A novel hydrothermal synthesis of nanohydroxyapatite from eggshell-calcium-oxide precursors

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ABSTRACT

Hydroxyapatite (HA) is a material that has been widely applied to replace the damaged bone as a bone implant. Different types of HA have been successfully synthesized by a hydrothermal method based on calcium oxide (CaO) which was originated from chicken eggshells and diammonium hydrogen phosphate (DHP)/(NH4)2HPO4 as their precursors. We present a novel approach to the hydrothermal synthesis of HA form eggshells as a new precursor via a one-step synthesis method. The influence of temperature was also observed to study the effect on the crystallinity, purity, and morphology of obtained HAs. The synthesis was carried out at two different temperatures, 200 °C (HA-200) and 230 °C (HA-230) for 48 h respectively. The structures, purities, and morphologies of hydroxyapatite were analyzed by X-ray Diffraction (XRD), Fourier Transform Infra-Red (FTIR), and Scanning Electron Microscopy- Energy Dispersive Spectroscopy (SEM-EDS), and Transmission electron microscopy (TEM). The XRD patterns show the HA main phase indicated the purity of 96.5% for HA-200 and 99.5% for HA-230. The TEM micrograph suggested a hexagonal-like of HA with an average particle size of 92.61 nm. Hexagonal-like of HAs are suitable for bone implants and further application.

1. Introduction

Hydroxyapatite (HA), Ca10(PO4)6(OH)2, is a mineral that has an important role in the component of hard tissue [1]. HA is a calcium phosphate-containing hydroxide that has a Ca/P ratio of 1.67. HA can help the growth of new bone without having to use fibrous tissue. HA can replace the damaged bone caused by the tumor, congenital diseases, and others as bone implants [2, 3]. HA is currently the most widely applied as materials in biomedical applications due to its stable chemical properties, composition, and crystal structure similar to human bone [4]. HA also can be the main-constituent materials of a composite for a bone implant such as gelatin/hydroxyapatite composite modified with osteoblast for bone bioengineering [5] and HA-based high-density polyethylene bio-composites [6].

Several studies have been performed to synthesize HA using natural materials as calcium sources such as coral, seaweed, and chicken eggshells [7, 8], shells [6], and starfish [2]. The chicken eggshells are a good source of calcium that was easily obtained than the other source of calcium. In addition, waste eggshells are abundant in Indonesia. The eggshells produced in a year are 138,956 tons [9], whilst the eggshells CaCO3 content could have reached a percentage of 94% [10]. The chicken eggshells showed a suitable potential for the synthesis of high-quality hydroxyapatite as a good implant material [11]. Hence, the chicken eggshells potentially become a source of calcium in the synthesis of HA.

HA can be synthesized using various methods including solid phase reaction method [12], precipitation [13], and hydrothermal [2, 11]. Pei et al. [12] synthesized HA using CaCl2 as a source of calcium by the solid-state reaction method at 775 °C. Yeten & Suat [13] synthesized HA from Ca(OH)2 as a source of calcium by precipitation method. The extraction process and natural waste source in the HA synthesis have an effect on the critical properties of the HA such as Ca/P ratio, crystallinity...
and phase assemblage, particle sizes, and morphology [14]. In the synthesis process, there are two phases: HA formation of Ca(OH)₂ and phosphoric acid at a temperature of 50–60 °C with a pH that must be controlled every last drop followed by a sintering process of HA synthesized at 900–1000 °C. Rodriguez-Lugo et al. [2] synthesized HA from CaO derived from starfish as a source of calcium and β-tricalcium phosphate by using the one-step hydrothermal method at 230 and 250 °C and produced homogeneous crystalline of HA. However, other phases of HA were still observed.

The hydrothermal method can be deployed for a synthesis of high-crystalline and doped apatite-based materials [15, 16]. Thus, we utilise the method to provide good quality, purity, and high yield in HA synthesis. The method can be fruitful to acquire an excellent structural building of HA. Some of the advantages with this method are low cost, short reaction steps, low-temperature synthesis, simple method, and obtaining good quality of HA. This method is also associated with a low energy dissolution process, so it is easy to form pure and homogeneous crystalline HA and unique morphology at relatively low temperatures [17, 18].

Hydrothermally HA synthesis can be influenced by several factors, such as temperature, time, and pH during the synthesis process. From the set of results obtained in this research, it can be established that the best synthesis conditions for HAS are a temperature of 250 °C during a reaction time of 48 h. These parameters yielded the best characteristics of HA, including a more uniform size, greater amounts, homogeneity with a Ca/P ratio near stoichiometric, and a higher crystalline degree [19]. HA in various pH condition were successfully synthesized by hydrothermal method [20]. At high pH at 9, the form of HA will grow into spherical shape, meanwhile at lower pH up to 6, will grow into one-dimensional nanorods or two-dimensional nanoplates. Additionally, more complicated shapes, including three-dimensional microfibers, three-dimensional microtubes, and three-dimensional featherly are only obtained if the pH value decreases to 4.

