Fragmentation of biomass-templated CaO-based pellets

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Abstract  
The use of biomass-templating materials with a cheap production method as an enhanced sorbent for CO\textsubscript{2} uptake has been proposed recently. However, the attrition and fragmentation behaviour of this type of material, which is a vital parameter for calcium looping sorbents, has not yet been investigated in detail. In this work the attrition and fragmentation behaviour of biomass-templated sorbents is investigated. Three types of materials were prepared using a mechanical pelletiser: 1. lime and cement (LC); 2. lime and flour (LF); and 3. lime, cement and flour (LCF). These samples were heat treated in a pressurised heated strip reactor (PHSR) and in a bubbling fluidised bed (BFB) and changes in particle size distribution were measured to assess fragmentation. Results indicated that the addition of biomass enhances the propensity to undergo fragmentation. Upon heat treatment in the PHSR the particle size of LC was not modified significantly; on the contrary the mean particle diameter of LF decreased from 520 \(\mu\text{m}\) to 116 \(\mu\text{m}\) and that of LCF from 524 \(\mu\text{m}\) to 290 \(\mu\text{m}\). Fragmentation tests in the BFB confirmed the trend: 67\% of the particles of LF fragmented, against 53\% of LCF and 18\% of LC samples. The addition of biomass to the LC samples partially counteracts this performance degradation with respect to attrition. However, calcium aluminate pellets (LC) showed the lowest rate of fragmentation amongst all of the samples tested.

Keywords: Calcium looping, pellets, biomass-templating, fragmentation
1 Introduction

Calcium looping (CaL) is a second-generation carbon capture technology, which uses a lime sorbent in dual fluidised-bed reactors; this technology depends on the following reversible exothermic calcium oxide carbonation reaction:

\[ \text{CaO} + \text{CO}_2 \leftrightarrow \text{CaCO}_3 \]  \hspace{1cm} (1)

The typical reactor set-up consists of two interconnected fluidised-bed reactors. In the first reactor (the carbonator) the CaO-based sorbent captures CO\(_2\) from power plant flue gas; this reaction occurs at a practical rate at 650-700°C [1-3]. The carbonated sorbent is then transferred to the second reactor (the calciner) where CO\(_2\) is released at high temperatures (850-950°C). The regenerated material is then returned to the carbonator for the next cycle. However, there are several challenges with CaO-based sorbent whose CO\(_2\) uptake decreases with increasing number of carbonation/calcination cycles. This decline in activity is mainly due to sintering during calcination because of the high temperatures necessary for calcination [4-8]. The CO\(_2\) capture capacity of the fresh sorbent drops quickly during the initial cycles until an asymptotic value is achieved after about 20 or 30 cycles, which then remains almost constant over subsequent cycles and adopts typical values of about 0.08 g CO\(_2\)/g sorbent in the case of limestone [9]. This reduction in performance can be partially compensated by increasing the Ca/C ratio in the reactor (by increasing the purge of spent sorbent and the make-up ratio) or by modifying the properties of the particles [10]. However, this deactivation can also be caused by sulphation or ash fouling [11].

Natural sorbents (limestone and dolomite) are attractive due to their low cost, ready availability and, in the case of limestones, the potential suitability of the CaL purge material for the cement industry [12, 13]. However, significant research efforts are being made to modify limestone sorbents or create new synthetic sorbents using techniques such as sol-gel combustion [14-18], organic acid modifications [19-23], co-precipitation [24, 25] and granulation [26-32]. Such materials exhibited higher CO\(_2\) uptake, in general, when compared to natural lime-based sorbents. However, the cost of these sorbents increases due to such complex production procedures and the cost of the additives and may become prohibitively expensive.

One of the methods proposed to improve the performance of calcium looping sorbents was biomass templating [30]. Such biomass is potentially a cheap material for increasing the porosity of pelletised sorbents. Ridha et al. [30] observed that the capture capacity was 0.41 g CO\(_2\)/g sorbent after 20 cycles in the presence of 15% steam for sorbents with 10% of powdered leaves incorporated into the sorbent. This was an increase of 33.3% when compared to the untemplated materials after both samples underwent 20 cycles. The use of flour as a biomass-templating material has been studied by Erans et al. [32]. These materials were tested in both a thermogravimetric analyser (TGA) and a bubbling fluidised bed (BFB), and the synthesised materials were shown to exhibit better performance than natural...
limestone. However, under BFB conditions the templated materials demonstrated higher rates of fragmentation and attrition compared to calcium aluminate pellets without the addition of biomass.

