 CO<sub>2</sub> capture efficiency obtained by increasing the CO<sub>2</sub> concentration in the flue gas leads to a  
 483 reduction of energy consumption in the CaL cycle. Thus, a SPECCA value of 2.37 MJ/kg  
 484 CO<sub>2</sub> is obtained for the Oxy-CaL-45 system, which is 28% below the SPECCA obtained for

485 the conventional CaL system. Nevertheless, the SPECCA for the complete oxy-CaL process  
486 is 3.62 MJ/kg CO<sub>2</sub>, which is 10% higher than in the CaL base case. On the other hand, the  
487 Oxy-CaL system allows for a reduction by 11% of energy consumption in comparison with  
488 the total oxy-combustion case. The next sections are devoted to a deeper comparative analysis  
489 on both capture efficiency and energy penalty resulting from the diverse capture systems with  
490 the goal of finding the most feasible choice to be implemented in practice.  
491

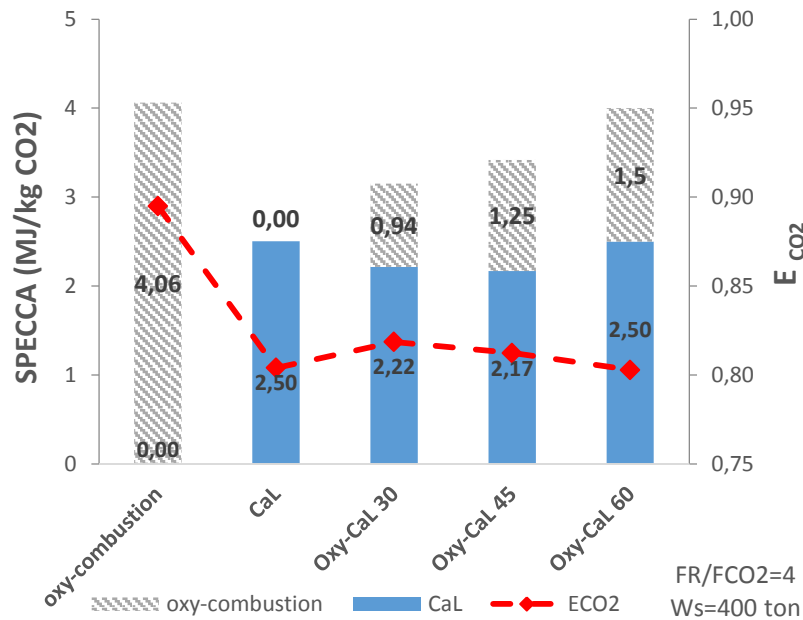
## 492 **5. CO<sub>2</sub> capture efficiency and energy consumption**

493  
494 As was shown in sections 2.b (TGA results) and 3.b (carbonator model), the CO<sub>2</sub> capture  
495 efficiency in the CaL process is remarkably enhanced by increasing the CO<sub>2</sub> concentration in  
496 the flue gas stream entering into the carbonator. This is reflected also in the SPECCA, which  
497 is decreased due to the higher efficiency of the CO<sub>2</sub> capture process. On the other hand,  
498 partial oxy-combustion carried out to increase the CO<sub>2</sub> concentration at the inlet of the  
499 carbonator in the CaL process contributes also to an additional energy penalty. The use of the  
500 CaL process, total oxy-combustion or a hybrid Oxy-CaL process for CO<sub>2</sub> capture in CFPP is  
501 carefully assessed below on the basis of the benefits and drawbacks of each one of these  
502 systems.  
503

504 Figure 10 shows the CO<sub>2</sub> capture efficiency and SPECCA values for total oxy-combustion,  
505 CaL, and Oxy-CaL systems obtained for the base cases analyzed in section 4. As seen in  
506 Figure 10a, the part of SPECCA that corresponds to the CaL process in the Oxy-CaL systems  
507 is decreased as the carbonation CO<sub>2</sub> vol% is increased if the oxy-CaL process is operated with  
508 a high solids recirculation flow rate (FR/F<sub>0</sub>=15 corresponding to  $\tau$  =2 min for the solids  
509 residence time and fixing W<sub>s</sub>=400 tons as solids inventory). Nevertheless, energy  
510 consumption in the partial oxy-combustion part of the process leads to global oxy-CaL  
511 SPECCA values somewhat higher than for the purely CaL process. On the other hand, the  
512 oxy-CaL 30 , 45 and 60 systems have a SPECCA smaller than total oxy-combustion. Note  
513 also that the CO<sub>2</sub> capture efficiency is notably increased for the oxy-CaL systems as  
514 compared to oxy-combustion and is especially increased over the CaL process. Figure 10b  
515 shows however that under prolonged solids residence times of  $\tau$  =10 min (corresponding to a  
516 reduced solids recirculation flow FR/F<sub>0</sub>=4 and fixing W<sub>s</sub>=400 tons as solids inventory) the  
517 capture efficiency of the oxy-CaL systems is hindered since carbonation in the solid-state  
518 diffusion controlled stage is not significant for carbonation under relatively high CO<sub>2</sub> vol% as  
519 was seen from the TGA tests (section 3.b).  
520



521



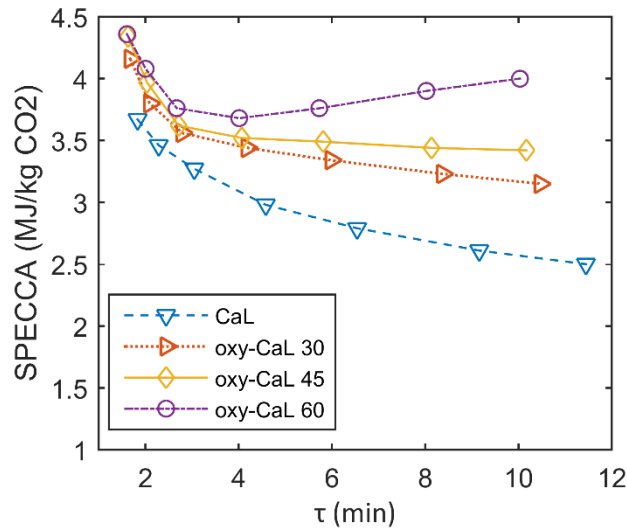
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523 Figure 9: SPECCA values and CO<sub>2</sub> capture efficiencies for the CO<sub>2</sub> capture systems analysed in this work  
 524 and using reference parameters shown in table 3. Figures a) and b) correspond to different values of the  
 525 solids residence time  $\tau$  in the carbonator for the CaL process. a) FR/FCO<sub>2</sub>=15 ( $\tau$ =2 min), b) FR/FCO<sub>2</sub>=4 ( $\tau$   
 526 =10 min).

527

528 As discussed above and according to recent works [13,47], energy consumption in the CaL  
 529 process is highly dependent on the carbonation rate in the solid-state diffusion controlled  
 530 phase (SDP), which determines the role of the solids residence time  $\tau$  in the carbonator.  
 531 Figure 11 shows how SPECCA evolves as  $\tau$  is increased for the CaL and hybrid oxy-CaL  
 532 processes. In the case of the CaL process, the rate of carbonation in the solid-state diffusion  
 533 controlled phase is not negligible as compared to the carbonation rate in the fast reaction  
 534 controlled phase, which leads to a considerable reduction of the energy consumption as  $\tau$  is  
 535 increased [13,47]. On the other hand, the rate of carbonation in the solid-state diffusion

536 controlled phase is decreased by increasing the CO<sub>2</sub> concentration in the flue gas as occurs in  
 537 the oxy-CaL systems (see Figure 1), which hinders a further reduction of SPECCA as  $\tau$  is  
 538 increased. For the Oxy-CaL-60 case, the SPECCA is even raised as  $\tau$  is prolonged beyond ~4  
 539 min. On the other hand, the SPECCA values for the CaL and oxy-CaL systems are similar for  
 540 short solids residence time (of about 2 minutes). Thus, it may be concluded that the optimum  
 541 operation of the oxy-CaL system is under short residence times.  
 542  
 543