Based on those studies, we conducted the HA synthesis via a one-step hydrothermal method using the variant source of CaO from chicken eggshell [2] and dissimilar phosphate source of DHP solution [9], more economically priced than β-tricalcium phosphate. The synthesis step reaction was noted in the following reaction (1):

\[ 10\text{CaO} + 6(\text{NH}_4)_2\text{HPO}_4 + 4\text{H}_2\text{O} \rightarrow \text{Ca}_6(\text{PO}_4)_2(\text{OH})_2 + 12\text{NH}_3\text{OH}(\text{aq}) \]

Meanwhile, the reaction underwent abasic pH between 9 and 10 to obtain a spherical shape for the application in the dental filling.

2. Experimental method

2.1. Conversion of CaCO₃ to CaO from chicken eggshell

Chicken eggshells were washed with water to remove the contaminants, then crushed by used planetary ball mill until 100 meshes. After being crushed, the material was calcined at 1000 °C for 5 h for the complete conversion of CaCO₃ to CaO. The phase composition of the material was characterized by XRF (Rigaku NEX CG).

2.2. Synthesis and characterization of HA

The synthesis was carried out at a different temperature of 200 °C (HA-200) and 230 °C (HA-230). CaO and DHP with the molar ratio of Ca/P = 1.67 were poured to distilled water (50mL). The mixture was then put into an autoclave 100mL and heated at 200 °C for 48 h. The obtained HA was filtered and washed with aquadest until pH = 7 to remove the NH₃OH and then dried at 110 °C for 2 h. The same procedure was repeated for the synthesis of HA at 230 °C. HA crystal structure identified by XRD (PANalytical XPERT PRO seri PW 3040/00) using CuKα radiation at 2θ 5°–70°, the functional group of HA was confirmed by FTIR spectrosopies (PerkinElmer Spectrum 100). HA morphology and elemental mapping analysis were characterized by SEM-EDS (Jeol jsm-6360LA) and Transmission Electron Microscope/TEM (JEOL JEM-1400plus TEM Single tilt holder TEM Grid Cu 400 mesh).

3. Result and discussion

3.1. Calcination of chicken eggshells

At the temperature of 750 °C, CaCO₃ started to decompose into CaO, then completely decomposed at 1000 °C with the loss of ignition (LoI) of 45.96% due to the decomposition of chicken eggshells from CaCO₃ to CaO, as shown in reaction (2) [2]. The phase compositions of chicken eggshells after calcination at 1000 °C, are shown in Table 1.

\[ \text{CaCO}_3(\text{s}) \rightarrow \text{CaO(})_1 + \text{CO}_2(\text{g)} \]

The CaO content was 97.80%, which was then used to calculate the molar ratio of Ca a source of the HA formation reaction with DHP as a phosphate source.

3.2. Structural characterization of HA by XRD

The XRD patterns of HA-200 and HA-230 are shown in Figure 1. The main peak of HAs was indicated at around 2θ ~31°. The portlandite phase was observed at 2θ ~17.2°.

The purity of HA-200 and HA-230 was 96.5% and 99.5% with 3.5% and 0.5% secondary phase portlandite (Ca(OH)₂) respectively. This fact suggests that the reaction temperature can affect portlandite Ca(OH)₂ content. The lower reaction temperatures produced more portlandite. The higher the reaction temperature, the higher a reaction rate between Ca(OH)₂ and DHP, the more HA was produced. HA-200 and HA-230 have a hexagonal structure with space group P63/m with slightly different lattice parameter as shown in Table 2.

Different temperatures of synthesis were also affected to the crystal size and crystallinity of HA. The crystallinity of synthesizing HA was calculated from the XRD data using the Debye-Scherrer equation, Eq. (3) below [19]:

\[ X_c = 100 \times (I_{100} - V_{112/300})/I_{100} \]

where I₁₀₀ is the intensity of the diffraction peak at the (300), V₁₁₂/₃₀₀ is the intensity of the (112) and (300). The crystallinity of HA-230 is 72.76%, meanwhile, HA-200 is 67.23%. This phenomenon complies with another study we found demonstrating that the increase in processing temperature had a great effect on the particle size and crystal structure of HA. The lower reaction temperatures produced more crystals. Colder solutions contract, adjusting atoms accelerating the crystal growth rate. The disparate temperatures produce different amounts of crystals. Colder solutions contract, adjusting atoms in the HA structure assemble forcing bonds to develop, engaging impurities in the structure simultaneously. These impurities subsequently interrupt the crystal pattern, supporting the formation of a larger number of smaller crystals. In warmer temperatures, the distance between molecules is wider, which allows crystals to form larger, purer shapes at a much more uniform rate than can occur in colder temperatures implying that the crystallinity would be larger by deploying a higher temperature [23].
3.3. FTIR characterization of HA