Attrition of lime based sorbents has been extensively investigated in previous studies [33-36]. There are several attrition/fragmentation mechanisms: primary fragmentation, which occurs when the sorbent is injected into the reactor due to thermal stresses and overpressures caused by CO$_2$ release from the calcination reaction; secondary fragmentation, which occurs due to mechanical stresses from collisions between particles and bed internals; and attrition by abrasion, which is also caused by mechanical stresses but generates finer particles when compared to secondary fragmentation [37]. It has also been reported that the attrition rate was higher during the initial cycles and then subsequently decreased [38, 39].

Previous results have shown the beneficial effect of biomass addition for the CO$_2$ uptake, as well as demonstrating the enhanced porous structure of the templated samples. However, there are discrepancies between TGA results and BFB results; and these differences are believed to be due to attrition and fragmentation. This work explores the effect of biomass templating in calcium aluminate pellets with regard to fragmentation. Three different types of materials: one with the addition of calcium aluminate cement (LC), another with flour addition (LF) and one with both (LCF) have been tested in two different types of reactors; namely a pressurised heated strip reactor (PHSR) and bubbling fluidised bed under different conditions.

2 Experimental

2.1 Materials
Longcal limestone from the UK was used as a lime precursor. Commercial calcium aluminate cement, CA-14, manufactured by Almantis, was used as a binder in the pelletisation process and as a source of Al$_2$O$_3$. Commercial flour was used as the biomass templating material.

2.2 Pellet preparation procedure
Three types of materials were produced: (i) 10% calcium aluminate cement and 90% calcined limestone (LC); (ii) 10% flour and 90% calcined limestone (LF); and (iii) 10% flour, 10% calcium aluminate cement and 80% calcined limestone (LCF). The particles were prepared introducing the desired proportional quantities in 1 kg batches into a pelletiser vessel (4 L). The mixing took place inside the vessel by means of a chopper and agitator under a continuous water spray. A more detailed explanation of this procedure can be found elsewhere [27]. After pelletisation of the samples, the particles were sieved to different particle sizes. The material was air dried for 24 h before storage. The weight percentage of materials used in each sample can be found in Table 1.
Table 1: Materials used

<table>
<thead>
<tr>
<th>Sample</th>
<th>Lime (wt %)</th>
<th>Calcium aluminate cement (wt %)</th>
<th>Flour (wt %)</th>
</tr>
</thead>
<tbody>
<tr>
<td>LC</td>
<td>90</td>
<td>10</td>
<td>0</td>
</tr>
<tr>
<td>LF</td>
<td>90</td>
<td>0</td>
<td>10</td>
</tr>
<tr>
<td>LCF</td>
<td>80</td>
<td>10</td>
<td>10</td>
</tr>
</tbody>
</table>

2.3 Fragmentation experiments

For the fragmentation experiments two experimental systems have been used: a pressurised heated strip reactor and a bubbling fluidised bed.

The apparatus used for the first round of tests (the PHSR) achieves a heating rate of 4000°C/s. Batches of particles of 500-710 μm are placed on the strip and heated up by physical contact with the strip and by thermal radiation from the semi-spherical cover of the reactor. Further details of this reactor are available elsewhere [40]. Tests have been carried out at 1 bar and 950°C in pure N₂. The final temperature was held for 30 s in all experiments. After the test, particles were recovered and the experiments were repeated on fresh particles numerous times (~10-15 times) in order to collect a sufficiently large amount of material to perform further particle size analyses.

Further fragmentation tests were carried out in a lab-scale BFB in order to reproduce conditions typical of the first calcinations step. The BFB had a 40 mm ID, and was operated at atmospheric pressure and heated to the desired temperature by means of an external electric furnace. Calcination of the particles was performed under two conditions: 100% vol air and 70% vol CO₂/30% vol air. The calcination time for all the experiments was 20 min to ensure complete calcination. These tests were repeated for two different particle size ranges: 500-710 μm and 250-500 μm.