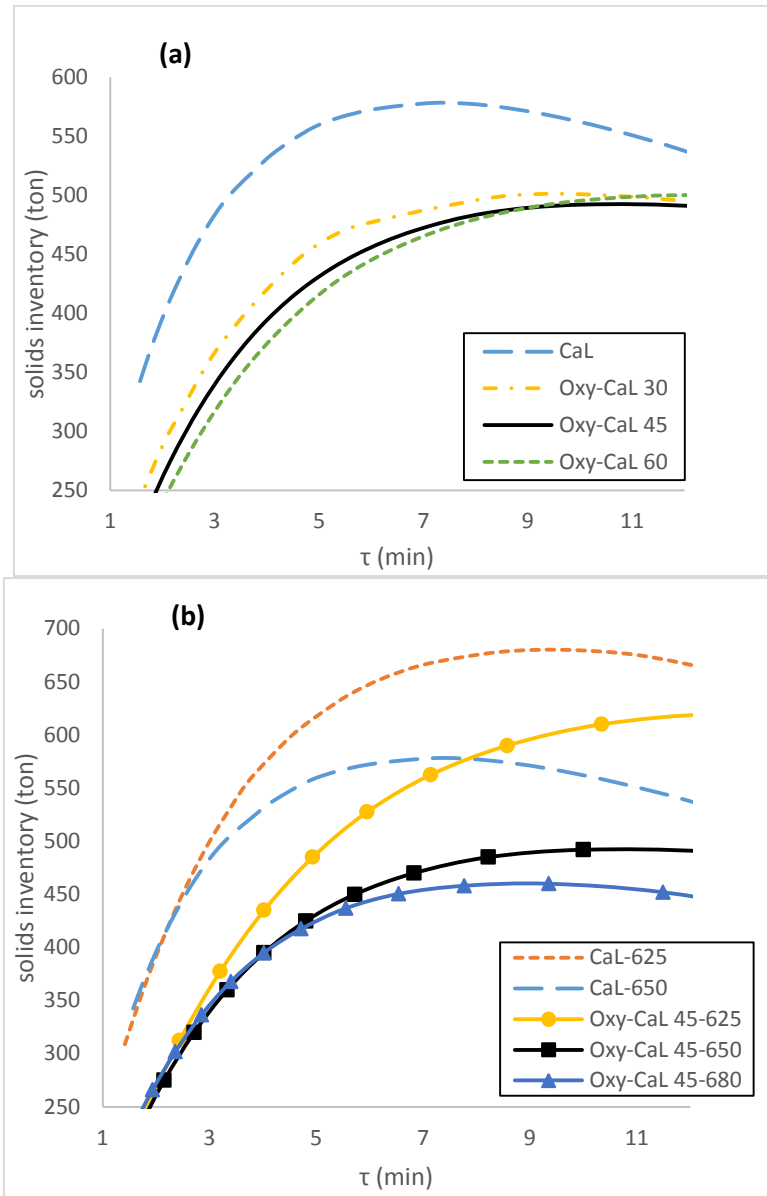
544  
 545 Figure 10: SPECCA values as a function of solids residence time in the carbonator for CaL and Oxy-CaL  
 546 systems operating at  $T_{carb}=650^{\circ}C$  (solids inventory fixed at  $W_s=400$  ton).

547  
 548  
 549

### 550 5.1 Role of solids inventory

551  
 552 A straightforward consequence of the improvement of CO<sub>2</sub> capture efficiency in the CaL  
 553 process as the CO<sub>2</sub> vol% in the carbonator is increased (Oxy-CaL) is the possibility of  
 554 reducing the solids inventory ( $W_s$ ) and the limestone makeup flow ( $F_0$ ), which could lead to a  
 555 potentially relevant capital and operational cost cutback. Figure 12a shows the variation of the  
 556 solids inventory ( $W_s$ ) in the CaL and oxy-CaL processes with the solids residence time for a  
 557 fixed value of the capture efficiency ( $E_{CO_2} = 90\%$ ). As can be seen, significantly lower solids  
 558 inventories are needed in the Oxy-CaL processes under carbonation residence times below  
 559 ~12 minutes that would conform to usual operation conditions in CFB reactors. This result  
 560 implies an important potential for reducing the CaL size in the hybrid oxy-CaL systems. It  
 561 must be reminded that system size is currently one of the main limitations for the CaL process  
 562 to reach a demonstration stage [4,16]. Moreover, a lower solids inventory, as would be  
 563 possible in the Oxy-CaL system, leads to a reduction of power consumption to overcome the  
 564 gas pressure drop across the reactor. The oxygen production and coal input needed for  
 565 heating-up the additional quantity of solids in calciner are also diminished.  
 566

567  
568



569

570

571 Figure 11: 90% capture iso-efficiency lines as a function of solids inventory and solids residence time in  
572 the carbonator for CaL and Oxy-CaL systems at a carbonator temperature  $T_{carb} = 650^\circ\text{C}$  (a) and for the CaL  
573 and Oxy-CaL-45 systems at carbonation temperatures of 625, 650, and 680°C (b).

574

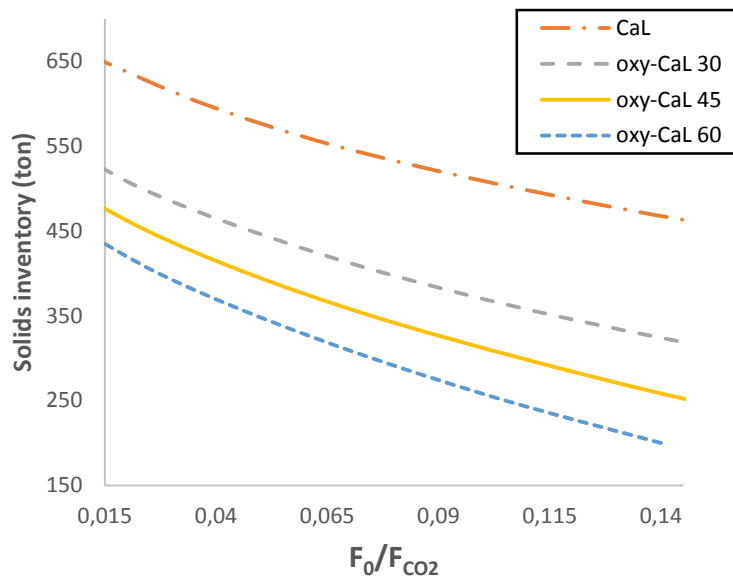
575 Figure 12b shows the evolution of  $W_s$  with the solids residence time in the carbonator for  
576 carbonation temperatures  $T = 625^\circ\text{C}$ ,  $650^\circ\text{C}$  and  $680^\circ\text{C}$  and a fixed capture efficiency ( $E_{CO_2} =$   
577  $90\%$ ). As may be observed, the relative decrease of the carbonation rate in the solid-state  
578 diffusion controlled stage when  $T$  is decreased to  $625^\circ\text{C}$  leads to a remarkable increase of  $W_s$   
579 for solids residence times beyond  $\sim 3$  min. Thus, the CaL process operated under long  
580 residence times, which would allow decreasing notably SPECCA [13,47], can be hampered  
581 by a small decrease of the carbonator temperature that might be expected locally under  
582 practical conditions [14,48] due to inefficient mass and/or energy transfer. Note that chemical  
583 equilibrium prevents in the case of the CaL process to achieve a 90% capture efficiency when

584 operating at 680°C (see the maximum capture value possible in Figure 6a). On the other hand,  
585 since the Oxy-CaL system would be ideally operated under short residence times, its  
586 performance would not be essentially affected by temperature changes.

587  
588 As seen in the results obtained from our TGA experiments (Figure 2), the rate of carbonation  
589 in the solid-state diffusion controlled phase depends critically on the carbonator temperature.  
590 Thus, carbonation in this phase is hindered if the temperature is decreased just from 650°C to  
591 625°C whereas it becomes enhanced by an increase of temperature up to 680°C. It is therefore  
592 interesting to assess the sensitivity of SPECCA and solids inventory needed in the CaL  
593 process to this small change of carbonation temperature that might occur in practice.

