Research article

Structure and phase analysis of calcium carbonate powder prepared by a simple solution method

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HIGHLIGHTS

• The engineered crystal structure and phase transformation of CaCO₃ were achieved at room temperature using Ca(NO₃)₂ and Na₂O₃ precursor.
• The vaterite phase formed at 15 min and 30 min reaction times with pH ~ 7.9 shows the spherical morphologies (2–5 µm in diameter).
• Different reaction times and pH vs. formation and transformation of various CaCO₃ phase was demonstrated experimentally.

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ABSTRACT

This paper focused on the analysis of the crystal structure and phase transformation of CaCO₃ synthesized by simple solution method from 0.5 M Ca(NO₃)₂ precursor and 0.5 M Na₂CO₃ precipitant at ambient temperature (300 K). The pH of the sample solution at various reaction times of 5, 10, 15, and 30 min were measured and correlated with the supersaturating condition in the presence of the Na₂CO₃ which is responsible for vaterite phase formation. The formation of the polymorph structure of obtained CaCO₃ powders was characterized using powder X-ray diffraction patterns and their crystal structure and phase transformation were evaluated using the Rietveld refinement method. Moreover, the qualitative analysis of the CaCO₃ powders phase was conducted by Fourier Transform Infrared (FTIR) spectroscopy to evaluate the effect of reaction time correlated with their crystal formation. The XRD analysis showed that the vaterite formation was 89 % at a reaction time of 15 min and confirmed also by FTIR that the amount of vaterite increased due to the effect of increasing reaction time. The crystallite size of vaterite was stable at 36 nm at the reaction time of 15 and 30 min. The morphology of the CaCO₃ powders obtained from Scanning Electron Microscope (SEM) was spherical with sizes of 2–5 µm. It was highlighted that the supersaturating condition started occurred at a reaction time of 15 min at pH 7.88 which was responsible for vaterite formation took place. It was concluded that the amount of precipitant (Na₂CO₃) and reaction times play an important role to determine the saturation of carbonate source to allow vaterite phase formation of CaCO₃ powders to occur.

1. Introduction

Calcium carbonate (CaCO₃) is an abundant inorganic biomaterial in the manner of different structures (calcite, vaterite, and aragonite) [1, 2]. Thermodynamically, the form of CaCO₃ under normal conditions is β-CaCO₃ (calcite). Other polymorphs of CaCO₃ such as λ-CaCO₃ (aragonite) and μ-CaCO₃ (vaterite) can be formed at certain synthesis temperatures [3]. The applications of CaCO₃ as a biomaterial are strongly dependent on their structure and phase transformation to become a concern of researchers in various fields. This material is increasingly used...
in many industries (i.e. in paint, rubber, plastics, and biomaterials) since its physical and chemical properties can be engineered [4, 5, 6, 7, 8].

The process of CaCO₃ crystals formation usually takes place in three important steps. Initially, the formation of amorphous calcium carbonate (ACC) begins, which then turns into calcite and vaterite phases at low temperatures (14–30 °C), while at 60–80 °C it becomes calcite and aragonite. In the third stage, the composition of vaterite and aragonite decreases along with the increase in calcite formation [9]. The formation of polymorphic CaCO₃ crystals results in different particle morphology. Calcite has a rhombohedral shape, aragonite has a needle-like particle shape, whereas vaterite has a spherical shape [10].

The polymorph structure and their morphology of CaCO₃ powder achieve by various synthesis techniques such as deposition of homogeneous solutions [11], water-in-oil emulsion [12, 13], mechano-chemical and sonochemical synthesis [14, 15], water-in-oil (W/O) microemulsion [16, 17], and high-pressure homogenizer (HPH) [7]. The standard engineering of CaCO₃ polymorphs, tuned shapes, and sizes are obtained by controlling over crystal growth mechanism [5]. The mechanism of vaterite crystal growth approaches is widely investigated and adopted by many researchers to synthesis vaterite crystals using additives such as organic additives and/or organic polymers to control the crystal growth, i.e. [10, 18, 19]. The influence of additives on the rate of CaCO₃ was reported by many researchers [19, 20, 21, 22]. A high concentration of mixed solvents of ethylene glycol and water affected the polymorphic composition and favoured the precipitation of the vaterite phase. The addition of ethylene glycol may reduce the CaCO₃ precipitation rate. In this condition, high supersaturation was preserved for a longer time and resulted in a higher concentration of vaterite phase, probably as a result of kinetic stabilization by slowing the growth rate of the more stable polymorphs [22]. Other researcher reports additional amino acids (glycine) to obtain the vaterite phase in the synthesis by precipitation process [23]. Recently reported that mesoporous vaterite CaCO₃ crystals synthesized by simple solution method without additive by mixing of Ca²⁺ and CO₃⁻ salts in aqueous media from Na₂CO₃ and CaCl₂ precursor [24].

