Carbon capture system and method for capturing carbon dioxide

A carbon capture system comprising a Carbonator (2) for adsorbing carbon dioxide with a carbon dioxide lean sorbent generating a carbon dioxide rich sorbent, a first Calciner (1) for thermally decomposing a carbon dioxide rich sorbent into a carbon dioxide lean sorbent and carbon dioxide, a supply of raw material (4) to be calcined into the first Calciner (1) containing a carbon dioxide rich sorbent, a connection (13) between the first Calciner (1) and the Carbonator (2), a second Calciner (15) for thermally decomposing a carbon dioxide rich sorbent into a carbon dioxide lean sorbent and carbon dioxide, a connection (16) between the Carbonator (2) and the second Calciner (15), a connection (17) between the second Calciner (15) and the Carbonator (2).
Description

TECHNICAL FIELD

[0001] The present invention relates to a carbon capture system and method for capturing carbon dioxide.
[0002] Several systems and methods have been described previously which detail the capture of CO₂ from flue gas. These systems and methods can be integrated into different applications, such as into an industrial application or into a power plant to capture carbon dioxide from flue gas. Due to the rapid deactivation of natural sorbent materials, such as limestone, a preferred and particularly advantageous application of the present system and method are in connection with the production of low carbon content feed material for downstream cement clinker production, such as in a rotary kiln. While in the following description particular advantage of the proposed scheme for cement production is made, it is anyhow clear that the application of the system and method are not limited to cement production.

BACKGROUND

[0003] Cement production occurs by clinkering a raw material typically including limestone CaCO₃ (as a source of calcium oxide CaO), and clay, marl or shale (as source of SiO₂, Al₂O₃, Fe₂O₃) and typically also other materials such as sand.
[0004] A coupled carbon capture and cement production occurs over the following steps:

1) Sorbent preparation; material milling and heat-treatment, such as the calcination of limestone to produce CaO (allowing capture of the chemically associated CO₂ during calcination in the Calciner reactor);
2) Exposure of the sorbent powder to flue gas (possibly generated for the above mentioned heat treatment but also possibly from an external process) in order to convert CaO to CaCO₃ by lowering the concentration of CO₂ in the combustion flue gas through gas solid contact in the Carbonator reactor;
3) Regeneration of the sorbent materials, in particular the restoration of the CO₂ capture capacity (quantity CO₂ / quantity sorbent);
4) A sorbent purge to close the material balance of the cycle and remove deactivated sorbent from the system which is utilized for cement production.

[0005] Considering the above steps, Figure 1 shows a simplified block flow diagram representing a state of the art process scheme for cement production coupled with a carbon dioxide capture process. It includes a Calciner 1, a Carbonator 2 and a kiln 3. Raw material containing limestone CaCO₃ is supplied from a raw material supply 4 into the Calciner 1 where the make-up sorbent (mainly CaCO₃) is calcined (decomposed by heat) along with CaCO₃ from the Carbonator according to the reaction

\[ \text{CaCO}_3 \rightarrow \text{CaO} + \text{CO}_2 \]

generating carbon dioxide CO₂ and carbon dioxide lean sorbent CaO. The lean sorbent CaO is supplied to the Carbonator 2 where carbon dioxide CO₂ is captured from flue gas 5 according to the carbonation reaction

\[ \text{CaO} + \text{CO}_2 \rightarrow \text{CaCO}_3 \]