594  
595 Despite the potential for reducing the CaL size in the Oxy-CaL process, the additional energy  
596 consumption due to partial oxy-combustion, and therefore O&M costs, must be also  
597 considered to assess the Oxy-CaL feasibility. In this regard, a further benefit of the Oxy-CaL  
598 hybrid system is that the higher capture efficiency achieved allows to reduce the fresh  
599 limestone fresh makeup, which yields a decrease of energy penalty. Figure 13 shows the  
600 relationship between the solids inventory and make-up flow for the diverse systems analyzed  
601 at a fixed capture efficiency of 90%.

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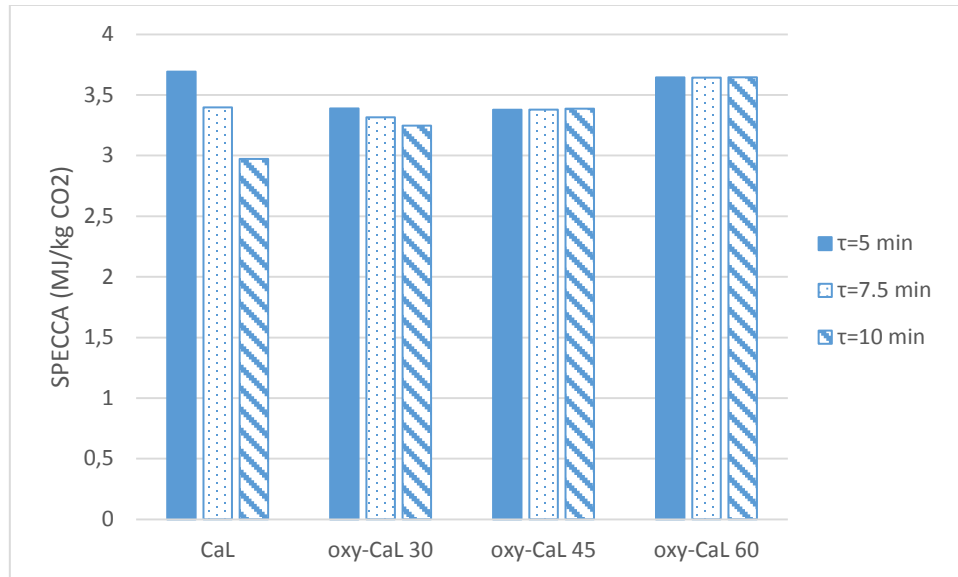
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604 Figure 12: Solids inventory as a function of makeup flow for CaL and Oxy-CaL systems at a carbonator  
605 temperature  $T_{carb}=650^\circ\text{C}$  and at capture efficiency of 90%. Solids recirculation flow rate is fixed to  
606  $F_R/F_{CO_2}=15$ .

607

608 As can be seen in Figure 13, by using the Oxy-CaL hybrid system it is possible to reduce  
609 considerably the fresh limestone makeup in comparison with the CaL process. For example, if  
610 we consider the Oxy-CaL 45 system with a solids inventory of 450 tons, the ratio of limestone  
611 makeup to  $\text{CO}_2$  flow rates ( $F_0/F_{CO_2}$ ) is just around 0.015, which is quite below the amount  
612 needed for the CaL process under the same operation conditions and capture efficiency  
613 ( $F_0/F_{CO_2}=0.12$ ). Figure 14 shows the SPECCA values obtained by fixing the solids inventory  
614 at 450 tons and varying the makeup flow to attain with each system a capture efficiency of  
615 90%.





617

618 Figure 13: SPECCA values for the CaL and Oxy-CaL systems obtained by considering a fixed solids  
 619 inventory of  $W_s=450$  tons and capture efficiency to 90%.

620

621 As shown in Figure 14, using a given solids inventory of 450 tons in the carbonator, SPECCA  
 622 values for Oxy-CaL systems are lower than for the CaL process when operating under solids  
 623 residence times below 7.5 min due to the lower makeup flow needed to reach a 90% capture  
 624 efficiency. Thus, the use of the Oxy-CaL systems would allow also for a reduction of O&M  
 625 costs associated to the capture process using typical reactor sizes. On the other hand, if the  
 626 solids residence time is prolonged to 10 min, the CaL process yields a lower SPECCA due to  
 627 promoted carbonation in the solid-state diffusion controlled phase.

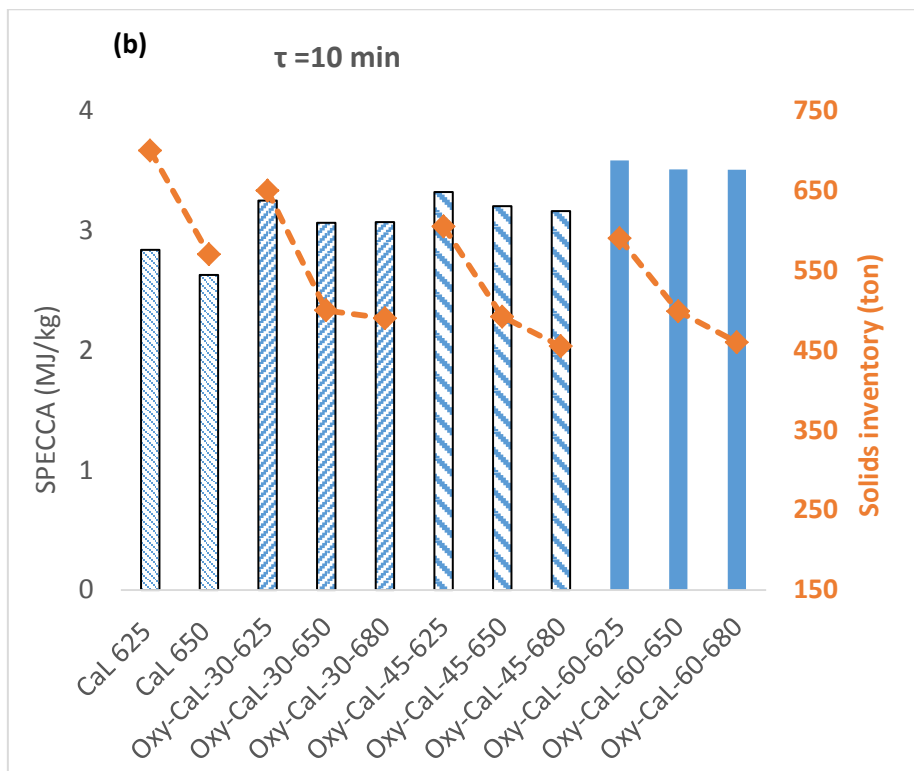
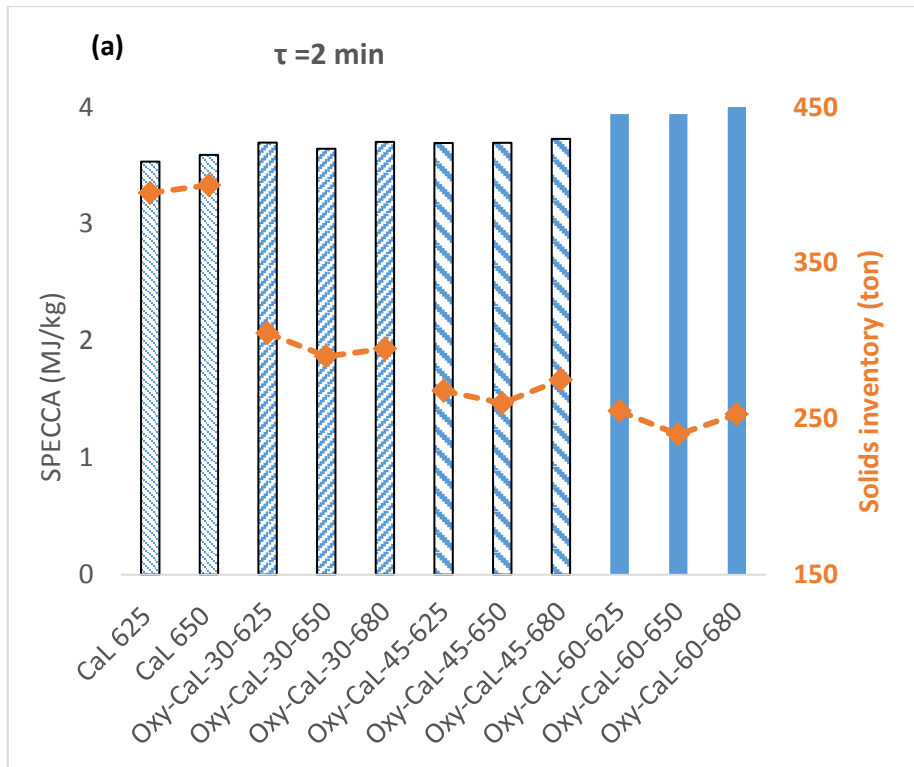
628

