The grain growth mechanism of nano-CaO regenerated by nano-CaCO₃ in calcium looping

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The grain growth mechanism of nano-CaO-based CO₂ adsorbents in calcium looping (CaL) process was studied to figure out the main factors affecting sorption durability. First, the thermal growth characteristics of nano-CaCO₃ grains and nano-CaO grains at 750–850 °C was measured to fit the grain growth kinetic models. The activation energy data of grain growth of nano-CaCO₃ and nano-CaO were obtained as 104.8 kJ mol⁻¹ and 212.8 kJ mol⁻¹ respectively, which indicated that the grain growth of nano-CaCO₃ was easier than that of nano-CaO. Then, the grain sizes of regenerated nano-CaO undergoing 10 CaL cycles were compared with those derived from nano-CaCO₃ suffering high temperature heat-treatment under the same looping temperature and time. It was found that CaCO₃–CaO chemical conversion could accelerate the grain growth of regenerated nano-CaO. Based on these results, the grain growth mechanism of regenerated nano-CaO grain in CaL process was proposed. The thermal growth of nano-CaCO₃ grain was the key issue to influence the grain growth of regenerated nano-CaO. Therefore, shortening the high temperature residence time as well as preventing the interface contact of nano-CaCO₃ grains were good for limiting the grain growth of regenerated nano-CaO.

1. Introduction

Using the reversible reaction of CaO-based adsorbents (as shown in eqn (1)) to capture greenhouse gas CO₂ at high temperature, termed CaL technology, is widely used in many fields such as zero-emission power generation technologies, and energy storage systems in chemical heat pumps. One of the bottleneck problems of CaL is that the sorption capacity of CaO-based adsorbents will always rapidly decay after several cycles, which is referred to as the “deactivation phenomenon”. It seriously affects the reversibility and economic feasibility of the CaL technology, so it is significant to study the deactivation mechanism of CaO-based adsorbents.

\[
\text{CaO} + \text{CO}_2 \rightleftharpoons \text{CaCO}_3, \Delta H_{298} = -178.8 \text{ kJ mol}^{-1}
\]  

(1)

The phenomenon of the growth of nano-CaO grain sizes can be often observed in CaL process, which means the sintering of CaO. Feng et al. found that the regenerated nano-CaO grain sizes in nano-CaO-based adsorbents increased from 37 nm to 62 nm after 6 CaL cycles. According to the research of Sun et al., the reason for deactivation of adsorbents was the sintering of CaO. It showed that the growth of nano-CaO grains led to the loss of specific surface area, which meant the reduction of the numbers of active CaO sites for adsorbing CO₂. Therefore, it is crucial to understand the grain growth mechanism of nano-CaO during CaL process. The growth of nano-CaO grains mainly occurs in the high temperature regeneration stage and the growth degree is controlled by regeneration temperature and time. Dash et al. found that the grain size of CaO would increase from 17 nm to 47 nm when the regeneration temperature was raised from 1000 °C to 1200 °C. Zhu et al. found that the CaO grain size would grow from 51.5 nm to 75.1 nm when the regeneration time was prolonged from 1 h to 4 h at 800 °C.

The grain size of regenerated nano-CaO is actually determined by the interaction of several structural changes including the thermal growth of CaO grains. The regeneration process of adsorbents can be roughly divided into three substeps: (1) thermal growth of nano-CaCO₃ grains, (2) CaCO₃ convert to CaO and (3) thermal growth of CaO grains. In recent researches, rapid growth of nano-CaCO₃ grain has been regarded as the fundamental reason for the growth of regenerated nano-CaO grains. Because the Tammann temperature of CaCO₃ (533 °C) is significantly lower than that of CaO (1154 °C), Müller et al. found that increasing the heating rate between sorption and regeneration steps to reduce the residence time of nano-CaCO₃ in the high temperature could slow down the deactivation trend of adsorbents. Nevertheless, the research on the grain growth mechanism of nano-CaO in CaL process is still incomplete. It is difficult to figure out which regeneration substep is the key issue to influence the grain growth of nano-CaO because of the coupling. Therefore, it is necessary to quantify and compare the impact of three regeneration substep on the grain growth of nano-CaO by designing decoupling experiments at the same regeneration conditions.