[0006] Gas 6 deprived from carbon dioxide CO₂ is discharged from the Carbonator 2. The calcium carbonate CaCO₃ generated at the Carbonator 2 is supplied back into the Calciner 1 allowing the release of captured Carbon dioxide and sorbent regeneration.
[0007] The carbon containing flue gas 5 may be attributed to a variety of thermally driven processes, such as power generation but a quantity of the flue gas will always be attributed to the cement process. At the kiln 3 flue gas is generated by combustion of a fuel with air; in addition carbon dioxide CO₂ is released during the clinkering reactions due to the residual carbon content of the feed material leaving the Carbonator 2 on route to the Kiln 3.
[0008] Figure 6 shows that after some adsorption/desorption cycles, the calcium oxide CaO loses its capacity to adsorb carbon dioxide CO₂, for this reason it must be purged.
[0009] According to the scheme of figure 1, purging is made by discharging calcined raw material containing calcium carbonate CaCO₃ from the Carbonator 2 and supplying it to the kiln 3 for calcination and thus clinkering.
[0010] As mentioned this scheme has the drawback that calcium carbonate CaCO₃ formed in the Carbonator is calcined at the kiln 3 which requires additional fuel combustion and produces additional CO₂ associated with the chemically bound CO₂ captured from the flue gas at the Carbonator 2. This causes an unnecessary carbon dioxide circulation between the Kiln and the Carbonator and consequently increased energy consumption and equipment dimensions and costs.
Figure 2 shows a second simplified block flow diagram representing a state of the art process scheme for cement production. It is similar to the scheme of figure 1 and same numbers indicates same or similar components. The scheme of figure 2 differs from the scheme of figure 1 in that purging is made by discharging raw material containing calcium oxide CaO from the Calciner 1 and supplying it to the kiln 3.

This second scheme reduces the amount of circulating carbon dioxide, because a reduced amount of carbon dioxide is released at the kiln (because additional calcination of CaCO₃ associated with carbon dioxide captured from the flue gas is avoided); nevertheless fresh and highly active calcium oxide CaO (sorbent) that can be most efficiently used for carbon capture is purged unselectively together with deactivated or spent sorbent (i.e. sorbent that has a reduced capacity to adsorb carbon dioxide).

SUMMARY

An aspect of the invention includes providing a system and a method that improve sorbent activity by avoiding or at least reducing unselectively purging of fresh and highly active sorbent and at the same time limit the circulation of carbon dioxide.

These and further aspects are attained by providing a system and a method in accordance with the accompanying claims.

BRIEF DESCRIPTION OF THE DRAWINGS

Further characteristics and advantages will be more apparent from the description of a preferred but non-exclusive embodiment of the system and method, illustrated by way of non-limiting example in the accompanying drawings, in which:

Figures 1 and 2 show examples of state of the art process schemes for Carbon capture coupled with cement production;
Figure 3 shows a simplified block flow diagram representing a process scheme for carbon capture coupled with cement production according to the invention;
Figure 4 shows a more detailed block flow diagram of an integrated carbon capture solution for cement production;
Figure 5 is diagram showing the fraction of sorbent in the system having completed a given number of calcination and recarbonation cycles. Figure 5 represents the situation for the plant schemes of figure 2 (curve A) and figure 3 (curve B) considering the material flows required to decarbonize cement production using the above mentioned methods;
Figure 6 is a diagram showing the reduction of sorbent activity with an increasing number of calcination and recarbonation cycles.

DETAILED DESCRIPTION OF EXEMPLARY EMBODIMENTS

With reference to figure 3, it shows a carbon capture system comprising a first Calciner 1 and a Carbonator 2. The Carbonator 2 is for adsorbing carbon dioxide from flue gas by employing a lean sorbent in order to generate a more rich sorbent. The composition of the lean and rich sorbent is dependent on system design and system operational parameters (such as the ratio of make-up flow to recirculation flow) in addition to the deactivation characteristic of the material. For deactivated rich sorbent the mole fraction of CO₂ bound to available Ca in the particle is typically less than 0.2 (corresponding to between 10 and 20 sorbent cycles in figure 8).

In the following reference to CaO as the carbon dioxide lean sorbent and CaCO₃ as the carbon dioxide rich sorbent is made; other lean/rich sorbent can anyway be used. In addition it is clear that lean and rich solvent are not made of only CaO and CaCO₃ but they are mixtures including mainly CaO or CaCO₃ and other materials; for example the lean sorbent has CaO as the main component, but may also contain CaCO₃ and similarly the rich sorbent may also contain CaO.

The first Calciner 1 is for desorbing the carbon dioxide rich sorbent CaCO₃ from the carbon dioxide CO₂ generating the carbon dioxide lean sorbent CaO.

The first Calciner 1 has a supply 4 of raw material to be calcined and a connection 13 between the first Calciner 1 and the Carbonator 2 for transferring the calcined raw material including the carbon dioxide lean sorbent CaO to the Carbonator 2.