629 Figure 15 shows data on SPECCA and solids inventory for the CaL and Oxy-CaL (carbonator  
 630 temperatures of 625°C, 650°C and 680°C) systems for a fixed capture efficiency ( $E_{CO_2}=90\%$ ),  
 631 and short (2 min) and prolonged (10 min) solids residence times. As may be seen, for operation  
 632 under short residence times, the Oxy-CaL systems (especially Oxy-CaL-30 and 45) lead to a  
 633 considerable reduction of the solids inventory (from 400 tons for the CaL-650 system to 286  
 634 tons for Oxy-CaL-45-650) and therefore to a reduction of the CaL system size whereas the  
 635 SPECCA is only slightly increased (from 3.59 MJ/kg for the CaL-650 system to 3.7 MJ/kg  
 636 for the Oxy-CaL-45-650). Regarding the effect of carbonator temperature on the CaL and  
 637 Oxy-CaL systems for a given solids residence time of 2 min, there is not a clear evidence on  
 638 the optimum system choice (lower SPECCA). On the other hand, for longer residence times  
 639 (Figure 15b), the CaL-625 system shows a better performance in terms of efficiency albeit a  
 640 considerable higher solids inventory is required in this case. Thus, the CaL process advantage  
 641 is lost by a modest decrease of the carbonator temperature, which would require a  
 642 considerable increase of the solids inventory to ensure a high capture efficiency. At this point,  
 643 it is important to note that both CaL and Oxy-CaL systems yield a lower energy consumption  
 644 than the conventional oxy-fuel combustion process regardless of the carbonator temperature  
 645 and carbonator CO<sub>2</sub> concentration.

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651 Figure 14: SPECCA and solids inventory for the CaL and Oxy-CaL systems operating under a) short (2  
652 min) and b) prolonged (10 min) solids residence times. The effect of changing the carbonator temperature  
653 (between 625°C to 680°C) is also shown. Capture efficiency is fixed to 90%.

654

655 Concerning the Oxy-CaL-30 system operated under short residence times, it would allow  
656 reducing the solids inventory from 400 tons (CaL-625) system to 300 tons (-33% relative

657 reduction) and thus the reactors (carbonator and calciner) size could be decreased, which  
658 should be quantitatively assessed in future works. Thermal inertia and flexible operation of  
659 the installation must be also carefully addressed. These effects are amplified when the  
660 residence time is increased to 10 min for the CaL process. The efficiency penalty is then  
661 clearly reduced around 1 MJ/kgCO<sub>2</sub> but the solids inventory is increased over a 200% up to  
662 near 700 ton. Care should be taken in this case to design and operate a system with such a  
663 large thermal inertia due to solids heat capacity. Starts-up and shuts-down would be also  
664 risky. This could be a suitable option if the CO<sub>2</sub> capture system is to be used in base load  
665 power plants. When power plant flexibility is required, the small inventory option allowed by  
666 the Oxy-CaL hybrid system is preferred to better accommodate the installation to load  
667 changes.  
668

## 669 **6. Conclusions**

670  
671 This work analyzes a novel CO<sub>2</sub> capture system (Oxy-CaL) based on the integration of the  
672 CaL process with partial oxy-combustion, the latter used to raise the CO<sub>2</sub> concentration in the  
673 flue gas thus enhancing the CaL capture efficiency. Results from thermogravimetric analysis  
674 experiments are used to simulate the hybrid capture system when integrated into a coal fired  
675 power plant (CFPP). Energy penalty and capture efficiency are analyzed for diverse Oxy-CaL  
676 systems to reach a CO<sub>2</sub> vol% in the flue gas in the range 30-60% and the results are compared  
677 with those obtained for the pure CaL and oxy-combustion processes.  
678

679 Main highlights concluded from the study are:

- 680  
681 - A higher CO<sub>2</sub> capture efficiency is achieved by means of the Oxy-CaL hybrid system  
682 while the specific energy consumption per kg of CO<sub>2</sub> avoided (SPECCA) is kept  
683 below 4 MJ/kg, which is a typical value usually reported for oxy-combustion or  
684 amine-based CO<sub>2</sub> capture systems. Thus, CO<sub>2</sub> capture using an Oxy-CaL system could  
685 be a potentially attractive alternative to total oxy-combustion for newly erected CFPP.
- 686 - The CaL process seems to be the most advantageous CO<sub>2</sub> capture process when  
687 operating under relative long solids residence time in the carbonator. However, its  
688 performance has a strong sensitivity to the carbonation rate in the solid-state diffusion  
689 controlled stage, which depends critically on temperature within the range of  
690 temperatures practically attainable in the carbonator.
- 691 - The effect of varying the carbonation temperature is more relevant when operating  
692 under long residence times due to the strong sensitivity of CaO carbonation to  
693 temperature in the solid-state diffusion controlled stage, especially for the pure CaL  
694 process.
- 695 - The higher CO<sub>2</sub> capture efficiency using Oxy-CaL systems allows to reduce the fresh  
696 limestone makeup flow, which leads to a reduction of energy consumption when  
697 operating under solids residence times below 7.5 minutes.
- 698 - In spite that SPECCA in the CaL process could be somewhat smaller than for the  
699 Oxy-CaL system when operating under prolonged solids residence times, the latter

700 shows potentially important benefits regarding plant operation flexibility.  
 701 Substantially smaller amounts of solids inventory are needed in the Oxy-CaL system,  
 702 which would allow a more efficient response to load changes in coal fired power  
 703 plants.

704 In a future work, a detailed techno-economic study must be developed in order to assess  
 705 quantitatively the cost of CO<sub>2</sub> capture by means of the Oxy-CaL system as compared to CaL  
 706 and oxy-combustion and for same values of SPECCA and capture efficiency.  
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709  
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## 716 Notation