For the BFB tests, 20 g of sorbent was diluted in 150 g of silica (particle size distribution of 850 -1000 μm) sand to avoid excessive decrease of temperature in the bed during calcination, due to the endothermic reaction of the sorbent. All the experiments were performed isothermally at 900°C.

It is important to note that in the present work the focus is on the first calcination step due to the fact that the highest attrition rate occurs in the first cycle rather than in the following cycles. Therefore, this first calcination is considered to provide a good indication of sorbent attrition behaviour [41, 42].

2.4 Sample characterisation

The biomass-containing materials (LF and LCF) were analysed by TGA in a Netzsch STA409 CD apparatus in order to investigate the effect of biomass pyrolysis and combustion throughout calcination of the templated sorbents. Two different types of tests have been carried out, namely, pyrolysis-calcination and combustion-calcination tests.

In pyrolysis-calcination tests the sample was dried in argon at 110°C, and then the temperature was taken to 900°C in argon with a ramp rate of 5°C/min. At 900°C the
gas was switched from argon to air and the samples were held at isothermal conditions for 60 min, before being cooled at a rate of 20°C/min. In combustion-calcination tests the sample was exposed to air flow from the very beginning. It was dried at 110°C and then taken to 370°C with the heating rate of 10°C/min. An isothermal step of 10 min in air was performed at 370°C to allow combustion of biomass. Finally, the sample was heated to 900°C at a similar heating rate and kept at this temperature for 10 min to allow calcination. Approximately 30 mg of sample were used for each experiment, with a gas flow rate of 200 mL/min. In the case of LCF, additional tests were performed for the following particle size ranges: 710-500 μm and 250-500 μm in order to investigate the effect of particle size.

Sample morphology was observed with a FEI Inspect S Scanning Electron Microscope (SEM) with 20 kV of accelerating voltage under high vacuum. The calcined samples were put in the SEM chamber together with the ceramic pan. Before the analysis, the samples were coated with gold to avoid excessive charging. The porosimetry was studied using an AutoPore IV 9500 with mercury intrusion.

The particle size distribution was measured using the Mastersizer 2000 (Malvern, UK) and acetone was used as a carrier liquid. For X-ray diffraction analysis (XRD) a D2 Phaser (Bruker, Germany) apparatus with Cu Kα radiation (30 kV, 10 mA) was used. Scattered X-ray intensities were recorded between 2θ = 5 and 75° with a scan velocity of 0.052θ s⁻¹.

3 Results and discussion

3.1 Sorbent characterisation

Figures 1 and 2 report the mass loss profiles obtained during TGA experiments of pyrolysis-calcination and combustion-calcination.

In the case of LF a first stage of mass loss is observed between 350-450°C followed by a second stage of mass loss between 650-700°C, which can be associated with flour pyrolysis and sorbent calcination, respectively. Interestingly, the mass appears to be higher under pyrolysis conditions than under combustion, possibly due to the concurrent uptake of H₂O/CO₂ from air. In the combustion-calcination test a more noticeable uptake of CO₂ is evident above 450°C. The release of CO₂ at still higher temperature supports the similar value of value of 60%, which is observed for the final residue after both pyrolysis-calcination and combustion-calcination of the flour templated samples.

In the case of LCF two stages of mass loss are observed associated with flour pyrolysis and sorbent calcination. The effect of CO₂ uptake in the combustion-calcination test is less striking for LCF than for LF, due to the smaller percentage of lime in LCF.
The fact that the final residue of the combustion-calcination tests exceeds that of the pyrolysis-calcination tests by 5% suggests that some structural changes in the cement phase might have occurred in air.