In addition, the system has a second Calciner 15 for desorbing a carbon dioxide rich sorbent CaCO₃ from carbon dioxide CO₂ generating a carbon dioxide lean sorbent CaO, and connections 16 and 17 between the Carbonator 2 and the second Calciner 15.
The connection 16 between the Carbonator 2 and the second Calciner 15 is for transferring the carbon dioxide rich sorbent CaCO₃ to the second Calciner 15, and the connection 17 between the second Calciner 15 and the Carbonator 2 is for transferring carbon dioxide lean sorbent CaO back to the Carbonator 2 for renewed capture of carbon dioxide CO₂.

The system can be used in a number of different applications, for this reason the Carbonator 2 can be supplied with flue gas containing carbon dioxide CO₂ to be captured and separated from the flue gas.

Preferably, the system is used in a cement production plant; for this reason there are also provided a kiln 3 for cement production and a connection 18 between the second Calciner 15 and the kiln 3 for transferring carbon dioxide lean sorbent CaO to the kiln 3.

Application of the system in a plant for cement production is particularly advantageous because in this case the amount of make-up CaCO₃ (supplied with fresh raw material) is very high compared to the amount of purged CaO (forwarded to the kiln). This can be shown with reference to the ratio of make-up molar sorbent flow to the molar sorbent circulation rate, defined by

\[
\text{Ratio} = \frac{\text{Moles of CaO obtained by calcining the make-up sorbent (mainly CaCO₃)}}{\text{Moles of sorbent (mostly CaO) required to capture CO₂ from a given flue gas}}.
\]

The ratio is typically above 1 for the schemes described in Figure 1-3, and is well below 1 typically less than 0.05 for post combustion carbon dioxide capture (for example in power plants or industrial applications). Anyhow, the system provides benefits also for applications to industrially generated flue gas or flue gas from power generation.

Figures 4 shows the system more in detail; this system can for example be applied as a carbon capture retrofit solution for an existing cement plant; when describing this figure reference to CaO and CaCO₃ is made as one of the main components used in cement production and as the preferred sorbent. It is clear that in applications different from cement production the sorbent can be any appropriate sorbent known in the art of natural or synthetic origin.

Figure 4 shows a raw material supply 4 (for e.g. raw meal containing CaCO₃) connected to one or more preheaters 21, 22; for example the preheater 21 represents a number of preheaters (for example four or five preheaters from an existing cement plant) for the raw material to be calcined. The preheater 21 heats the raw material by cooling a mixture of carbon dioxide rich flue gas originating from the kiln 3 via a line 23 and carbon dioxide lean flue gas from the Carbonator 2 via a line 24. For example the kiln gas is bypassed to the preheater 21 of the existing plant to ensure the required operating pressure (under atmospheric pressure, vacuum conditions).

The preheater 22 heats the raw material by cooling the carbon dioxide discharged from the first and second Calciners 1 and 15 via the lines 29, 30.

The preheater 22 is connected to the first Calciner 1 for supplying the raw material (preheated raw material).

Before the first Calciner 1 and the Carbonator 2, a solid/solid heat exchanger 31 is provided, for heating the carbon dioxide rich sorbent CaCO₃ moving from the Carbonator 2 to the second Calciner 15 by cooling the carbon dioxide lean sorbent CaO moving from the first Calciner 1 and/or the second Calciner 15 to the Carbonator 2.

The second Calciner 15 is connected with the kiln 3 via the connection 32 and with the Carbonator 2 via the connection 33.

In addition, preferably a separator 34 is provided connected on the line 23, between the kiln 3 and the preheater 21; the separator 34 separates the solid particles possibly contained in the flue gas from the kiln 3 and forwards them back to the kiln 3 via a line 52.

In order to heat the first and second Calciners 1 and 15, a furnace 35 (for example air fired furnace) is provided; the furnace 35 heats solids that are circulated via connection lines 36 between the furnace 35 and the first Calciner 1 and via connection lines 38 between the furnace 35 and the second Calciner 15. Naturally also different schemes are possible for heating the Calciners 1 and 15. For example each Calciner 1, 15 can have a furnace or indirect heat transfer schemes without solids circulation, or direct fired (oxy-fuel) calcination can be implemented.