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$[CO_2]$	average CO <sub>2</sub> concentration, mol/m <sup>3</sup>	$T_{calc}$	calciner temperature, °C
$[CO_2]_{eq}$	equilibrium concentration of CO <sub>2</sub> , mol/m <sup>3</sup>	$T_{carb}$	carbonator temperature, °C
$D^*_{eff}$	equivalent diffusion constant, m <sup>2</sup> /(mol · s)	$T_{vv}$	live steam temperature, °C
$E$	emissions ratio after CO <sub>2</sub> capture, kg CO <sub>2</sub> /kWh	$t_0$	time lag of TGA multicycle test, s
$E_{CO_2}$	carbon capture efficiency	$t_{FRP}$	time of the carbonation fast phase, s
$E_{ref}$	emissions ratio before CaL, kg CO <sub>2</sub> /kWh	$t_{max}$	total carbonation time, s
$E_{max}$	maximum capture efficiency	$v_{CO_2}$	CO <sub>2</sub> v/v concentration at CFPP outlet
$F_{CO_2}$	mole flow rate of CO <sub>2</sub> in flue gas entering the carbonator, kmol/h	$v_{O_2}$	O <sub>2</sub> v/v concentration at CFPP outlet
$F_0$	mole flow rate of fresh makeup limestone, kmol/h	$W_s$	solid inventory in the carbonator per MW of a typical power plant, kg
$F_R$	mole flow rate of CO <sub>2</sub> in flue gas entering the carbonator, kmol/h	$\dot{W}_{ASU}$	power consumption in the ASU, MW
$F_{fg}$	flue gas molar flow rate, kmol/s	$\dot{W}_{sec}$	net power production in secondary steam cycle, MW
$FRP$	fast reaction controlled phase	$\dot{W}_{comp,CO_2}$	power consumption in CO <sub>2</sub> compression, MW
$SDP$	solid-state diffusion controlled phase	$\dot{W}_{comp,fg}$	power consumption in flue gas compression, MW
$k_s$	intrinsic kinetic constant m <sup>4</sup> /(mol · s)	$\dot{W}_{solid}$	power consumption in solids transport, MW
$k$	deactivation constant of a sorbent particle	$X$	carbonation degree of a CaO particle
$\dot{m}$	mass flow, kg/s	$X_{ave}$	average conversion of the sorbent
$\dot{m}_{CO_2,calc}$	CO <sub>2</sub> mass flow exiting the calciner, kg/s	$X_{ave,SDP}$	average conversion of the sorbent in the diffusion phase
$\dot{m}_{gas,calc}$	Total gas mass flow exiting the calciner, kg/s	$X_{ave,FRP}$	average conversion of the sorbent in the kinetic phase
		$W_s$	solid inventory in the carbonator per MW of a

			typical power plant, kg
$N_{Ca}$	mol of Ca in the carbonator, mol	$x_{CO_2}$	CO <sub>2</sub> Molar fraction exiting the plant
$P$	Pressure, bar	$x_{O_2}$	O <sub>2</sub> Molar fraction exiting the plant
$P_{vv}$	live steam pressure, bar	$y_{CO_2,in}$	CO <sub>2</sub> molar fraction at carbonator inlet
$P_{HE,solids}$	solid-solid thermal power exchanged, MW	$y_{CO_2,eq}$	CO <sub>2</sub> molar fraction at carbonation equilibrium
$SPECCA$	energy consumption for kg CO <sub>2</sub> avoided, MJ/Kg CO <sub>2</sub>	$\eta_{boiler}$	boiler efficiency
$r_{ave,SDP}$	average reaction rate in the diffusion regime, s <sup>-1</sup>	$\eta_{CFPP}$	coal fire power plant efficiency
$r_{ave,FRP}$	average reaction rate in the kinetic regime, s <sup>-1</sup>	$\eta_{ref}$	reference plant efficiency
$S_{ave}$	average surface area available for reaction, m <sup>-1</sup>	$\eta_{int}$	new global efficiency (CFPP-capture system)
$t$	time, s	$\tau$	average residence time in the carbonator, s

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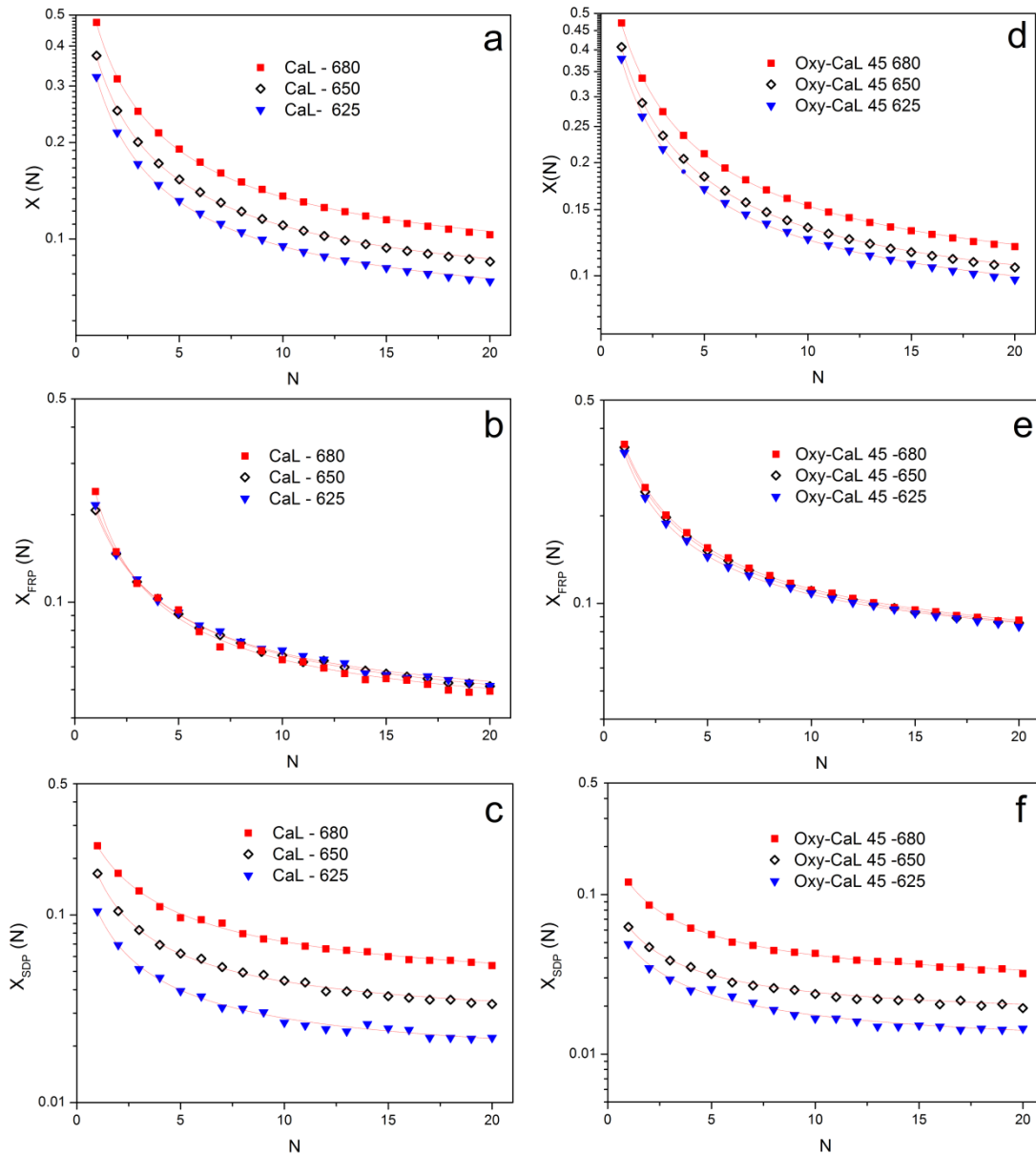
## 719 **Appendix A**

720

721 Figure 15 shows data on the multicycle overall CaO conversion ( $X(N)$ ),  $X_{FRP}$  and  $X_{SDP}$  as a  
722 function of the cycle number for the tests carried out under CaL and Oxy-CaL-45 conditions  
723 and for the diverse carbonation temperatures employed (625, 650°C and 680°C). Values of the  
724 deactivation rate constant and residual conversion obtained from the fittings of Eq. 1 to  
725 experimental data are also shown in Table 4.

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728  
 729 Figure 15. (a, d) Overall CaO conversion versus the cycle number for carbonation/calcination cycles  
 730 carried out under CaL and Oxy-CaL-45 conditions. (b, e) Conversion in the fast reaction controlled  
 731 phase. (c, f) Conversion in the solid-state diffusion controlled phase. Carbonation is carried out under  
 732 15% vol CO<sub>2</sub> (CaL conditions) and 45% vol CO<sub>2</sub> (Oxy-CaL-45 conditions) at 625°C, 650°C and 680°C  
 733 as indicated. Calcination is performed at 950°C (70% CO<sub>2</sub>/30% air vol/vol) in all the tests.