Figure 1: TG (%) and temperature of tests in air and argon of LF

Figure 2: TG (%) and temperature of tests in air and argon of LCF

Figures 3, 4 and 5 show typical SEM images of LC, LF and LCF, respectively, after calcination. There are clear differences in structure: it is evident that the LC has a more compact structure, whilst LF has a more porous surface, as can be seen in Figure 4, due to the addition of flour, which creates mesopores in the structure. LCF displays a mixture of both structures, and it is definitely denser than LF; however, it has smaller pores and a more porous structure than LC.
Figure 3: SEM images of calcined LC at 20 kV and different magnifications
In Figure 6 the mercury intrusion pore volume (dV/dD) is shown. From these data, it can be inferred that there is a difference in the pore size distribution. In LC, there is a greater amount of larger pores (around 100 nm) than in LF and LCF and this was
expected due to the addition of biomass; this addition creates smaller pores, as has
been mentioned above. However, it can be seen that the total pore surface area is
lower in the sample with biomass-only templating (LF) with a total pore area of 13.2
m²/g compared to 15.0 m²/g (LC). Further, this seems to be mitigated when adding
cement, as the pore area is increased to 17.12 m²/g (LCF). This increase in area is
believed to be related to the mesoporous Al₂O₃ phase formed by the addition of the
calcium aluminate cement in the pelletisation process [43].

Figure 6: dV/dD pore volume vs pore diameter for calcined LC, LF and LCF

XRD analysis of the samples was also carried out for LC, LF and LCF. The results of
this analysis can be seen in Figure 7. The differences in composition of the samples
are mainly in the mayenite (Ca₁₂Al₁₄O₃₃) phase that forms from the reaction between
calcium aluminate cement and lime in the production process. These outcomes have
been well documented in earlier investigations [44, 45].
3.2 Fragmentation tests

3.2.1 PHSR tests

The PHSR tests were performed repeatedly with raw samples until enough material was collected to analyse particle size distribution (PSD). In Figure 8, the PSD for LC is shown, comparing the raw material with the material after injection into the reactor, simulating the conditions at the entrance of the calciner. It can be seen that the PSDs are similar. However, it should be noted that the fine particles present in the raw sample probably became finer still, and could not be recovered after the PHSR experiment. Nonetheless, the mean diameter is almost equal in both the raw material and the treated sample.
Figure 8: Particle size distribution of LC before and after the PHSR tests

The same distributions are shown for LF and LCF in Figures 9 and 10, respectively. For LF the difference in the particle size distribution before and after the fragmentation experiment is quite significant, with major fragmentation occurring when the particles are treated at 950°C, and the mean diameter decreases substantially from 520 µm to 116 µm. However, for LCF the change is less pronounced, although there is some fragmentation occurring when compared to LC; the change in mean diameter in LCF is smaller than for LF with a decrease from 524 µm to 290 µm. From these results, it can be inferred that the addition of biomass has a negative effect on the mechanical strength of the particles, making them more prone to fragmentation in the early stages of calcination presumably due to thermal stresses they experience. Nevertheless, the introduction of cement seems to have a positive effect in the biomass-templated particle with respect to fragmentation.

Figure 9: Particle size distribution of LF before and after the PHSR tests
3.2.2 BFB experiments

Figures 11-14 report the results of fragmentation tests in the fluidised bed reactor. Figure 11 reports the PSDs after treating the samples of LC, LF and LCF of original size 500-710 µm in the fluidised bed reactor in air. There is a clear difference in the behaviour among samples; LC undergoes less fragmentation, with 82% of the samples retaining the initial size range, as compared to LF, with 33% of the sample in the initial size range and LCF, with 47% of the sample in the initial size range. However, as noted in the PHSR experiments, cement addition has a positive effect making the particles less susceptible to fragmentation than the biomass-only templated material (LF).
Figure 12: Weight distribution percentage of recovered material (500-710 µm, 900 °C in 70% vol CO₂, 30% vol air)

Figure 12 reports the weight distribution of the material after calcination of the same samples in 70% CO₂ with balance of air. It can be seen that the material that fragments the most is LF with only 30% of the particles remaining in the initial size range, followed by LCF with 43% and LC with 90%.

The results of the same tests carried out on samples of smaller particle size (250-500 µm), are reported in Figures 13-14. For calcination in air, LF has the highest fragmentation with only 67% of particles retaining the initial size, followed by LCF with 74% and LC with 94%. It can be seen that smaller particle sizes lead to less fragmentation of the material recovered in the reactor at the end of the tests.

Figure 13: Weight distribution percentage of recovered material (250-500 µm, 900 °C in air)

For the smaller size particles (250-500 µm) calcined in 70% CO₂ with balance of air, the results were qualitatively similar to the other results. Nevertheless, the particles that remained in the reactor underwent less fragmentation than LF with 62% of particles in the initial size range, as was expected. The amount of fines (53-125 µm
and 125-180 µm) is higher for LF (0.9%wt) and LCF (0.08%wt) and LC (0.04%wt),
due to increased fragmentation created by biomass addition.