Moreover, the polymorphism of the obtained CaCO₃ powder depends also on the chosen precursors. Most reported research uses Calcium Chloride (CaCl₂) as a salt precursor as a source of calcium ions [25] and Sodium carbonate (Na₂CO₃) or potassium carbonate K₂CO₃ as anions source [26]. The advantage of CaCl₂ is very easily dissolved in water to generate calcium ions and react with the carbonate source to form a solid precipitate of calcium carbonate due to an exothermic process [27]. However, governing the crystal growth in the exothermic process is quite difficult without controlling the temperature. Also, irregular cubic particles are obtained when the crystal growth process by stirring the solution for a longer time [28]. Another important factor that affects the formation of CaCO₃ polymorphs is the pH of the solution during the precipitation process. At particular temperature and concentration of the solution. At 24 °C and a low concentration of CaCl₂ solution (0.0134 M), pure calcite is obtained at a pH of more than 12, aragonite is obtained at a pH of 11, and vaterite is obtained at a pH less than 10 [29]. Mitsuaki Kogo et. al. [30] reported the adjustment of the pH from 2.0–11.3 to the initial pH of 0.1 M CaCl₂ and Na₂CO₃ solution of 6.5 by adding HCl and NH₃ to investigate the polymorph process. From the result of X-ray diffraction, single-phase vaterite is formed at initial pH 2.5 with spherical shape and particle size of 1–10 μm [30]. In contrast, Radek Sevčik et. al. obtain high pure vaterite (99 wt%) from high concentration solution (2 M CaCl₂ precursor and K₂CO₃ precipitant) provided higher temperature (60°) and higher stirring speed of 600 rpm [31]. However, the morphology of the obtained particles are irregular in shape and mainly as big bodies up to 10 μm in size may be due to massive crystallization of many nuclei that joined together during growth at high concentrations of the solution. To the best knowledge of the author and review from reported research, the use of calcium nitrate Ca(NO₃)₂ as a source of cations is rarely used as a precursor for CaCO₃ synthesis. Interestingly, the reaction of Ca(NO₃)₂ in water is endothermic consequently no need for effort to control the temperature process if compared to CaCl₂.

Recently, the fabrication of CaCO₃ crystals with sizes beyond the conventional range without additives is still a challenge [24]. Our current study provides the polymorphic transformation from calcite stable phase to vaterite metastable phase without any additives and templates by a simple method. In this study, we analyzed the crystal structure and phase transformation of CaCO₃ synthesized by simple solution method from Ca(NO₃)₂ precursor and Na₂CO₃ precipitant. The crystal structure and phase transformation were analyzed using the Rietveld refinement method [32]. This method is a sophisticated procedure for quantitative analysis of CaCO₃ powder containing a mixture of calcite and vaterite from powder X-ray diffraction (XRD) data at various reaction times. The investigation on the structure and phase transformation also correlated with the pH of the solution at various reaction times. The final particles are spherical in shape with porous morphology. The formation of porous polycrystalline vaterite was markedly different from the conventional particle to particle formation in which nucleation occurred.

2. Material and method

2.1. Material

In this research, all of the materials we use are high-quality commercial products. Sodium carbonate (Na₂CO₃) and calcium nitrate tetrahydrate (Ca(NO₃)₂·4H₂O) were purchased from Merck (Indonesia).

2.2. Synthesis of CaCO₃ powder

The synthesis of CaCO₃ by a simple solution method is schematically shown in Figure 1. Since higher mixing temperature might initiate the formation of aragonite and calcite phases [6], the favoured range of temperatures for vaterite phase precipitation in various experiments is up to 40 °C [26]. In the current study, the temperature of 27 °C has been chosen for the experiment by mixing the solutions under 800 rpm to obtain homogenous, high-shear, and sufficient agitation to enable vaterite phase precipitation.