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744 Table 4: Values of the deactivation rate constant  $\kappa$  and residual conversion  $X_r$  obtained from the best fits of  
745 Eq. (1) to TGA experimental data for carbonation at different temperatures (625, 650, and 680°C) and CO<sub>2</sub>  
746 concentrations (15% vol in the CaL tests; 30%, 45%, and 60% vol in the Oxy-CaL tests).

<b>T carb =625°C</b>		<b>CaL-625</b>	<b>Oxy-CaL30-625</b>	<b>Oxy-CaL45-625</b>	<b>Oxy-CaL60-625</b>
$X_{overall}$	$X_1$	0.321	0.338	0.378	0.400
	$\kappa$	0.753	0.665	0.675	0.623
	$X_r$	0.053	0.067	0.072	0.077
	$R_{sqr}$	0.999	0.999	0.999	0.999
$X_{FRP}$	$X_1$	0.216	0.260	0.329	0.350
	$\kappa$	0.737	0.697	0.683	0.647
	$X_r$	0.038	0.056	0.062	0.067
	$R_{sqr}$	0.999	0.998	0.999	0.999
$X_{SDP}$	$X_1$	0.105	0.078	0.049	0.050
	$\kappa$	0.792	0.573	0.615	0.464
	$X_r$	0.015	0.011	0.010	0.0103
	$R_{sqr}$	0.997	0.996	0.990	0.994
<b>T carb =650°C</b>		<b>CaL-650</b>	<b>Oxy-CaL30-650</b>	<b>Oxy-CaL45-650</b>	<b>Oxy-CaL60-650</b>
$X_{overall}$	$X_1$	0.373	0.388	0.407	0.425
	$\kappa$	0.731	0.667	0.651	0.633
	$X_r$	0.061	0.070	0.076	0.081
	$R_{sqr}$	0.999	0.999	0.999	0.999
$X_{FRP}$	$X_1$	0.207	0.293	0.344	0.374
	$\kappa$	0.660	0.661	0.641	0.674
	$X_r$	0.037	0.053	0.060	0.068
	$R_{sqr}$	0.999	0.999	0.999	0.998
$X_{SDP}$	$X_1$	0.166	0.095	0.063	0.051
	$\kappa$	0.828	0.686	0.711	0.633
	$X_r$	0.025	0.016	0.016	0.014
	$R_{sqr}$	0.998	0.998	0.993	0.980
<b>T carb =680°C</b>		<b>CaL-680</b>	<b>Oxy-CaL30-680</b>	<b>Oxy-CaL45-680</b>	<b>Oxy-CaL60-680</b>
$X_{overall}$	$X_1$	0.474	0.472	0.472	0.474
	$\kappa$	0.737	0.681	0.636	0.636
	$X_r$	0.074	0.081	0.085	0.086
	$R_{sqr}$	0.999	0.999	0.999	0.999
$X_{FRP}$	$X_1$	0.240	0.299	0.352	0.391
	$\kappa$	0.876	0.716	0.630	0.673
	$X_r$	0.037	0.057	0.060	0.066
	$R_{sqr}$	0.997	0.999	0.999	0.999
$X_{SDP}$	$X_1$	0.233	0.173	0.120	0.083
	$\kappa$	0.609	0.627	0.655	0.478
	$X_r$	0.037	0.025	0.025	0.020
	$R_{sqr}$	0.998	0.999	0.999	0.990

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	parameter	Reference CFFP (air combustion)	oxy- combustion	CaL	Oxy- CaL 30	Oxy- CaL 45	Oxy- CaL 60
CFPP	$\dot{m}_{coal}$ (kg/s)	42.20	55.05	42.20	46.10	47.50	48.15
	$\dot{m}_{air}$ (kg/s)	475	-	475	208.90	100.20	43.54
	$\dot{m}_{O_2}$ (kg/s)	-	136.91	-	68.85	96.35	110.503
	$\gamma_{fg}$	-	0.78	-	0.63	0.72	0.75
	$F_{fg}$ (kmol/s)	17.12	3.85	17.12	10.13	7.25	5.74
	$F_{CO_2}$ (kmol/s)	2.60	3.39	2.60	2.84	2.93	2.96
	$v_{CO_2}$	0.15	0.89	0.15	0.30	0.45	0.60
	$v_{O_2}$	0.023	0.025	0.023	0.025	0.025	0.025
	$\eta_{boiler}$	0.90	0.90	0.90	0.90	0.90	0.90
	$\eta_{CFPP}$	0.377	0.287	0.377	0.351	0.344	0.337
	net work (MW)	490.47	488.80	490.47	498.30	502.30	499.75
	Penalty	-	<b>9.02%</b>	-	<b>2.64%</b>	<b>3.47%</b>	<b>4.03%</b>
SPECCA (MJ/kgCO <sub>2</sub> )	-	<b>4.06</b>	-	<b>0.94</b>	<b>1.25</b>	<b>1.50</b>	
CaL	$F_R/F_{CO_2}$	-	-	15	15	15	15
	$F_0/F_{CO_2}$	-	-	0.05	0.05	0.05	0.05
	$\tau$ (min)	-	-	3.05	2.79	2.71	2.68
	$T_{calc}$ (°C)	-	-	950	950	950	950
	$T_{carb}$ (°C)	-	-	650	650	650	650
	$E_{CO_2}$	-	-	0.827	0.950	0.976	0.981
	$\dot{m}_{coal}$ (kg/s)	-	-	18.48	22.34	23.40	23.84
	$\dot{m}_{O_2}$ (kg/s)	-	-	48.00	58.41	61.00	62.49
	$\gamma_{fg}$	-	-	0.7	0.7	0.7	0.7
	$\dot{m}_{gas,calc}$ (kg/s)	-	-	165.07	203.6	214	218.34
	$\dot{m}_{CO_2,calc}$ (kg/s)	-	-	150.45	185.70	195.40	199.19
	$x_{CO_2}$	-	-	0.030	0.019	0.017	0.026
	$x_{O_2}$	-	-	0.028	0.032	0.042	0.058
	$P_{HE,solids}$	-	-	640.92	697.60	717.80	727.29
	$\dot{W}_{sec}$ (MW)	-	-	75.80	113.90	126.46	130.88
	$\dot{W}_{solids}$ (MW)	-	-	45.70	49.72	51.16	51.82
	$\dot{W}_{ASU}$ (MW)	-	-	34.56	42.05	43.92	44.99
	$\dot{W}_{comp,CO_2}$ (MW)	-	-	57.55	71.01	74.59	75.20
	$\dot{W}_{comp,FG}$ (MW)	-	-	12.53	7.27	4.90	3.66
	$\eta_{int}$	-	-	0.303	0.291	0.288	0.285
Penalty	-	-	<b>7.43%</b>	<b>6.04%</b>	<b>5.48%</b>	<b>5.27%</b>	
SPECCA (MJ/kgCO <sub>2</sub> )	-	-	<b>3.28</b>	<b>2.63</b>	<b>2.37</b>	<b>2.29</b>	
Total	$\dot{m}_{coal,total}$	-	-	60.68	68.44	70.90	71.99
	Penalty <sub>total</sub>	-	<b>9.02%</b>	<b>7.43%</b>	<b>8.68%</b>	<b>8.95%</b>	<b>9.30%</b>
	SPECCA <sub>total</sub> (MJ/kgCO <sub>2</sub> )	-	<b>4.06</b>	<b>3.28</b>	<b>3.56</b>	<b>3.62</b>	<b>3.79</b>



