Carbonated calcium hydroxyapatite (CHAp) exhibits excellent biocompatibility with bone and teeth, making it an ideal candidate for orthopedic and dental application. However, the study of CHAp synthesis from natural material is still scarce. The purpose of this research is to synthesize and characterize CHAp, using Rebon shrimp (Acetes erythraeus) as a calcium source. The synthesis was conducted by hydrothermal method with the variation of Ca/P ratios 1.61; 1.67; 1.73. The as-prepared CHAp was characterized by Fourier transform infrared spectroscopy (FT-IR), X-ray diffraction (XRD), and scanning electron microscopy-energy dispersive X-ray (SEM-EDX). The FT-IR results show that synthesized material exhibited characteristic CHAp band of hydroxide at 3448 and 1635 cm⁻¹, carbonate at 872 and 1427 cm⁻¹, and phosphate at 1049; 606; and 570 cm⁻¹. The diffractogram pattern assigned the all observed peak of CHAp are in good agreement compared to CHAp database with the nano-scale size. It also observed that the high Ca/P ratio will decrease the crystallinity of CHAp. The as-prepared CHAp micrograph is agglomerates spherical form with size between 5-20 nm which build up from 18–26 nm crystallite particles. The result of this research confirmed that Rebon shrimp is the promising materials for calcium source in CHAp production.

**Keywords:** Dental restoring candidate material, Carbonated calcium hydroxyapatite (CHAp), Nanoparticles, Calcium Source Rebon shrimp (Acetes erythraeus).

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1. **INTRODUCTION**

In present decade, there has been observed a huge progress in the field of dentistry (Deb & Chana, 2015). Not only from dental techniques and methods of therapy, the progress has been occurred in significant development of biomaterial engineering (Matinlinna, 2013). The science of the biomaterials is consistently increasing due to ingenious modification of previously known materials or completely novel materials for application in dentistry. Bio-glass, biopolymers, bioactive ceramics, and above all, composite materials are the leading directions in the development of dental materials (Vishwakarma et al., 2014).

One of the more immensely studied groups of materials comprises inorganic calcium apatite known as hydroxyapatite (HAp). HAp is grows in hexagonal crystals and naturally occurring as the calcium phosphate minerals (Vallet-Regi, 2014). The color of pure HAp is white and it build up most of the tooth enamel, bone structure, and small amount in part of the human brain (Xie et al., 2012). The first use of synthetic HAp to repair and regenerate bone tissue was reported by Albee (1920), who utilized crystalline HAp to repair surgically induced bone defects in rabbits (LeGeros, 2008). In 1960s, bio-ceramics based on HAp was begun to greater interest when it seen as biomaterials that possibly induces the reconstruction and repair
of bone defects (Dorozhkin, 2011). In 1970s, the researchers began to pay attention to the use of HAp in dental surgery (Vallet-Regi, 2014). In particular, the work of Denissen and de Groot in which were used the natural HAp as dental implant (Pajor & Pajchel, 2019). The research of synthetic HAp has accelerated when it start permanently introduced as dental-restoring implant in 1980s (Xie et al., 2012). Recently, the composition in HAp structure can be modified to contain carbonate thereby providing a composition more similar to the bone and teeth material (Garskaite et al., 2014; Odusote et al., 2019).

Carbonated calcium hydroxyapatite (CHAp) is found in the mineral component of natural bones and teeth and when the concentration of CO$_3^{2-}$ ions in the HAp lattice is in the range of 3–8 wt% (Garskaite et al., 2014). The OH or PO$_4^{3-}$ group can be replaced by the CO$_3^{2-}$, leading the A- or B-type of CHAp, respectively. The chemical formula for A-type CHAp has been defined as Ca$_{10}$(PO$_4$)$_6$(OH)$_2$$_2$(CO$_3$)$_3$, where 0 ≤ y ≤ 1, while for the B-type Ca$_{10-x}$(PO$_4$)$_6$$x$(CO$_3$)$_3$(OH)$_2$$_2$, where 0 ≤ x ≤ 2 and sodium ions are located in the calcium sites, inducing a favourable electrical charge balance. The carbonate ion can also replace both of PO$_4$$^{3-}$ and OH as AB-type CHAp Ca$_{10-x}$(PO$_4$)$_6$$x$(CO$_3$)$_3$(OH)$_2$$_2$(CO$_3$)$_3$ that found in many biological apatites (Garskaite et al., 2014).

The biocompatible, bio-active, and non-toxic character of CHAp makes the materials has highly suitable for dental-restoring materials (Okada & Matsumoto, 2015). In general, the manufacturing of synthetic CHAp has been conducted by using several methods: precipitation, sol-gel, hydrothermal, and mechanochemical to solid solution (Sianipar, 2016). The materials used in the manufacture of synthetic CHAp are generally natural substances containing high amount of calcium, such as fish bones, crab shells, shrimp shells, and so forth. Several publications have reported the use of those natural substances as a basic material for synthetic HAP production: clam shell (Sianipar, 2016), fish bone (Anggresani et al., 2021; Hanura et al., 2017), and shrimp shell (Sulistioso et al., 2012). To the best our knowledge there has no publication reported the Rebon shrimp (Acetes erythraeus) from Jambi province as a basic material for synthetic CHAp production.