The calcium carbonate synthesis process started by mixing between 0.5 M Na₂CO₃ and 0.5 M Ca(NO₃)₂·4H₂O solutions. The solution was mixed continuously at a temperature of 27 °C (T = 300 K), with the fixed stirring speed at 800 rpm with different reaction times (5, 10, 15, and 30 min). The solution was filtered to get the precipitated powder. The obtained powders were washed with ethanol to remove the remaining contaminant indicated by the white appearance of the powder. The powder was then dried in a vacuum desiccator for 48 h and subjected to analysis. The pH of the initial solution (0 min) and after the reaction time at various reaction times (5, 10, 15, and 30 min) was measured as a parameter in the formation of the structure and phase of the CaCO₃ powder.

2.3. Characterizations

We used X-ray diffraction measurements (PANalytical) to analyze the crystal structure of the synthesized CaCO₃ powder. Data were collected over a 2-theta in the range of 20–80° with a time step of 2.90 s. Cu-Kα X-ray radiation at a wavelength of 1.5406 Å was used. Synthesized CaCO₃ powders were measured using Fourier Transform Infrared Spectroscopy (FTIR, Nicolet iS5, Thermo Scientific) to determine the vibration mode. The measured infrared spectrum region was in the range 600–1000 cm⁻¹ represented the fingerprint area that was used in analyzing the polymorphous vibration mode of the synthesized CaCO₃ powder [33]. The scanning electron microscope (SEM, Hitachi SU3500) with an accelerating voltage of 10 kV was used to characterize the morphology of CaCO₃ powders. The morphology of obtaining CaCO₃ powder used to identify the shape of the particle at various reaction times indicated the presence of rhombohedral (calcite) and spherical (vaterite) shapes.  

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2.4. Crystal structure analysis and phase identification

The structural analysis including the crystallite size and phase identification of the synthesized powder employed open-source software, MAUD [34]. We used the crystal structure of calcite as trigonal [35] and the vaterite is orthorhombic [36] in the entire analysis. The unit cells of calcite and vaterite are represented three-dimensionally by the atoms of each crystal as shown in Figure 2 were visualized by the VESTA software package [37]. The detailed quantitative results based on Rietveld refinement method analysis were then used for crystallite size calculation using the Scherer method [38]. Qualitative analysis of the synthesized CaCO₃ powder was carried out by comparing the vibration bending mode ratio between the vaterite phase in the 744 cm⁻¹ area and calcite which was located at the wave number 711 cm⁻¹ [39].

3. Results and discussions

3.1. Structural analyses of CaCO₃ powder

The XRD pattern of the polymorphs of CaCO₃ is shown in Figure 3. The XRD patterns characteristic peaks corresponding to the stable phase (calcite) and metastable phase (vaterite) were observed at T = 300 K in various reaction times from 5–30 min. It was observed the transformation of the calcite phase to the vaterite phase occurred at various reaction times. When the reaction time for calcium carbonate precipitation was only 5 min, the majority of the crystals formed were calcite (94%) and contained only a small phase of vaterite (6%). After the reaction time was increased to 10 min, the precipitate contained a mixture of calcite (52%) and vaterite (48%) crystals. It showed that dissolution re-precipitation was occurred to form polymorph crystal of calcium carbonate [40]. At the reaction time of 15 min, it provided a condition to create the new solid particles, i.e., vaterite (90%) as a major phase and calcite (10%) as a secondary phase. This result indicated that reaction time 15 min appropriate condition to form calcium carbonate precipitate with a vaterite crystal structure. In 30 min of reaction time, the vaterite phase was decreased slightly (89%) due to the crystal formation process of a stable phase of calcite about 11%. In conclusion, the major calcite phase was obtained rapidly at 5 min of reaction times, and we could find the appropriate reaction time of the vaterite phase at 15 min, and it was highlighted the reaction occurred at room temperature.

The crystallite sizes of calcite decreased significantly due to the defects that occurred in the transformation process (Table 1). While the crystallite sizes of vaterite showed only slight changes. The result described that the crystallite size for both calcite and vaterite were stable occurred at the reaction time of 15 and 30 min. The characteristic of crystal also determined by the volume of unit cell represented of the density of the unit cell might be affected by reaction time. The effect of reaction time on the formation of CaCO₃ precipitation on the unit cell volume of calcite and vaterite crystals phase was shown in Table 2. At the reaction time of 5 min, the volume of the unit cell of the two crystals

![Figure 1. Schematic illustration of synthesis CaCO₃ by simple solution method.](image1)