A connection 39 is provided between the furnace 35 and the Carbonator 2, for supplying flue gas generated in the furnace 35 by combusting a fuel into the Carbonator 2. Preferably, a heat exchanger 40 is provided on the connection 39, for preheating the air (or oxidizer) supplied into the furnace 35 via a line 42 by cooling flue gas discharged from the furnace 35 via the line 39; an additional line 43 can be provided to supply preheated air (or oxidizer, generated from cooling clinker exiting the kiln 3) to the furnace 35, bypassing the heat exchanger 40.

A heat exchanger 45 is also preferentially provided in addition to or in place of the heat exchanger 40, in order to preheat solids supplied via a line 46 to the furnace 35 by cooling the flue gas discharged from the furnace 35 via the line 39. The preheater 45 functions in a counter current manner heating the hot solids which enter at the top via line 46 and fall via gravity through the rising cooling flue gas from furnace 35. The line 46 can originate from the line 36 forwarding solids from the Calciner 1 into the furnace 35 and/or from the line 38 forwarding the solids from the calciner 15 into the furnace 35.

The kiln 3 is connected to a heat exchanger 48 for cooling the clinker 49 discharged from the kiln 3 by heating the fresh oxidizer such as air; the heat exchanger 48 is then connected via the connections 42, 43 to the furnace 35 for supplying the heated air (or oxidizer) into the furnace 35, and via a connection 50 to a fuel drying system, for drying solid fuel such as coal. The fuel drying system is then connected to the furnace 35 and kiln 3 in order to supply the required fuels.
The operation of the system is apparent from that described and illustrated and is substantially the following.

Raw material supply 4 supplies raw material to the preheaters 21 and 22 where it is heated and then supplied to the first Calciner 1. At the first Calciner 1 raw material is calcined (for example raw meal for cement production, it can be different for different applications). Since the raw material supplied to the first Calciner is fresh raw material additional time is required to calcine the sorbent. The Calciner 1 is designed accordingly so that the conversion CaCO₃ into CaO is almost complete (i.e. almost all CaCO₃ is converted into CaO). This conversion occurs according to the calcination reaction

\[ \text{CaCO}_3 \rightarrow \text{CaO} + \text{CO}_2 \]

usually at a temperature above 900°C, preferably about slightly above 900°C.

The Calciner 1 is designed so that the conversion CaCO₃ into CaO is almost complete (i.e. almost all CaCO₃ is converted into CaO). This conversion occurs according to the calcination reaction

\[ \text{CaCO}_3 \rightarrow \text{CaO} + \text{CO}_2 \]

usually at a temperature above 900°C, preferably about slightly above 900°C.

The CaO is supplied to the Carbonator 2 where it adsorbs carbon dioxide from the flue gas coming from the furnace 35 (and possibly from flue gas generated in an industrial application or power plant); adsorption takes place according to the reaction

\[ \text{CaO} + \text{CO}_2 \rightarrow \text{CaCO}_3 \]

usually at a temperature between about 600-750°C.

At the second Calciner 15 carbon dioxide is released by thermal decomposition according to the calcination reaction

\[ \text{CaCO}_3 \rightarrow \text{CaO} + \text{CO}_2 \]

usually at a temperature above 900°C, preferably about slightly above 900°C.

CaO is thus supplied to the kiln 3 via the line 32 and partly to the Carbonator 2 via the line 33.

Since at the Carbonator 2 there is a large amount of CaO coming from the first Calciner 1 (this is fresh CaO) and the amount of CaO from the second Calciner 15 is limited (because CaO is only purged from the second Calciner 15), the carbonation reaction at the Carbonator 2 is very efficient and the CaO can adsorb a large amount of carbon dioxide.

Figure 6 shows the cycles that the CaO has undergone in a system according to figure 2 (curve A) and according to figures 3 and 4 (curve B). This figure shows that according to the invention (figures 3 and 4) the CaO supplied to the Carbonator 2 undergoes less calcination / carbonation cycles and is therefore more reactive.