In this paper, the grain growth kinetic model of nano-CaCO₃ was fitted under the heat-treatment conditions (temperature...
750–850 °C, time 0–240 min) with the protective atmosphere of CO2. Also, the grain growth kinetic model of nano-CaO would be got at the same conditions with N2 protection. The dominant one for the increase of regenerated nano-CaO grain sizes could be found by comparing their activation energy for grain growth. Moreover, the grain sizes, pore structure and sorption performance of regenerated nano-CaO were compared separately through the CaL process and the heat-treatment looping (HtL) process without any reactions to investigate the effect of CaCO3–CaO conversion on the grain growth of regenerated nano-CaO. Finally, the grain growth mechanism of regenerated nano-CaO in CaL process was proposed.

2. Experimental
2.1 Heat-treatment of nano-CaCO3 and nano-CaO samples
Commercial nano-CaCO3 (>95% purity, Huzhou Ling Hua Ltd. China) and nano-CaO derived from commercial nano-CaCO3 decomposition were adopted. The schematic diagram of a laboratory-scale fixed bed reactor was shown as Fig. 1. The diameter of the stainless tube in the reactor was \( \Phi 12 \times 1 \) mm and the length was 400 mm. The temperature stable section was filled with 2 g nano-CaCO3 (or nano-CaO). The temperature was adjusted by a temperature controller. The feed flow rates of CO2 and N2 were regulated by the mass flow meters. First, the fixed bed was heated to a set heat-treatment temperature (750–850 °C) at a heating rate of 15 °C min\(^{-1}\), then held for a set period of time (60–240 min). When the nano-CaCO3 was heated, the reactor was filled with CO2 atmosphere to keep CaCO3 state. On the contrary, when the nano-CaO was heated, the reactor was filled with N2 atmosphere to keep CaO state. The heat-treatment conditions and the corresponding samples’ name were shown in Table 1. CaC-a/b and CaO-a/b (a = 0, 750, 800 and 850; b = 0, 60, 120, 240) referred to the nano-CaCO3 and nano-CaO sample under heat-treatment temperature of \( \alpha \) °C and heat-treatment time of \( \beta \) min, respectively. CaC-0/0 and CaO-0/0 referred to fresh nano-CaCO3 and fresh nano-CaO without heat-treatment.

2.2 CaL and HtL processes of nano-CaCO3
2 g commercial nano-CaCO3 was filled in the temperature stable section of the fixed bed reactor. The CaL process was operated at atmospheric pressure. First, heating the reactor to 800 °C and holding for 50 min in N2 gas flow (100 mL min\(^{-1}\)) to complete decomposition of CaCO3. Then cooling the reactor to 600 °C and maintaining for 25 min with CO2–N2 (20 mL min\(^{-1}\) CO2, 80 mL min\(^{-1}\) N2) atmosphere to complete carbonation of CaO. The regeneration–carbonation cycle was repeated to 10 times and the adsorbent samples were collected after carbonation period at different cycles. In contrast to the above CaL process, HtL process indicated that nano-CaCO3 was thermally cycled between 600 °C (25 min) and 800 °C (50 min) under the CO2 atmosphere protection without any reactions. HtL-N and CaL-N represented samples after the \( N \)th HtL cycle and the \( N \)th CaL cycle, respectively (\( N = 1, 5, 10 \)).

2.3 Characterization
In this paper, the X-ray diffraction pattern of adsorbents was tested by XRD (D/MAX-RA, Rigaku, Japan). The characteristic
peaks of $2\theta = 29.5^\circ$, $36.2^\circ$ and $39.5^\circ$; $2\theta = 53.9^\circ$, $64.2^\circ$ and $67.4^\circ$ were used to calculate the average grain size of nano-CaCO$_3$ and nano-CaO, respectively.$^{14}$ It could be determined by the Scherrer equation, as follows:

$$d = \frac{K\lambda}{\beta \cos \theta}$$  \hspace{1cm} (2)

