# A high temperature, lithium orthosilicate based solid absorbent for post combustion CO<sub>2</sub> capture

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#### ABSTRACT

A lithium orthosilicate (Li<sub>4</sub>SiO<sub>4</sub>) based solid absorbent supplied by Toshiba Corporation was found to be promising for post combustion CO<sub>2</sub> capture. The absorbent reacts chemically with CO<sub>2</sub> at elevated temperatures and very large capacities can be achieved. The presence of water vapor greatly enhanced CO<sub>2</sub> absorption rates without affecting capacity. Exposure to flue gas containing even low levels of SO<sub>2</sub> resulted in an irreversible reaction with the absorbent and a decrease in CO<sub>2</sub> capacity. Breakthrough experiments indicated capacities as high as 6.7 mmol CO<sub>2</sub>/g. The absorbent will likely require a thermal swing process with absorption at 550°C and regeneration at 650°C.

*Keywords*: carbon dioxide, post combustion capture, lithium orthosilicate, solid absorbent

#### **1 INTRODUCTION**

Capture of carbon dioxide from combustion processes presents a unique and challenging technical problem arising from low CO<sub>2</sub> partial pressures, high flow rates, and the presence of water vapor and reactive contaminants such as SO<sub>2</sub>. In addition to these technical challenges, the scale of post combustion capture is daunting. A 500 MW coal fired power plant burns enormous quantities of coal, about 2 rail cars/h, and emits over 4 billion kg CO<sub>2</sub> in one year.[1] Based on the Department of Energy's target of 90% capture, 3.6 billion kg CO<sub>2</sub> must be separated each year. The current technology of choice, aqueous monoethanolamine (MEA), can effectively capture CO<sub>2</sub> but regeneration requires substantial heat input, 63-84 kJ/mol CO<sub>2</sub>, resulting in plant inefficiencies and higher costs[1] even though recent efforts have achieved some cost and energy reductions.[2] Solid sorbents have been proposed as an alternative to aqueous MEA for post combustion CO<sub>2</sub> and numerous such materials have been evaluated.[3-5] We have identified a lithium orthosilicate (Li<sub>4</sub>SiO<sub>4</sub>) based absorbent supplied by Toshiba Corporation[6-10] as a promising candidate for post combustion capture.

#### 2 RESULTS AND DISCUSSION

The Li<sub>4</sub>SiO<sub>4</sub> based absorbent was provided by Toshiba Corporation as 2 and 5 mm diameter spheres and are referred to as LS 2 mm and LS 5 mm, respectively. X-ray diffraction (XRD) analysis indicated that both contained Li<sub>4</sub>SiO<sub>4</sub> as the major phase along with Li<sub>2</sub>SiO<sub>3</sub> and Li<sub>2</sub>TiO<sub>3</sub>. Elemental compositions as determined by x-ray fluorescence (XRF) are listed in Table 1 with the exception of lithium which is not detected by XRF. It is assumed that potassium was present as amorphous K<sub>2</sub>CO<sub>3</sub> and, thus, not observed by XRD. Absorption of CO<sub>2</sub> occurs by formation of lithium carbonate and lithium metasilicate as in reaction 1[6-9] with K<sub>2</sub>CO<sub>3</sub> functioning as a promoter.[10]

$$Li_4SiO_4 + CO_2(g) = Li_2CO_3 + Li_2SiO_3$$
(1)

XRD analysis of the absorbent exposed to  $CO_2$  confirmed the expected product,  $Li_2CO_3$ , along with  $Li_2SiO_3$  and  $Li_2TiO_3$ . The absence of a  $Li_4SiO_4$  phase implies near quantitative reaction with  $CO_2$ . Capacities at 650°C and 1 atm  $CO_2$  were 5.51 mmol/g for LS 5 mm and 6.47 mmol/g for LS 2 mm. The heat of absorption for LS 2 mm was 82.0 kJ/mol  $CO_2$  at 650°C as determined by differential scanning calorimetry.

Table 1. Elemental compositions as determined by XRF.

	concentration, weight %			
absorbent	Si	Ti	Κ	Na
LS 5 mm	33.66	14.28	3.15	-
LS 2 mm	33.77	12.92	4.38	0.491