Considering the decline in the activity of the sorbent according to figure 6, the method of the invention provides an increased active fraction of circulating sorbent for the same sorbent molar flow (molar flow of CaO and CaCO₃). Considering the make-up rate of fresh sorbent (required for cement production), the method shown in figure 3 increases sorbent activity (mol CaCO₃/mol CaO and CaCO₃ in the sorbent) by between 5 to 20%, typically above 10%. Ultimately this translates into a lower required sorbent circulation rate or smaller equipment.

The furnace 35 combusts a fuel and generates hot solids that are circulated via lines 36 and 38 between the furnace 35 and the first and second Calculiners 1 and 15. Preferably a part of the cooled solids returning from the Calculiners (1 or 15) is diverted via the line 46 to be heated against flue gas discharged from the furnace 35 in the heat exchanger 45. The flue gas discharged from the furnace 35 is supplied to the Carbonator 2 (for carbon dioxide capture) after cooling via the heat exchangers 45 and 40.

The carbon dioxide rich flue gas from the kiln 3 is supplied to the separator 34 via the line 23, where solid particles are separated from the gas and forwarded back to the kiln 3 via the line 52.

The present invention also refers to a method for capturing carbon dioxide.

The method comprises supplying a raw material containing a carbon dioxide rich sorbent such as CaCO₃ (but other sorbents are possible, CaCO₃ is preferably used when the method is implemented in connection with a kiln for cement production) into a first Calciner 1. At the first Calciner 1, the raw material is calcined according to the calcination (decarbonation reaction)

\[ \text{CaCO}_3 \rightarrow \text{CaO} + \text{CO}_2 \]

that usually takes place at a temperature above 900°C, preferably about slightly above 900°C.

Following the calcination reaction, a calcined raw material containing a carbon dioxide lean sorbent CaO is generated; in addition, also carbon dioxide is generated.

The calcined raw material containing the carbon dioxide lean sorbent CaO is supplied to the Carbonator 2. In addition, also flue gas is supplied into the Carbonator 2; the flue gas comes from the furnace 35, but in other applications
can also come from an industrial process or a furnace of a power plant for electricity production.

[0054] At the Carbonator 2, carbon dioxide from flue gas is adsorbed by the carbon dioxide lean sorbent CaO according to the carbonation reaction

\[
\text{CaO} + \text{CO}_2 \rightarrow \text{CaCO}_3
\]

that takes place at a temperature between about 600-750°C.

[0055] Following the carbonation reaction, a raw material containing a carbon dioxide rich sorbent CaCO_3 is generated.

[0056] The raw material containing carbon dioxide rich sorbent, namely CaCO_3, is in turn supplied to the second Calciner 15.

[0057] At the second Calciner 15, the raw material containing carbon dioxide rich sorbent CaCO_3 is calcined according to the calcination (decarbonation reaction)

\[
\text{CaCO}_3 \rightarrow \text{CaO} + \text{CO}_2
\]

that usually takes place at a temperature above 900°C, preferably about slightly above 900°C.

[0058] Following the calcination reaction, calcined raw material containing carbon dioxide lean sorbent CaO and carbon dioxide are generated.

[0059] A part of the calcined raw material containing carbon dioxide lean sorbent CaO is supplied to the Carbonator 2.

[0060] When the method is used in connection with a kiln for cement production (preferred application) a part of the calcined raw material containing carbon dioxide lean sorbent CaO is transferred from the second Calciner 15 to the kiln 3 for cement production.

[0061] The additional Calciner 15 provides increased flexibility to consider increased calcination times ensuring complete calcination of make-up sorbent and ultimately increasing the sorbent activity (quantity CO_2 / quantity sorbent).

[0062] Naturally the features described may be independently provided from one another.

[0063] In practice the materials used and the dimensions can be chosen at will according to requirements and to the state of the art.