The sorption performance of the adsorbent was measured using TGA (Pyris1, PerkinElmer, USA). The operation conditions of carbonation stage and regeneration stage were 600 °C–20% CO$_2$/80%N$_2$–20 min and 750 °C–100%N$_2$–10 min, respectively. The sorption capacity and sorption rate were calculated according to eqn (3) and (4):

$$\text{Sorption capacity} = \frac{\text{CO}_2 \text{ sorption mole amount}}{\text{adsorbent weight}} \left( \text{mol kg}^{-1} \right)$$ \hspace{1cm} (3)

$$\text{Sorption rate} = \frac{\text{sorption capacity}}{\text{time}} \left( \text{mol kg}^{-1} \text{ min}^{-1} \right)$$ \hspace{1cm} (4)

In addition, the specific surface area (Brunauer–Emmer–Teller, BET) and desorption average pore diameter (Barrett–Joyner–Halenda, BJH) analyses were conducted by nitrogen physisorption at $-196^\circ$C by means of a Micrometrics ASAP2020 apparatus. The morphology of the adsorbent was investigated by scanning electron microscopy (SEM, S-4800, Hitachi, Japan).

### 3. Results and discussion

#### 3.1 The grain growth characteristics of nano-CaCO$_3$ and nano-CaO

Prior to this research, the XRD of nano-CaCO$_3$ and nano-CaO samples with the most severe heat-treatment condition of 850 °C–240 min were tested to verify the feasibility of atmosphere protection, which was shown in Fig. 2. There were only their own characteristic peaks in the spectrum of nano-CaCO$_3$ and nano-CaO samples, indicating that the nano-CaCO$_3$ sample did not undergo thermal decomposition and the carbonation of the nano-CaO sample did not occur either after the heat-treatment. Therefore, CO$_2$ and N$_2$ could meet the requirements of inert atmosphere protection during heat-treatment and this exploration of grain growth process was feasible.

The grain sizes of nano-CaCO$_3$ and nano-CaO obtained under the different heat-treatment conditions were shown in Fig. 3 to compare their grain growth characteristics and fit their grain growth kinetic models. It could be found that within the range of examining conditions, as the heat-treatment temperature and the time increased, the grain sizes of nano-CaCO$_3$ and nano-CaO grew significantly. The nano-CaCO$_3$ grain size increased from initial 61.7 nm to more than 100 nm, and the nano-CaO grain size grew from initial 44.9 nm to about 70 nm. The growth rate of both kinds of grains would remarkably develop with the increase of heat-treatment temperature. However, it gradually decreased with the prolongation of heat-treatment time, which showed a parabolic trend. In general, the growth rate of the nano-CaCO$_3$ grain was greater than that of nano-CaO.

$$G^2 - G_0^2 = Ct$$ \hspace{1cm} (5)

![Fig. 2 XRD patterns of nano-CaCO$_3$ and nano-CaO after heat-treatment of 850°C-240 min.](image-url)
Table 2. The values of the grain size and pore structure of adsorbents during CaL and HtL process.

| Sample | Grain size (nm) | Specific surface area (m² g⁻¹) | Pore volume (cm³ g⁻¹) | Mean pore diameter (nm) |
|--------|----------------|-------------------------------|----------------------|------------------------|
|        | Experimental value | Calculated value |                |                        |                        |
| CaCO₃  | CaO             | CaO*                      |                     |                        |                        |
| HtL-1  | 61.0            | 46.9                      | 1.60                | 0.007                  | 20.9                   |
| HtL-5  | 74.2            | 57.1                      | 0.96                | 0.007                  | 57.5                   |
| HtL-10 | 95.1            | 73.2                      | 0.55                | 0.006                  | 83.4                   |
| CaL-1  | 36.7            | 64.3                      | 14.73               | 0.092                  | 27.6                   |
| CaL-5  | 29.0            | 76.3                      | 5.45                | 0.049                  | 37.1                   |
| CaL-10 | 26.5            | 81.9                      | 3.71                | 0.041                  | 38.3                   |