Utilization of a solid absorbent for post combustion  $CO_2$  capture requires knowledge of  $CO_2$  uptake as a function of temperature. A high operating temperature is preferable so as to maximize absorption rates but lower temperatures favor absorption. An evaluation using a synthetic flue gas containing 15%  $CO_2$  showed that the maximum absorption temperatures for 90% capture were 585°C for LS 5 mm and 571°C for LS 2 mm and 550°C was chosen as the absorption temperature. Thermogravimetric analysis (TGA) was used to determine absorption properties for a

feed gas containing approximately ~15% CO<sub>2</sub> in N<sub>2</sub> at 1 atm total pressure. Although CO<sub>2</sub> was absorbed to its equilibrium capacity, absorption was discouragingly slow at 550°C. Quite surprisingly, however, the use of humidified gas mixtures led to a substantial enhancement of absorption rates. Figure 1 compares CO<sub>2</sub> absorption as a function of time for LS 5 mm exposed to 14.7% CO<sub>2</sub> in N<sub>2</sub>, either dry or containing ~2.6% water. The dramatic effect of water vapor is illustrated, for example, by comparing the quantity of CO<sub>2</sub> absorbed after 5 min at 550°C: dry gas, 1.05 mmol/g; humidified gas, 3.67 mmol/g. The enhanced absorption rate in the presence of water vapor does not result from water adsorption as shown by a 0.02 wt% increase upon exposure to humidified N<sub>2</sub> at 550°C.



Figure 1. LS 5 mm CO<sub>2</sub> absoprtion rates at 550°C. Feed: 14.7% CO<sub>2</sub> in N<sub>2</sub>, dry (dashed curve) or humidified to  $\sim$ 2.6% H<sub>2</sub>O (solid curve).

The cyclic stability of the absorbent was demonstrated by a TGA experiment in which LS 5 mm was repeatedly exposed to humidified 14.7% CO2 in N2 for 10 min followed by a 30 min dry N<sub>2</sub> purge. The 1<sup>st</sup> cycle capacity of 4.12 mmol/g was lower than the equilibrium capacity due to the limited exposure time. A modest capacity decline to 3.81 mmol/g after 5 cycles was obtained but this was followed by a gradual increase in capacity and a final capacity of 4.31 mmol/g after 45 cycles. This is unusual compared to other high temperature solid sorbents that typically undergo a decreasing in capacity with cycling. Solid sorbents often undergo a decrease in crush strength with repeated cycling but this was not the case for the Li<sub>4</sub>SiO<sub>4</sub>-based absorbents. Also unusual was the higher crush strength of 45.1 lb after 50 cycles versus 16.2 lb before cycling.

A major concern for any post combustion sorbent is the effect of flue gas contaminants, particularly SO<sub>2</sub>, on its performance.[11-13] The reactivity of SO<sub>2</sub> was investigated utilizing a Rubotherm magnetic suspension balance for which the sample was completely isolated from the instrument electronics permitting the use of reactive or corrosive gases to high pressures and temperatures.[13] Exposure of the LS 5 mm absorbent to 0.002, 0.03 or 0.95 vol.% SO<sub>2</sub> in N<sub>2</sub> for 6 hours at 550°C and 1 atm resulted in the weight changes shown in Figure 2. Exposure to SO<sub>2</sub> concentrations as low as 0.002% resulted in a weight

increase. Higher SO<sub>2</sub> concentrations resulted in faster absorption and a 40 wt% increase was obtained for a 0.95% SO<sub>2</sub> containing gas mixture. Purging the SO<sub>2</sub>-loaded absorbent with 15%  $H_2O/N_2$  resulted in no weight change consistent with an irreversible reaction.

The effects of SO<sub>2</sub> on CO<sub>2</sub> absorption were determined by cycling between an SO<sub>2</sub> containing feed gas and an inert purge gas (25% H<sub>2</sub>O in N<sub>2</sub>), both at 1 atm and 550°C. In each experiment, a baseline cyclic capacity was established using a SO<sub>2</sub>-free feed for 10 cycles followed by repeated cycling using a SO<sub>2</sub>-containing feed (Figure 3). Both SO<sub>2</sub> and  $CO_2$  were absorbed during the feed step but only  $CO_2$ desorbed under the inert purge. Absorption of SO<sub>2</sub> was irreversible. The weight changes in Figure 3 reflect those arising from  $CO_2$  absorption alone. A feed containing 0.002% SO<sub>2</sub> led to a very modest decrease in the CO<sub>2</sub> capacity from 22.8 to 21.4 wt% over 25 cycles. Much more dramatic changes were observed for higher SO<sub>2</sub> concentrations: 0.03 vol.% SO<sub>2</sub>, capacity 16.2 wt% after 20 cycles; 0.95 vol.% SO<sub>2</sub>, capacity 1.4 wt% after 12 cycles.



Figure 2. SO2 uptake for various concentrations of SO2 in  $N_2$  at 550°C and 1 atm.