REFERENCE NUMBERS

[0064]

1 Calciner
2 Carbonator
3 kiln
4 raw material supply
5 flue gas
6 gas deprived from carbon dioxide
13 connection
15 second Calciner
16 connection
17 connection
18 connection
21 preheater
22 preheater
23 line
24 line
29 line
30 line
31 heat exchanger
32 connection
33 connection
34 separator
35 furnace
36 connection line
38 connection line
39 connection
40 heat exchanger
A carbon capture system comprising
a Carbonator (2) for adsorbing carbon dioxide with a carbon dioxide lean sorbent generating a carbon dioxide rich sorbent,
a first Calciner (1) for thermally decomposing a carbon dioxide rich sorbent into a carbon dioxide lean sorbent and carbon dioxide,
a supply of raw material (4) to be calcined into the first Calciner (1), the raw material containing a carbon dioxide rich sorbent,
a connection (13) between the first Calciner (1) and the Carbonator (2) for transferring at least the carbon dioxide lean sorbent to the Carbonator (2), characterised by further comprising
a second Calciner (15) for thermally decomposing a carbon dioxide rich sorbent into a carbon dioxide lean sorbent and carbon dioxide,
a connection (16) between the Carbonator (2) and the second Calciner (15) for transferring at least carbon dioxide rich sorbent to the second Calciner (15),
a connection (17) between the second Calciner (15) and the Carbonator (2) for transferring carbon dioxide lean sorbent (CaO) to the Carbonator (2).

2. A carbon capture system of claim 1, characterised by further comprising
a kiln (3) for cement production,
a connection (18) between the second Calciner (15) and the kiln (3) for transferring carbon dioxide lean sorbent to the Carbonator (2).

3. A carbon capture system of claim 1, characterised by further comprising a preheater (21, 22) for the raw material to be calcined.

4. A carbon capture system of claim 1, characterised by further comprising a solid/solid heat exchanger (31) for heating the carbon dioxide rich sorbent moving from the Carbonator (2) to the second Calciner (15) by cooling the carbon dioxide lean sorbent moving from the first Calciner (1) and the second Calciner (15) to the Carbonator (2).

5. A carbon capture system of claim 1, characterised by further comprising a furnace (35) for heating solids, and connection lines (36, 38) between the furnace (35) and the first and/or the second Calciner (1, 15), for circulating the solids between the first and second Calciner (1, 15) and the furnace (35).

6. A carbon capture system of claim 5, characterised by further comprising a connection (39) between the furnace (35) and the Carbonator (2), for supplying flue gas generated in the furnace (35) by combusting a fuel into the Carbonator (2).

7. A carbon capture system of claim 1, further comprising a heat exchanger (48) for cooling clinker discharged from the kiln (3) by heating fresh oxidizer, and at least a connection (42, 43) of the heat exchanger (48) to at least one of:
   - the furnace (35), for supplying the heated oxidizer into the furnace (35),
   - a fuel drying system.

8. A carbon capture system of claim 7, wherein at least one of the at least a connection (42) to the furnace (35) is provided with a heat exchanger (40) for preheating the oxidizer supplied into the furnace (35) by cooling flue gas
discharged from the furnace (35).

9. A method for capturing carbon dioxide comprising supplying a raw material containing a carbon dioxide rich sorbent into a first Calciner (1), at the first Calciner (1), thermally decomposing the carbon dioxide rich sorbent into a carbon dioxide lean sorbent and carbon dioxide, supplying the calcined raw material containing the carbon dioxide lean sorbent to a Carbonator (2), at the Carbonator (2), adsorbing carbon dioxide from flue gas generating a raw material containing a carbon dioxide rich sorbent, **characterised by** supplying the raw material containing the carbon dioxide rich sorbent into a second Calciner (15), at the second Calciner (15), thermally decomposing the carbon dioxide rich sorbent into carbon dioxide lean sorbent and carbon dioxide, transferring at least a part of the calcined raw material containing carbon dioxide lean sorbent to the Carbonator (2).

10. The method of claim 9, **characterised in that** the carbon dioxide rich sorbent includes and the carbon dioxide lean sorbent includes CaO.

11. The method of claim 9, **characterised by** transferring at least a part of the calcined raw material containing carbon dioxide lean sorbent from the second Calciner (15) to a kiln (3) for cement production.
Fig. 4
Fraction vs. Cycle Number

Fig. 5

Typical Sorbent Activity vs. Cycle Number

Fig. 6
## Relevant Documents

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### Technical Fields Searched (IPC)

- B01D
- C04B

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The present search report has been drawn up for all claims

Place of search: Munich

Date of completion of the search: 8 December 2014

Examiner: Persichini, Carlo

### Category of Cited Documents

- **X**: particularly relevant if taken alone
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