3.2 Regenerated nano-CaO grain growth during CaCO₃-CaO conversion

In this section, the effects of CaL and HtL processes on the grain size, pore structure and reactive sorption performance of nano-CaO adsorbents were compared to study the relationship between CaCO₃-CaO conversion and regenerated nano-CaO grain growth. The evolution of adsorbent's structural parameters during 10 cycles of CaL and HtL processes were shown in Table 2. Fig. 4 showed the composition change of the adsorbent during CaL process. There was only CaCO₃ component in the adsorbent during the HtL process. However, the adsorbent included “skeleton CaO” that failed to participate in the carbonation and CaCO₃ product generated by the carbonation of “active CaO” in the CaL process (Fig. 4). According to the grain sizes calculated by XRD patterns and Scherrer equation, the specific surface area of CaCO₃ molar volume was about 2.2 times as much as the CaO molar volume. CaO* for CaL process was calculated as the sum of the grain size of “active CaO” estimated by CaCO₃ product and “skeletal CaO”. The calculation results of CaO* were shown in Table 2. It could be found that the values of CaO* obtained in CaL process (from 61.0 nm to 95.1 nm) at the same cycle time of HtL and CaL processes, a new parameter CaO* was set to represent the total grain size of CaO in the adsorbent after regeneration of nano-CaCO₃ without any further sintering. For HtL process, the value of CaO* could be directly calculated by the molar volume difference between CaCO₃ and CaO (the CaCO₃ molar volume was about 21.28 kJ mol⁻¹). The activation energy of the grain growth during CaL process was 120 kJ mol⁻¹, which was located between E_CaCO₃ and E_CaO. It illustrated that the regeneration process of adsorbents involved both nano-CaCO₃ and nano-CaO grain growth, in which the influence of nano-CaCO₃ was dominant.

The grain growth model of solid-phase sintering was shown as eqn (5)–(7). First, the data in Fig. 3 were used to linearly fit $\dot{G}^2$ with $t$ in eqn (5). The nano-CaCO₃ and nano-CaO grain growth rate constants $C$ at different heat-treatment temperatures could be obtained. Then, by linearly fitting ln $C$ and $-1/T$, the nano-CaCO₃ activation energy of the grain growth $E_{\text{CaCO}_3}$ was got as 104.8 kJ mol⁻¹, and the nano-CaO activation energy of the grain growth $E_{\text{CaO}}$ was got as 212.8 kJ mol⁻¹. It could be observed that nano-CaCO₃ grains was much easier to grow than that of nano-CaO grains. In addition, according to the research of Zhu et al., the activation energy of the grain growth of nano-CaO adsorbents (the same raw material as this research) through regeneration process was 120 kJ mol⁻¹, which was located between $E_{\text{CaCO}_3}$ and $E_{\text{CaO}}$. It illustrated that the regeneration process of adsorbents involved both nano-CaCO₃ and nano-CaO grain growth, in which the influence of nano-CaCO₃ was dominant.

\[
C = C_0 \exp \left( -\frac{E}{RT} \right) \tag{6}
\]

\[
\ln C = \ln C_0 - \frac{E}{R} \times \frac{1}{T} \tag{7}
\]
In addition, the evolution of adsorbent’s pore structural parameters and pore volume distribution during 10 cycles of CaL and HtL processes were shown as Table 2 and Fig. 5 respectively. According to the results, the capillary porosity of nano-CaCO₃ was very low because the specific surface area \( \xi_{BET} \) (0.55–1.60 m² g⁻¹) and pore volume \( V_p \) (0.006–0.007 cm³ g⁻¹) measured by the BET method were both small. After CaL cycles, \( \xi_{BET} \) (3.71–14.73 m² g⁻¹) and \( V_p \) (0.041–0.092 cm³ g⁻¹) were significantly improved, as the content of nano-CaO in the adsorbent increased and more capillary pores appeared (Fig. 5). HtL process had no obvious influence on the pore structure of the nano-CaCO₃. However, a large amount of microporous loss happened with the prolongation of CaL cycles for the adsorbent, leading to the decrease of \( \xi_{BET} \) and \( V_p \) as well as the increase of the average pore diameter \( r \) (from 27.6 nm to 38.3 nm).

Fig. 6 and 7 depicted the sorption capacity and sorption rate of adsorbent during CaL and HtL processes. It could be seen that as the increase of the number of CaL cycles and HtL cycles, the sorption capacity and the sorption rate would obviously decrease. However, the extent of adsorbents’ deactivation through HtL process (from 15.34 mol kg⁻¹ to 11.95 mol kg⁻¹) was less than that of CaL process (from 10.83 mol kg⁻¹ to 4.5 mol kg⁻¹). After the same times of cycle, the sorption capacity and sorption rate of adsorbent through HtL process were higher than that of adsorbent through CaL process.