Figure 3. Weight change upon exposure to 15% CO<sub>2</sub>, 10% H<sub>2</sub>O in N<sub>2</sub>, cycles 1-10, followed by 15% CO<sub>2</sub>, 10% H<sub>2</sub>O, in N<sub>2</sub> containing 0.002% SO<sub>2</sub> ( $\blacktriangle$ ), 0.03% SO<sub>2</sub> ( $\blacklozenge$ ), or 0.95% SO<sub>2</sub> ( $\blacklozenge$ ).

XRF analysis of absorbent samples after SO<sub>2</sub> exposure confirmed the presence of sulfur with loadings of 1.43, 4.84, and 12.15 wt% S for the 0.002, 0.03, and 0.95% SO<sub>2</sub> containing feeds, respectively. XRD analysis showed that SO<sub>2</sub> absorption occurred by formation of a sulfate

containing product. For the sample exposed to 0.95% SO<sub>2</sub>, Li<sub>2</sub>SO<sub>4</sub>, KLiSO<sub>4</sub>, and K<sub>2</sub>SO<sub>4</sub> phases were found along with the inert phases Li<sub>2</sub>SiO<sub>3</sub> and Li<sub>2</sub>TiO<sub>3</sub>. The absence of Li<sub>4</sub>SiO<sub>4</sub> implies that near quantitative reaction with SO<sub>2</sub> had occurred. Energy dispersive spectroscopy of the cross-sectioned SO<sub>2</sub> exposed pellets revealed the expected elements: O, Si, K, Ti, S (Li is not detected). Although there were slight gradients from the outer edge to the center, sulfur was found throughout the pellets, even at the lowest loading. This means that virtually all the Li<sub>4</sub>SiO<sub>4</sub> was available for contacting with gaseous SO<sub>2</sub>.

The formation of  $\text{Li}_2\text{SO}_4$  as the major product requires the oxidation of SO<sub>2</sub>. Excluding oxidation via exposure to air during sample handling, it is reasonable that CO<sub>2</sub> and/or H<sub>2</sub>O function as oxidizing agents at 550°C. If oxidation to Li<sub>2</sub>SO<sub>4</sub> occurs in the absence of O<sub>2</sub>, it will very likely occur in the presence of an actual flue gas containing ~3% O<sub>2</sub>. The resulting formation of Li<sub>2</sub>SO<sub>4</sub> precludes any reasonable process for regeneration of the SO<sub>2</sub> exposed absorbent. In this regard, the Li<sub>4</sub>SiO<sub>4</sub> based absorbent is no different than other materials that chemically absorb CO<sub>2</sub>. All such absorbents, including aqueous MEA, will have higher affinities for SO<sub>2</sub> than CO<sub>2</sub> and desulfurization of flue gas prior to contacting the absorbent will be required or spent material will need to be replaced.

Breakthrough capacities were determined using a feed gas consisting of 14.7% CO2 in N2, either dry or containing water vapor, generally 10 vol%. Typical feed flows were 40-50 sccm with contact times generally near 4-6 sec. Because of the relatively small vessel diameter, the LS 5 mm spheres were crushed and sieved to 25-35 mesh prior to testing while LS 2 mm was used as intact 2 mm spheres. Figure 4 compares breakthrough curves at 550°C for LS 5 mm obtained with a dry gas feed and one containing 10% water vapor. Capacities at 100% breakthrough were about 4 times greater in the presence of water than in its absence, 4.18 mmol/g versus 0.90 mmol/g, respectively. The LS 2 mm absorbent exhibited larger breakthrough capacities with a similar enhancement of capacity in the presence of water vapor, 4.45 mmol/g for a dry feed and 6.77 mmol/g for a feed containing 10% water. The higher capacities in the presence of water vapor are a consequence of the enhance CO<sub>2</sub> absorption rates as demonstrated by the TGA results described above. Breakthrough capacities as a function of temperature were evaluated between 350 and 650°C for LS 5 mm and a maximum capacity was achieved near 550°C.

Critical to the successful implementation of any sorbent for  $CO_2$  capture is its regeneration properties. Two modes of regeneration can be considered. One is a pressure swing or, more accurately, a concentration swing process in which regeneration is carried out at the same temperature as sorption by purging with steam to reduce the  $CO_2$  partial pressure above the sorbent. The  $CO_2$  product is subsequently obtained by condensing liquid water. The concentration swing process has the advantage that no heating or cooling is required and cycle times can be short (<hours). The second approach is a temperature swing

process in which the sorbent is heated above the sorption temperature resulting in  $CO_2$  desorption. This can be coupled with a steam purge. A temperature swing process has the disadvantage that the sorbent must be heated and cooled resulting in long cycle times (hours) and, thus, larger vessels and sorbent inventory are required. For both LS 2 mm and LS 5 mm, a temperature swing process with absorption at 550°C and regeneration at 650°C was sufficient to maintain capacity as demonstrated by largely constant capacities over three cycles. A concentration swing regeneration, however, in which the absorbents were purged with 45% H<sub>2</sub>O in N<sub>2</sub> at 550°C for 2 h resulted in decreased capacities over three cycles. This implies that regeneration was incomplete and a temperature swing process will likely be required. As shown in Figure 5, desorption at 650°C was fast and much slower at 550°C.