Figure 14: Weight distribution percentage of recovered material (250-500 µm, 900 °C in 70% vol CO₂, 30% vol air)

The percentage of material loss in the course of the BFB experiments is reported in Table 2. It can be seen that around 45% weight of LF was lost in the first calcination for smaller particles and around 30% for larger particles. This loss would be unacceptable for any real system. For LCF the losses were less significant for larger particles (between 4-8%), whilst for the smaller particles LCF behaved very similar to LF. The material that performed best was LC with a loss of around 3% for the smaller sample and 30% for the larger sample. Interestingly, the gas composition used in the calcination process did not affect the results significantly.

It has been noted above that LCF and LF have similar behaviour for the smaller particle range; this could be due to the different composition in the larger LCF particles, which might incorporate more cement during the pelletisation process. This hypothesis was supported by comparing the results of pyrolysis-calcination TGA experiments carried out on LCF samples of different size cuts. The TG curves, reported in Figure 15 show a smaller water content for the larger particles which suggests that they contained more calcium aluminate cement than the smaller size range particles. However, it should be noted that the particle size effect observed in Table 2 could be related to elutriation. The contribution of elutriation to the loss of bed material is in fact more noticeable the smaller the particle size is. Another point worth reiterating is that the main difference between both size ranges is the amount of cement present in each size cut. The effect of adding cement has been extensively studied in previous papers. There is a negative impact of adding cement because the particles have less active (CaO) material in them so that would negative influence there CO₂ uptake, also cement reacts with lime to form mayenite (Ca₁₂Al₁₄O₃₃). However, a positive effect has also been found in which the addition of calcium aluminate cement stabilizes the structure of the particle due to a mesoporous Al₂O₃ phase that delays sintering and therefore decreases the reactivity decay over cycles [26, 44].
Table 2: Loss in sample weight during the BFB experiments

<table>
<thead>
<tr>
<th>Sample</th>
<th>Loss of mass in weight (%) for 100% vol air</th>
<th>Loss of mass in weight (%) for 70% vol CO₂ and 30% vol air</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>250-500 µm</td>
<td>500-710 µm</td>
</tr>
<tr>
<td>LC</td>
<td>31.3</td>
<td>3.3</td>
</tr>
<tr>
<td>LF</td>
<td>45.4</td>
<td>27.4</td>
</tr>
<tr>
<td>LCF</td>
<td>41.5</td>
<td>3.4</td>
</tr>
</tbody>
</table>

Figure 15: TG Results for LCF (250-500 µm) and LCF (500-710 µm) for the first calcination

4 Conclusions

Biomass-templated pellets for calcium looping appear to be a cheap and scalable alternative option to achieve high CO₂ uptake when compared with other synthetic materials for Ca looping. The reactivity of these templated materials has been previously investigated in both TGA and BFB. Here, LF experienced the highest fragmentation in PHSR tests with a reduction in mean diameter of 404 µm compared to 234 µm for LCF and no change for LC. Moreover, in BFB, LF displayed the worst performance with a mass loss as high as 45.4% wt for smaller particles in the air fluidisation case. The weight loss of this material was significantly higher than for LC with 31.3% wt for the same case; this suggests that addition of biomass has a detrimental effect with regards to fragmentation. This effect is partially counteracted by the addition of calcium aluminate cement, which augments the resistance to fragmentation for LCF. However, the composition of LCF varied with particle size with the smaller range (250-500 µm) showing more elutriation of fines due to the pelletisation process, in which the larger particle size appeared to have incorporated more cement, leaving the smaller particles with a less stable structure. In consequence, smaller LCF particles were more prone to elutiate than the larger ones (500-710 µm). However, the particles that stayed in the reactor for the duration of the test underwent less fragmentation compared to the larger size range. It is clear that LC is the best material as regards fragmentation in all the cases explored, with a loss in mass as low as 3.3% for 500-710 µm particles in air calcination. The results for both techniques, BFB and PHSR, agree on the effects that biomass-templating...
has on the fragmentation behaviour of these sorbents under calcium looping conditions, although the addition of cement partially mitigated this negative effect.

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