According to the above experiment results, the structural evolution of the CaCO₃–CaO conversion during the CaL process could accelerate the growth of regenerated nano-CaO grain and the disappearance of micropores. It reduced the numbers of active CaO sites, so the sorption performance of the adsorbent would rapidly decay during CaL process.

3.3 The grain growth mechanism of regenerated nano-CaO in CaL

Based on the above experimental results and the SEM images of the adsorbent through CaL process (Fig. 8(a)), the grain growth
mechanism of the regenerated nano-CaO was summarized in Fig. 8(b). According to the SEM image (a)-I, fresh nano-CaCO₃ grains were dispersed stacked cubic microcrystals (calcite). During the first regeneration step, the grain boundaries among the nano-CaCO₃ with surface contact were gradually eliminated and the grain size increased (transition state). Subsequently, the conversion between CaCO₃ and CaO with a large difference in the molar volume as well as the CO₂ gas flow among the grains would cause the internal microstructure of the adsorbent to rearrange constantly. The structure of nano-CaO became tighter with more capillary pores (SEM image (a)-II). Then, in the first carbonation stage, only small particle sizes’ nano-CaO and the surface of large particle sizes’ nano-CaO could be converted back to nano-CaCO₃, which indicating the initial decline of sorption capacity. Because the molar volume of CaCO₃ was bigger than that of CaO, many micropores were filled up. It resulted in more surface contact of nano-CaCO₃ and created conditions for the growth of nano-CaCO₃ during the next regeneration step. In the subsequent CaL processes, these steps were iteratively repeated and the proportion of “skeletal CaO” continuously increased until the micropores in the adsorbent almost disappeared (SEM image (a)-III). Finally, the growth of regenerated nano-CaO grain stopped and the sorption capacity decreased to a fairly low level.

In summary, there were two necessary conditions triggering the grain growth of regenerated nano-CaO. One was sufficiently high ambient temperature (much higher than the Tammann temperature of CaCO₃), which could be called as the external factor. The other one was the surface contact of nano-CaCO₃ grains, which could be called as the internal factor. Therefore, two strategies for limiting the regenerated nano-CaO grain growth was reducing the high-temperature residence time and preventing interface contact of nano-CaCO₃ grains.

4. Conclusions

In CaL process, the thermal decomposition of nano-CaCO₃ to produce nano-CaO was the main stage affecting the growth of regenerated nano-CaO grains, which would lead to the sintering and deactivation of adsorbents. In this work, the grain growth mechanism of the regenerated nano-CaO in CaL process was studied experimentally. By comparing the activation energy data of grain growth of nano-CaCO₃ and nano-CaO, it indicated that the grain growth of nano-CaCO₃ was easier during the high-temperature looping process. Moreover, the chemical conversion between CaCO₃ and CaO would accelerate the sintering of adsorbents by increasing the interface contact among nano-CaCO₃ grains in CaL process based on the experimental results. Among the three substeps of regeneration process of the adsorbent, the thermal growth of nano-CaCO₃ grain was the key issue to influence the grain growth of regenerated nano-CaO.

Conflicts of interest

There are no conflicts to declare.
Nomenclature

d  Average grain size, nm
K  Scherrer constant, 0.89
\(\lambda\)  X-ray wavelength, 0.154 nm
\(\theta\)  Diffraction angle, degree
\(\beta\)  FWHM of the characteristic peak in the XRD spectra
\(\bar{G}\)  Instantaneous average grain size, nm
\(t\)  Heat-treatment time, min
\(G_0\)  Initial grain size, nm
C  Grain growth rate constant, nm\(^2\) min\(^{-1}\)
T  Heat-treatment temperature, K
E  Activation energy of the grain growth, kJ mol\(^{-1}\)
\(S_{\text{BET}}\)  Specific surface area, m\(^2\) g\(^{-1}\)
V\(_p\)  Pore volume, cm\(^3\) g\(^{-1}\)
\(r\)  Average pore diameter, nm

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