To confirm the functional groups of two different types of HA, FTIR spectroscopy analysis was also conducted. Figure 2 shows the FTIR spectra of HA-200 (top) and HA-230 (bottom). The HA-200 showed the stretching of the P–O bond from the PO$_4^{3-}$ at 566, 962, and 1025 cm$^{-1}$. The bending of the P–O bond from the PO$_4^{3-}$ of HA was observed at 602 cm$^{-1}$, the O–H stretching and bending appeared at 3571 and 632 cm$^{-1}$, respectively [2]. The formation of carbonated apatite with B-type substitution (tetrahedral positions) was indicated by the presence of a peak at 1401 cm$^{-1}$ (H–O–H) [11]. Overall the FTIR HA-200 and HA-230 spectra were very similar. This implies that the difference in the temperature of synthesis does not influence the dissimilarity of HA functional groups. The functional groups detected on HA-200 and HA-230 are shown in Table 3. We can perceive that HA-200 forms a broader band at ~1000 cm$^{-1}$ than the HA-230 spectrum which indicates a difference of their PO$_4^{3-}$ asymmetric stretching.

The HA formed in this study is slightly different from the HA obtained from a previous paper [2] which exhibited five of the nine bands of the PO$_4$ ion and one corresponding to the OH ions. The differences in source precursors (in this study, DHP was used as a phosphate source, while reference studies were used β-TCP), might cause small differences in shifts and intensities for P–O vibrational modes in PO$_4$ groups. The bands emanate from 910 cm$^{-1}$ and 715 cm$^{-1}$ which correlates to the P–O vibration mode of the PO$_4$ ion. Furthermore, the formation of a band localized at 3650 cm$^{-1}$, which perhaps due to the emergence of HO hydroxyls, suggested the Ca(OH)$_2$ formation. It would also be feasible to characterize a band due to the H–O–H mode, which was related to moisture located at 3460 cm$^{-1}$ which may also be distinct the HA spectrum in this study.

3.4. Morphology characterization of HA using SEM-EDS

Both the HA-200 and HA-230 morphologies are agglomerates with spherical and porous particle shapes. The morphology SEM of HA-230 is shown in Figure 3 (b) denoted a more homogenous flock than the HA-200 (Figure 3. (a)). It suggests that the increase in temperature allows the size of the particles to grow larger (in terms of diameter) and the form homogeneity is quite high [2]. Higher synthesis temperatures will produce particles with a more regular form because at higher temperatures they have higher crystallinity [24]. The regular form could be perfectly suitable for bone implantation [24]. It is known that the surface shape and texture of implants play a significant role in regulating tissue response and cell activities [25]. Hulbert et al. [26] studied the tissue reaction of three ceramics with porous and non-porous structures in soft and live tissue. The results showed that tissue around porous implants healed quickly.

Semi-quantitative analysis of EDS shows that the Ca/P molar ratio of HA-200 is 2.44 and the Ca/P molar ratio of HA-230 is 2.29. Table 4 shows that calcium contents in HA-200 were slightly higher (47.02%) than HA-230 (45.98%) while the phosphor contents were lower that prompted a larger ratio of Ca/P in HA-200 than in HA-230. This result exceeds the Ca/P molar ratio of stoichiometric HA. This is due to the possibility of contamination of carbonate (CO$_3^{2-}$) as shown in FTIR spectra from the air which can react with HA, causing a decrease in PO$_4^{3-}$. In addition, it can be caused because the HA phase produced is not 100% but there is a by-product, Ca(OH)$_2$ as shown in the XRD result so that it changed the

### Table 1. Phase Composition of chicken eggshells after calcination.

| Component     | Composition (%) |
|---------------|-----------------|
| CaO           | 97.80           |
| P$_2$O$_5$    | 0.29            |
| Fe$_2$O$_3$   | 0.01            |
| SiO$_2$       | 0.16            |
| K$_2$O        | 0.18            |
| SO$_2$        | 0.16            |
| Al$_2$O$_3$   | 0.18            |
| MgO           | 1.17            |

### Table 2. The lattice parameter, crystallite size and percentage of crystallinity of HA-200 and HA-230.

| Samples     | Lattice parameter | Crystallite size (nm) | Crystallinity (%) |
|-------------|-------------------|-----------------------|-------------------|
|             | a – b (Å)         | c – (Å)               |                   |
| HA-200      | 9.4155(8)         | 6.8786(3)             | 24.57             | 67.23            |
| HA-230      | 9.4168(1)         | 6.8794(1)             | 35.28             | 72.76            |

Figure 1. XRD pattern of HA-200 and HA-230.