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753 **References**

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- 755 [1] IEA. Energy Technology Perspectives 2012. 2012. doi:10.1787/energy\_tech-2012-en.
- 756 [2] Tokimatsu K, Konishi S, Ishihara K, Tezuka T, Yasuoka R, Nishio M. Role of  
757 innovative technologies under the global zero emissions scenarios. *Appl Energy*  
758 2016;162:1483–93. doi:10.1016/j.apenergy.2015.02.051.
- 759 [3] Goto K, Yogo K, Higashii T. A review of efficiency penalty in a coal-fired power plant  
760 with post-combustion CO<sub>2</sub> capture. *Appl Energy* 2013;111:710–20.  
761 doi:10.1016/j.apenergy.2013.05.020.
- 762 [4] Bevan N, Boston A. CCS Forum Report. 2016.
- 763 [5] Wang M, Lawal A, Stephenson P, Sidders J, Ramshaw C, Hill W, et al. Post-  
764 combustion CO<sub>2</sub> capture with chemical absorption: A state-of-the-art review. *Chem*  
765 *Eng Res Des* 2011;89:1609–24. doi:10.1016/j.cherd.2010.11.005.
- 766 [6] Perejon A, Romeo LM, Lara Y, Lisbona P, Valverde JM. The Calcium-Looping  
767 technology for CO<sub>2</sub> capture: On the important roles of energy integration and sorbent  
768 behavior. *Appl Energy* 2015;162:787–807. doi:10.1016/j.apenergy.2015.10.121.
- 769 [7] Politecnico di Milano – Alstom UK (CAESAR project). European best practice  
770 guidelines for assessment of CO<sub>2</sub> capture technologies. 2011.
- 771 [8] Aaron D, Tsouris C. Separation of CO<sub>2</sub> from Flue Gas: A Review. *Sep Sci Technol*  
772 2005;40:321–48. doi:10.1081/SS-200042244.
- 773 [9] Wang M, Joel AS, Ramshaw C, Eimer D, Musa NM. Process intensification for post-  
774 combustion CO<sub>2</sub> capture with chemical absorption: A critical review. *Appl Energy*  
775 2015;158:275–91. doi:10.1016/j.apenergy.2015.08.083.
- 776 [10] Luis P. Use of monoethanolamine (MEA) for CO<sub>2</sub> capture in a global scenario:  
777 Consequences and alternatives. *Desalination* 2016;380:93–9.  
778 doi:10.1016/j.desal.2015.08.004.
- 779 [11] Rey A, Gouedard C, Ledirac N, Cohen M, Dugay J, Vial J, et al. Amine degradation in  
780 CO<sub>2</sub> capture. 2. New degradation products of MEA. Pyrazine and alkylpyrazines:  
781 Analysis, mechanism of formation and toxicity. *Int J Greenh Gas Control*  
782 2013;19:576–83. doi:10.1016/j.ijggc.2013.10.018.
- 783 [12] Shimizu T, Hiramata T, Hosoda H, Kitano K, Inagaki M, Tejima K. A Twin Fluid-Bed  
784 Reactor for Removal of CO<sub>2</sub> from Combustion Processes. *Chem Eng Res Des*  
785 1999;77:62–8. doi:10.1205/026387699525882.
- 786 [13] Ortiz C, Valverde JM, Chacartegui R. Energy Consumption for CO<sub>2</sub> Capture by means  
787 of the Calcium Looping Process: A Comparative Analysis using Limestone, Dolomite,  
788 and Steel Slag. *Energy Technol* 2016:1–12. doi:10.1002/ente.201600390.
- 789 [14] Ströhle J, Junk M, Kremer J, Galloy A, Eppe B. Carbonate looping experiments in a 1  
790 MWth pilot plant and model validation. *Fuel* 2014;127:13–22.  
791 doi:10.1016/j.fuel.2013.12.043.
- 792 [15] Arias B, Diego ME, Abanades JC, Lorenzo M, Diaz L, Martínez D, et al.

- 793 Demonstration of steady state CO<sub>2</sub> capture in a 1.7MWth calcium looping pilot. *Int J*  
794 *Greenh Gas Control* 2013;18:237–45. doi:10.1016/j.ijggc.2013.07.014.
- 795 [16] Hanak DP, Anthony EJ, Manovic V. A review of developments in pilot-plant testing  
796 and modelling of calcium looping process for CO<sub>2</sub> capture from power generation  
797 systems. *Energy Environ Sci* 2015;8:2199–249. doi:10.1039/C5EE01228G.
- 798 [17] Abanades JC, Grasa G, Alonso M, Rodriguez N, Anthony EJ, Romeo LM. Cost  
799 structure of a postcombustion CO<sub>2</sub> capture system using CaO. *Environ Sci Technol*  
800 2007;41:5523–7. doi:10.1021/es070099a.
- 801 [18] Scheffknecht G, Al-Makhadmeh L, Schnell U, Maier J. Oxy-fuel coal combustion-A  
802 review of the current state-of-the-art. *Int J Greenh Gas Control* 2011;5:16–35.  
803 doi:10.1016/j.ijggc.2011.05.020.
- 804 [19] Escudero AI, Espatolero S, Romeo LM, Lara Y, Paufigue C, Lesort A-L, et al.  
805 Minimization of CO<sub>2</sub> capture energy penalty in second generation oxy-fuel power  
806 plants. *Appl Therm Eng* 2016;103:274–81. doi:10.1016/j.applthermaleng.2016.04.116.
- 807 [20] Jin B, Zhao H, Zheng C. Thermo-economic cost analysis of CO<sub>2</sub> compression and  
808 purification unit in oxy-combustion power plants. *Bo. Energy* 2015;83:416–30.  
809 doi:10.1016/j.energy.2015.02.039.
- 810 [21] Posch S, Haider M. Optimization of CO<sub>2</sub> compression and purification units  
811 (CO<sub>2</sub>CPU) for CCS power plants. *Fuel* 2012;101:254–63.  
812 doi:10.1016/j.fuel.2011.07.039.
- 813 [22] Escudero AI, Espatolero S, Romeo LM. Oxy-combustion power plant integration in an  
814 oil refinery to reduce CO<sub>2</sub> emissions. *Int J Greenh Gas Control* 2016;45:118–29.  
815 doi:10.1016/j.ijggc.2015.12.018.
- 816 [23] Kather A, Scheffknecht G. The oxycoal process with cryogenic oxygen supply.  
817 *Naturwissenschaften* 2009;96:993–1010. doi:10.1007/s00114-009-0557-2.
- 818 [24] Vega F, Sanna A, Maroto-Valer MM, Navarrete B, Abad-Correa D. Study of the MEA  
819 degradation in a CO<sub>2</sub> capture process based on partial oxy-combustion approach. *Int J*  
820 *Greenh Gas Control* 2016;54:160–7. doi:10.1016/j.ijggc.2016.09.007.
- 821 [25] Ylätaalo J, Parkkinen J, Ritvanen J, Tynjälä T, Hyppänen T. Modeling of the oxy-  
822 combustion calciner in the post-combustion calcium looping process. *Fuel*  
823 2013;113:770–9. doi:10.1016/j.fuel.2012.11.041.
- 824 [26] Martínez I, Grasa G, Murillo R, Arias B, Abanades JC. Modelling the continuous  
825 calcination of CaCO<sub>3</sub> in a Ca-looping system. *Chem Eng J* 2013;215–216:174–81.  
826 doi:10.1016/j.cej.2012.09.134.
- 827 [27] Koga N, Criado JM. The influence of mass transfer phenomena on the kinetic analysis  
828 for the thermal decomposition of calcium carbonate by constant rate thermal analysis  
829 (CRTA) under vacuum. *Int J Chem Kinet* 1998;30:737–44. doi:10.1002/(SICI)1097-  
830 4601(1998)30:10<737::AID-KIN6>3.0.CO;2-W.
- 831 [28] Alonso M, Criado Y a., Abanades JC, Grasa G. Undesired effects in the determination  
832 of CO<sub>2</sub> carrying capacities of CaO during TG testing. *Fuel* 2014;127:52–61.  
833 doi:10.1016/j.fuel.2013.08.005.
- 834 [29] Grasa G, Murillo R, Alonso M, Abanades JC. Application of the random pore model to  
835 the carbonation cyclic reaction. *AIChE J* 2009;55:1246–55. doi:10.1002/aic.11746.