Figure 4. Breakthrough curves for LS 5mm at 500 °C. Feed: 14.7% CO<sub>2</sub>, dry or humidified to 10% H<sub>2</sub>O as indicated.



Figure 5.  $CO_2$  concentration (dry basis) in reactor exit for regeneration of LS 2 mm at 550°C (dashed line) and 650°C (solid line); 700 sccm 45%  $H_2O/N_2$  purge gas.

The scale of post combustion capture using the  $Li_4SiO_4$ based absorbent in a fixed bed process was calculated based on the most optimistic assumptions. Considered was a 500 MW coal fired power plant with 90% CO<sub>2</sub> capture corresponding to about 419,000 kg CO<sub>2</sub>/h.[1] It was assumed that the equilibrium breakthrough capacity could be achieved and this value was used to calculate the minimum quantity of absorbent required. A temperature swing process was used with a total cycle time of 8 h, 4 h each for absorption and regeneration. A 4.27 m diameter by 6.10 m high vessel was used (volume 87.2 m<sup>3</sup>) and the number of vessels needed was taken to be twice that required for absorption. Based on this "best case" analysis, the scale of capture is immense; greater than 200 vessels and 5,600,000 kg of absorbent would be required.

It is clear that a fixed bed process using a sorbent that reacts chemically with CO<sub>2</sub> is unlikely to be practical for post combustion capture. This does not mean that solid sorbents cannot be applied to post combustion capture, rather an alternative to a conventional temperature swing fixed bed process is required. For example, if the  $Li_4SiO_4$ based absorbent could be used with a cycle time of 30 min rather than 8 h, the number of vessels and quantity of absorbent become much more reasonable, 791,000 kg and 14 vessels, based on the same optimistic assumptions. A process that heats and cools vessels and contents rapidly could result in shorter cycle times, thus minimizing absorbent inventory and numbers of vessels. Briefly, three such alternatives are: (1) a "rapid thermal swing chemisorption" (RTSC) process that utilizes shell and tube type vessels to minimize heating and cooling times [14] (2) a moving bed process in which the sorbent is circulated between a sorption vessel and a regeneration vessel[15] and (3) an "adsorbent wheel" in which the sorbent is contained within a rotating wheel or disc that rotated through absorption and desorption zones.[16]

## **3 EXPERIMENTAL**

The  $Li_4SiO_4$  based absorbents were supplied by Toshiba Corporation as 5 mm or 2 mm diameter spheres. For use in breakthrough experiments, the 5 mm spheres were crushed and sieved into 25-35 mesh particles under an inert atmosphere.

Thermogravimetric analysis was performed using a TA Instruments TGA Q5000 instrument contained within a N<sub>2</sub> purge box to minimize exposure to atmospheric water or oxygen. Humidified gases were obtained by passage through a water bubbler at room temperature. Water concentrations were not measured and were assumed to be equal to the water vapor pressure at room temperature, 20 torr or 2.6% at ambient pressure. Heats of absorption were obtained using a TA Instrument 2960 SDT V3.0F. XRF and XRD analysis were performed as described elsewhere.[13] Sulfur dioxide experiments were performed using a Rubotherm magnetic suspension balance at 550°C and 1 atm. Details of the balance and its operation are described elsewhere.[13] In cyclic exposure experiments, the absorbent was first exposed to 10 cycles of absorption and regneration using an SO<sub>2</sub>-free feed gas (15% CO<sub>2</sub>/10%  $H_2O/N_2$ ) for 1 h followed by purging with 25 vol.%  $H_2O$  in  $N_2$  for 3 hours. An additional 25 cycles using the above feed containing 0.002 to 0.95% SO<sub>2</sub> were then performed.

Breakthrough experiments were performed using a stainless steel tube measuring 0.37 in internal diameter and 7.25 in long. Following an initial purge with  $N_2$ , the absorbent was exposed to 14.7% CO<sub>2</sub> in  $N_2$  generally humidified by passage through a water bubbler at 46°C.

Gases exiting the absorbent vessel were passed through a chilled trap to condense water vapor prior to analysis using a Horiba  $CO_2$  analyzer. Breakthrough capacities were obtained by mass balance as the difference between the total quantity of  $CO_2$  in the inlet and effluent gas. Regeneration was performed by purging with ~45%  $H_2O/55\%$  N<sub>2</sub> stream.

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