![Figure 2. Space-filling representation of the crystal structure of trigonal calcite (a) and orthorhombic vaterite (b) in three dimensional lattice vectors; blue, brown and red sphere represent Ca atom, C atom, and O atom.](image2)

![Figure 3. XRD patterns of synthesized CaCO₃ powder at various reaction times.](image3)
shown in Figure 4. For precipitated CaCO₃ samples with a reaction time weak (minutes)

\begin{table}[h]
\centering
\caption{Unit cell volume of synthesized CaCO₃ powder at various reaction time.}
\begin{tabular}{cccc}
\hline
Reaction time (minutes) & Calcite Unit Cell volume (Å\(^3\)) & Vaterite Unit Cell volume (Å\(^3\)) &
\hline
Ref. data (AMCS) & 367.9162 & 250.4102 &
5 & 367.1829 & 249.3785 &
10 & 367.9314 & 250.4102 &
15 & 367.4410 & 249.6451 &
30 & 367.4110 & 249.6411 &
\hline
\end{tabular}
\end{table}

(calcite and vaterite) was still close to reference [35, 36]. The smallest of the unit cell volume for both calcite and vaterite were formed at the reaction time of 15 min. Theoretically, we know that the volume of the unit cell is inversely proportional to the density of the unit cell. Thus, we conclude that the highest density of calcite and vaterite unit cells occurred at the reaction time of 15 min.

3.2. Vibrational mode analyses

Infrared spectroscopy serves as a useful tool to study different calcium carbonate phases (calcite, aragonite, and vaterite) since each phase shows characteristic absorption bands in its infrared spectrum [33]. The regions of the absorption bands corresponding to the two fundamental vibrational modes of carbonate ion at ~700 cm\(^{-1}\) (in-plane bending, \(\nu_4\)) and 877 cm\(^{-1}\) (out of plane bending, \(\nu_2\)) are assigned [31]. The \(\nu_4\) band at ~700 cm\(^{-1}\) is a fingerprint region and distinct for crystalline phases (calcite, aragonite, and vaterite) provides useful information for CaCO₃ polymorph assignment. The FTIR spectroscopy of CaCO₃ powder is shown in Figure 4. For precipitated CaCO₃ samples with a reaction time of 5 min, the absorption peak of calcite was observed at 711 cm\(^{-1}\). The weak \(\nu_2\) band at 877 cm\(^{-1}\) for calcite indicate defective crystal structure [41]. The absorption peaks at 711 and 744 cm\(^{-1}\) were assigned for mixture calcite-vaterite precipitate at reaction time reach up to 10 min. At the prolonged reaction time (10–30 min), the phases of vaterite (\(\nu_4\) band at 744 cm-1) was increased, while the \(\nu_4\) band for calcite at 711 cm\(^{-1}\) was suppressed. The concentration ratio of vaterite–calcite in the calcium carbonate samples from FTIR spectra can be determined by using the absorbance peak ratio at \(\nu_4\) band which was given in Table 3. Qualitatively, the amount of vaterite increased due to the effect of increasing reaction time. It was shown that the optimum reaction time for the vaterite phase had occurred at 15 min.

3.3. Morphology of CaCO₃ powder

The morphology of CaCO₃ powder which was influenced by the reaction time was shown in a grayscale image showing the topography of the sample in Figure 5a–d in a combination of rhombohedral and spherical particles [42]. Figure 5a indicated that the calcium carbonate precipitate was a rhombohedral shape, which was the morphology of calcite. Figure 5b, rhombohedral calcite particles, and spherical vaterite particles were precipitated. This occurred because the reaction time was increased up to 10 min, the saturation level of calcium carbonate increases so that the shape becomes irregular, calcite–vaterite mixture. The calcite phase started to decrease with increasing reaction time and the vaterite phase started to appear. Figure 5c, the morphology of the calcium carbonate precipitate started to be spherical but still showed a rough surface. Under this condition, the majority of the powders synthesized were shown to be in the vaterite phase. In Figure 5d, the calcium carbonate precipitated were spherical with a smoother surface, which indicated the reaction time was appropriate for vaterite crystal growth. The TEM images showed that the synthesized CaCO₃ powders were spherical in shape with sizes of 2–5 μm. More spherical vaterite particles obtained at 15 and 30 min reaction times (Figure 5) confirmed the presence of the vaterite phase as the dominant phase obtained from XRD quantitative analysis (Figure 3). The presence of the rhombohedral calcite phase was also remarkably reduced. Our results are consistent with the work studied by Baqiya et al. [43]. They found that when the weight fraction of the vaterite phase is quite high (~95%), the particles tend to have spherical morphology due to a decrease of concentration between CO\(^2\) and Ca\(^{2+}\) which may be the reason for the formation of the vaterite phase.