- 836 [30] Bhatia SK, Perlmutter DD. Effect of the product layer on the kinetics of the carbon  
837 dioxide-lime reaction. *AIChE J* 1983;29:79–86. doi:10.1002/aic.690290111.
- 838 [31] Sun Z, Luo S, Qi P, Fan LS. Ionic diffusion through Calcite (CaCO<sub>3</sub>) layer during the  
839 reaction of CaO and CO<sub>2</sub>. *Chem Eng Sci* 2012;81:164–8.  
840 doi:10.1016/j.ces.2012.05.042.
- 841 [32] Bhatia SK, Perlmutter DD. Effect of the product layer on the kinetics of the CO<sub>2</sub>-lime  
842 reaction. *AIChE J* 1983;29:79–86. doi:10.1002/aic.690290111.
- 843 [33] Anderson TF. Self-diffusion of carbon and oxygen in calcite by isotope exchange with  
844 carbon dioxide. *J Geophys Res* 1969;74:3918–32. doi:10.1029/JB074i015p03918.
- 845 [34] Grasa GS, Abanades JC. CO<sub>2</sub> Capture Capacity of CaO in Long Series of  
846 Carbonation/Calcination Cycles. *Ind Eng Chem Res* 2006;45:8846–51.  
847 doi:10.1021/ie0606946.
- 848 [35] Valverde JM. A model on the CaO multicyclic conversion in the Ca-looping process.  
849 *Chem Eng J* 2013;228:1195–206. doi:10.1016/j.cej.2013.05.023.
- 850 [36] Blamey J, Anthony EJ, Wang J, Fennell PS. The calcium looping cycle for large-scale  
851 CO<sub>2</sub> capture. *Prog Energy Combust Sci* 2010;36:260–79.  
852 doi:10.1016/j.peccs.2009.10.001.
- 853 [37] Zhu Y, Wu S, Wang X. Nano CaO grain characteristics and growth model under  
854 calcination. *Chem Eng J* 2011;175:512–8. doi:10.1016/j.cej.2011.09.084.
- 855 [38] Ortiz C, Chacartegui R, Valverde J, Becerra J, Perez-Maqueda L. A new model of the  
856 carbonator reactor in the calcium looping technology for post-combustion CO<sub>2</sub>  
857 capture. *FUEL* 2015;160:328–38. doi:10.1016/j.fuel.2015.07.095.
- 858 [39] USA National Energy Technology Laboratory. Detailed Coal Specifications Quality  
859 Guidelines for Energy System Studies. Off Progr Perform Benefits 2012.  
860 doi:[http://www.netl.doe.gov/File%20Library/research/energy%20analysis/publications/  
861 QGESS\\_DetailCoalSpecs\\_Rev4\\_20130510.pdf](http://www.netl.doe.gov/File%20Library/research/energy%20analysis/publications/QGESS_DetailCoalSpecs_Rev4_20130510.pdf).
- 862 [40] Romano MC. Ultra-high CO<sub>2</sub> capture efficiency in CFB oxyfuel power plants by  
863 calcium looping process for CO<sub>2</sub> recovery from purification units vent gas. *Int J  
864 Greenh Gas Control* 2013;18:57–67. doi:10.1016/j.ijggc.2013.07.002.
- 865 [41] Zhang X, Bauer C, Mutel CL, Volkart K. Life Cycle Assessment of Power-to-Gas:  
866 Approaches, system variations and their environmental implications. *Appl Energy*  
867 2017;190:326–38. doi:10.1016/j.apenergy.2016.12.098.
- 868 [42] Perez-Fortes M, Schoneberger JC, Boulamanti A, Tzimas E. Methanol synthesis using  
869 captured CO<sub>2</sub> as raw material: Techno-economic and environmental assessment. *Appl  
870 Energy* 2016;161:718–32. doi:10.1016/j.apenergy.2015.07.067.
- 871 [43] Pei X, He B, Yan L, Wang C, Song W, Song J. Process simulation of oxy-fuel  
872 combustion for a 300 MW pulverized coal-fired power plant using Aspen Plus. *Energy  
873 Convers Manag* 2013;76:581–7. doi:10.1016/j.enconman.2013.08.007.
- 874 [44] Romano M, Martínez I, Murillo R, Arstad B. Guidelines for modeling and simulation  
875 of Ca-looping processes. 2012.
- 876 [45] Martínez I, Murillo R, Grasa G, Abanades JC. Integration of a Ca-looping system for  
877 CO<sub>2</sub> capture in an existing power plant. *Energy Procedia* 2011;4:1699–706.  
878 doi:10.1016/j.egypro.2011.02.043.

- 879 [46] Hanak DP, Powell D, Manovic V. Techno-economic analysis of oxy-combustion coal-  
880 fired power plant with cryogenic oxygen storage. *Appl Energy* 2017;191:193–203.  
881 doi:10.1016/j.apenergy.2017.01.049.
- 882 [47] Ortiz C, Chacartegui R, Valverde JM, Becerra JA. A new integration model of the  
883 calcium looping technology into coal fired power plants for CO<sub>2</sub> capture. *Appl Energy*  
884 2016;169:408–20. doi:10.1016/j.apenergy.2016.02.050.
- 885 [48] Ylätaalo J, Ritvanen J, Tynjälä T, Hyppänen T. Model based scale-up study of the  
886 calcium looping process. *Fuel* 2014;115:329–37. doi:10.1016/j.fuel.2013.07.036.
- 887 [49] Romano MC. Modeling the carbonator of a Ca-looping process for CO<sub>2</sub> capture from  
888 power plant flue gas. *Chem Eng Sci* 2012;69:257–69. doi:10.1016/j.ces.2011.10.041.
- 889 [50] Vorrias I, Atsonios K, Nikolopoulos A, Nikolopoulos N, Grammelis P, Kakaras E.  
890 Calcium looping for CO<sub>2</sub> capture from a lignite fired power plant. *Fuel* 2013;113:826–  
891 36. doi:10.1016/j.fuel.2012.12.087.
- 892 [51] Dieter H, Bidwe AR, Varela-duelli G, Charitos A, Hawthorne C. Development of the  
893 calcium looping CO<sub>2</sub> capture technology from lab to pilot scale at IFK , University of  
894 Stuttgart. *Fuel* 2014;127:23–37. doi:10.1016/j.fuel.2014.01.063.
- 895 [52] Charitos A, Rodríguez N, Hawthorne C, Alonso M, Zieba M, Arias B, et al.  
896 Experimental Validation of the Calcium Looping CO<sub>2</sub> Capture Process with Two  
897 Circulating Fluidized Bed Carbonator Reactors. *Ind Eng Chem Res* 2011;50:9685–95.  
898 doi:10.1021/ie200579f.
- 899 [53] Kunii D, Levenspiel O. The K-L reactor model for circulating fluidized beds. *Chem*  
900 *Eng Sci* 2000;55:4563–70. doi:10.1016/S0009-2509(00)00073-7.
- 901 [54] Kunii D, Levenspiel O. Circulating fluidized-bed reactors. *Chem Eng Sci*  
902 1997;52:2471–82. doi:10.1016/S0009-2509(97)00066-3.
- 903 [55] Edwards SEB, Materić V. Calcium looping in solar power generation plants. *Sol*  
904 *Energy* 2012;86:2494–503. doi:10.1016/j.solener.2012.05.019.
- 905 [56] Martínez I, Grasa G, Parkkinen J, Tynjälä T, Hyppänen T, Murillo R, et al. Review and  
906 research needs of Ca-Looping systems modelling for post-combustion CO<sub>2</sub> capture  
907 applications. *Int J Greenh Gas Control* 2016;50:271–304.  
908 doi:10.1016/j.ijggc.2016.04.002.
- 909 [57] Hanak DP, Kolios AJ, Manovic V. Comparison of probabilistic performance of  
910 calcium looping and chemical solvent scrubbing retrofits for CO<sub>2</sub> capture from coal-  
911 fired power plant. *Appl Energy* 2016;172:323–36. doi:10.1016/j.apenergy.2016.03.102.

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