Figure 2. The FTIR spectra of HA-200 (a) and HA-230 (b).
The molar ratio of Ca/P from both HA-200 and HA-230. To confirm the difference of Ca/P ratio in both HA-200 and HA-230, the elemental mapping was generated as shown in Figure 4.

Elementary mapping analysis is performed to determine the distribution of elements in HA. The colors besides the SEM morphology express the intensity of the distribution of certain elements. The color order from top to bottom indicates a decrease in the distribution of these elements at the specified hydroxyapatite spot. For example, in HA-200 the highest oxygen distribution is in the middle to the bottom (red) and appears to decrease in the left side (blue). Oxygen and phosphorus distribution in HA230 is higher than that HA200, while the distribution of calcium is almost the same in HA -200 and HA-230. The analysis of oxygen, calcium and phosphorus mapping on both HA supports the results of the XRF analysis which confirms the Ca/P ratio at HA-230 is lower than the Ca/P ratio at HA-200.

The grain shape of HA particles was confirmed using TEM as shown in Figure 5. The micrograph indicated hexagonal-like-to-round shapes of HA. The average particle size is 92.61 nm with the largest to the lowest size denoted 128.77 nm and 63.89 nm respectively. We found that the agglomeration occurred at HA-230 which shows larger particles than HA-200. This corroborates the evidence of higher crystallite size (35.28 nm) and crystallinity (72.76%) of HA-230.

Table 3. Comparison of FTIR spectra of HA-200 and HA-230.

| No | Functional Groups          | Wavenumber at 200 °C (cm⁻¹) | Wavenumber at 230 °C (cm⁻¹) |
|----|----------------------------|------------------------------|------------------------------|
| 1  | PO₄²⁻ stretching           | 566 and 962                  | 565 and 962                  |
| 2  | PO₄²⁻ asymmetric stretching| 1025                         | 1043                         |
| 3  | PO₄²⁻ asymmetric bending   | 602                          | 602                          |
| 4  | Free O-H stretching        | 3571                         | 3571                         |
| 5  | H−O−H                     | 3422                         | 3422                         |
| 6  | O−H bending                | 632                          | 632                          |
| 7  | CO₃²⁻                     | 1403                         | 1401                         |

Table 4. Chemical Composition of HA-200 and HA-230 was analyzed by using SEM.

| Element | HA-200 | HA-230 |
|---------|--------|--------|
| Ca      | 47.02  | 45.98  |
| P       | 14.93  | 15.56  |
| O       | 38.05  | 38.45  |

Figure 3. SEM image of HA-200 (a) and HA-230 (b) surface.

Figure 4. The elemental mapping of oxygen (O), calcium (Ca), and phosphor (P), respectively in HA-200 and HA-230. The distribution of each element is shown in a different colour.
4. Conclusion

HA-200 and HA-230 were successfully synthesized using the hydrothermal method from the chicken-eggshells calcium oxide. It can be concluded that the hydrothermal synthesis method of HA performed excellent results with a one-step synthesis using eggshells source of calcium as the precursor. HA-230 demonstrated better results with a lower Ca/P ratio (2.29) other than HA-200 with Ca/P ratio (2.44). The XRD resulted in an additional peak to HA-200 which corresponded to Ca(OH)₂ impurities. This suggested that HA-230 had a better structure than that HA-200. Therefore, HA-230 is considered most appropriate as a biomaterial for implantation due to the proximity of the structural and composition to the commercial HA. In addition, to obtain HA with 92 nm of average particle size, the novel-one-step synthesis of HA is also suggested. The hexagonal-like structure would be considered suitable for biomaterial applications such as a bone implant. The developments of chicken eggshells are expected to be prevalently applicable in miscellaneous implementation.

Declarations

Author contribution statement

Atiek Rostika Noviyanti: Conceived and designed the experiments.
Nur Akbar, Retna Fauzia Purti: Performed the experiments.
Yusi Deawati, Yoga Trianzar Malik: Analyzed and interpreted the data.
Risdiana: Contributed reagents, materials, analysis tools or data.
Engela Evi Ernawati: Analyzed and interpreted the data.

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Competing interest statement

The authors declare no conflict of interest.

Additional information

No additional information is available for this paper.
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