TEM analysis confirmed morphology of obtained vaterite microstructure as shown in Figure 6. The TEM images showed the CaCO₃ microsphere consisted of primary particles with relatively homogeneous in size around 150 nm. A multipoint BET measurement was conducted. BJH (Barret, Joyner, and Halenda) method was conducted to calculate pore surface area, pore size from experimental isotherm using adsorption and desorption technique. Only BJH desorption results were shown in the table as the desorption technique gives a narrower pore size distribution than the adsorption technique. BET analysis proved the presence of porous

\begin{table}[h]
\centering
\caption{The bending mode vibration of absorbance ratio of synthesized CaCO₃ powder at various reaction times.}
\begin{tabular}{ccc}
\hline
Reaction time (minutes) & \(A_{\nu_4}/A_{\nu_1}\) ratio &
\hline
5 & 0.3385 &
10 & 1.4803 &
15 & 3.4225 &
30 & 3.1540 &
\hline
\end{tabular}
\end{table}
particles that correspond to the particle as shown in the SEM image (Figure 5). The particles possess an average pore diameter in the size of 3.58 nm, which is typical for the vaterite phase [24]. The particles possess a specific surface area of 10.33 m²g⁻¹. This result confirmed the polymorphic transformation from calcite stable phase to vaterite meta-stable phase without any additives and templates.

The analysis of crystal structure and phase transformation, and morphology of CaCO₃ powder at various reaction times in agreement with the pH solution parameter of carbonate formation is shown in Figure 7. The initial pH of the solution was 10.2 indicated that constituent precursors were well dissolved. At 5 min reaction time, the pH of the solution was 8.76. In this condition, the CaCO₃ powder was predominantly calcite in the form of rhombohedral particles. At 10 min reaction time, the pH of the solution was 8.1 indicate the mixture of calcite-vaterite. It was highlighted that when the pH solution dropped closer

Figure 5. SEM images of synthesized CaCO₃ particles at various reaction times (a) 5 min, (b) 10 min, (c) 15 min, (d) 30 min.

Figure 6. TEM (a) and SAED (b) images of synthesized CaCO₃ particles at 30 min reaction time.

Figure 7. The pH of the solution at various reaction times (5–30 min).
transformation of the synthesized CaCO3 powder can be utilized to our general conclusion, the analysis of crystal structure and phase at alkaline pH ~ 7.9 with spherical morphology (2 μm) [3]. According to Ostwald’s step rule [45], at higher supersaturation, the difference between interfacial energy of polymorphs is prevalent and the metastable vaterite will be formed predominantly. Our result was well aligned with Han et al. result [46] which showed that an increase in the pH value (above 8) resulted in a decrease in the vaterite concentration. When pH decreased from 9.7 to 7.7, almost pure vaterite was produced.

4. Conclusions

We have successfully analyzed the crystal structure and phase transformation of CaCO3 synthesized by simple solution method at room temperature (T = 300 K) using a Ca(NO3)2 and Na2CO3 as a precursor. The vaterite phase was formed within the reaction time of 15 and 30 min at alkaline pH ~ 7.9 with spherical morphology (2-5 μm) in diameter. In our general conclusion, the analysis of crystal structure and phase transformation of the synthesized CaCO3 powder can be utilized to investigate the formation of various carbonate at different reaction times and pH of the solution.

Declarations

Author contribution statement

Renny Febrida: Conceived and designed the experiments; Performed the experiments; Analyzed and interpreted the data; Wrote the paper.

Setianto Setianto: Performed the experiments; Analyzed and interpreted the data; Contributed reagents, materials, analysis tools or data; Wrote the paper.

Elyza Herda: Conceived and designed the experiments; Wrote the paper.

Arief Cahyanto: Analyzed and interpreted the data; Wrote the paper.

I Made Joni: Conceived and designed the experiments; Analyzed and interpreted the data; Contributed reagents, materials, analysis tools or data; Wrote the paper.

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Data availability statement

Data included in article/supplementary material/referenced in article.

Declaration of interests statement

The authors declare no conflict of interest.

Additional information

No additional information is available for this paper.

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