Jambi province is the abundance of marine animals including the Rebon shrimp (Acetes erythraeus) which found in coastal areas such as in Tanjung Jabung Barat (Tungkal) and Tanjung Jabung Timur (Muaro Sabak) regencies. Rebon shrimp is a type of tiny shrimp whose length ranges from 1-3 cm with 69.45% of calcium (Permatasari et al., 2016; Sulistiyono et al., 2017). The high calcium content of Rebon shrimp that has been reported by Sulistiono and Permatasi can be used as a basic material for synthetic CHAp for candidate material in dental-restoring application. This work aims to synthesize the CHAP from Rebon shrimp as a candidate for dental-restoring application. The CHAP synthesis was conducted by the hydrothermal method with the variation of Ca/P ratio. The analysis of functional group, crystallinity, composition, surface morphology, and particle size of synthesized CHAp has been deeply discussed in this paper.

2. MATERIALS AND METHODS

Materials

The Rebon shrimp as the basic materials was obtained from local fisherman in Tanjung Jabung Barat (Tungkal) and Tanjung Jabung Timur (Muaro Sabak) regencies. The reagents used in this study were pro analytis (p.a) grade produced by Merck i.e. HNO$_3$ (aq) 65%, NH$_3$ (aq) 25%, and (NH$_4$)$_2$HPO$_4$ (aq). The CO$_2$ gas produced from PT. AGII, and distilled water produced from G-Force brand.

Instruments

The instruments analysis used were PANalytical Minipal 4 type X-Ray Fluorescence (XRF), Shimadzu Prestige 21 Fourier transform infrared (FTIR), Bruker D2 Phaser X-ray diffraction (XRD), and Phenom Pro X Scanning electron microscopy-energy dispersive X-Ray (SEM-EDX). The preparation tools used in this study were glassware, porcelain dishes, scales, analytical balance, sieve, autoclave, magnetic stirrer, furnace, desiccator, filter paper, aluminum foil, burette, static, clamp, blender, hot plate, pH meters, stirring rods, ovens and hydrothermal vessels.

Preparation of precipitated carbonate carbon (PCC)

A total of 20 g of dry 100 mesh sieved Rebon shrimp was calcined for 3 hours at a
temperature of 900 °C to produce the calcium oxide (CaO). The CaO powder was allowed to cool in the furnace for 24 hours. The powder was then weighed to get the yield. The preparation of PCC was conducted by CaO carbonation method. The process was carried out by dissolving 17 g of CaO in 300 mL of HNO$_3$ 2M. The mixture was stirred at 250 rpm of magnetic stirring for 30 min to get the filtrate. The filtrate was heated to 60 °C and the pH was set to 12 by carefully addition of NH$_4$OH 25%. The filtrate obtained at the process was flowed by CO$_2$ gas continuously until the pH becomes 8. The white solid obtained in this process was filtered and washed with distilled water until the pH is neutral. The dry solid of 100 mesh powder was labeled as PCC. The PCC powder was analyzed by XRF to determine the % elemental composition.

**Synthesis of CHAp**

The CHAp synthesis was performed by hydrothermal method by Hien et al. (2010) and Azis et al. (2015) through the mixing of PCC and (NH$_4$)$_2$HPO$_4$ saturated solution with a variation of Ca/P ratio of 1/1.61; 1/1.67; and 1/1.73. The mixture was dripped carefully by NH$_3$ (aq) 25% until pH 11 was reached. This mixture was put into the hydrothermal vessel at 120°C for 12 hours reaction time. The CHAp formed was separated from the rest of reactants and then washed by distilled water until the pH of filtrate was neutral. The synthesized CHAp was dried by 50 °C heating in oven to remove the water, sieved with 100 mesh sieves, and characterized by the FT-IR, XRD, and SEM.

### RESULTS AND DISCUSSION

**Characterization of PCC**

The synthesis of PCC was carried out using CaO powder as a calcination product of Rebon shrimp powder as basic materials. The equation for the reaction that occurs can be written as et al., 2015). Elemental analysis result of PCC by XRF (Figure 1) showed that PCC dominated by calcium (70.8%), phosphor (12.3%), and potassium (4.32%). The result was in line with the previous work by Sulistiyono et al. (2017).

Calcination: $2\text{CaCO}_3(s) + \text{heat} \rightarrow 2\text{CaO}(s) + \text{CO}_2(g)$ (1)

Hydration: $\text{CaO}(s) + 2\text{HNO}_3(l) \rightarrow 2\text{Ca(NO}_3)_2(aq) + \text{H}_2\text{O(l)}$ (2)

Precipitation: $\text{Ca(NO}_3)_2(aq) + 2\text{NH}_3(l) + 2\text{H}_2\text{O(l)} \rightarrow \text{Ca(OH)}_2(aq) + 2\text{NH}_4\text{NO}_3(aq)$ (3)

Precipitation: $\text{Ca(OH)}_2(aq) + \text{CO}_2(g) \rightarrow \text{CaCO}_3(s) + \text{H}_2\text{O(l)}$ (4)

![Figure 1. Elemental analysis of PCC by XRF of the synthesized PCC](image-url)
The functional group analysis of PCC can be seen in Figure 2. The IR spectrum of PCC in this work exhibits two characteristic carbonates bands: $\nu_2$ at 856 cm$^{-1}$ and $\nu_3$ at 1489 cm$^{-1}$ (Figure 1b). The wide peak at 3448 cm$^{-1}$ is assigned to O-H stretching of hydroxyl band. As already known, at undisturbed state, the VSEPR of carbonate ion is planar with three identic symmetrically C-O bonds, associating to the $D_{3h}$ point group which is three-fold rotational axis, a horizontal plane of symmetry and three vertical planes of symmetry at $60^\circ$ to one another. Among the regular vibrational form of planar carbonate ion, $\nu_2$ (C-O out-of-plane bending), $\nu_3$ (C-O asymmetric stretching), and $\nu_4$ (C-O planar bending) are IR active, while $\nu_1$ (C-O symmetric stretching), $\nu_3$, $\nu_4$ are Raman active (Ren et al., 2014).

XRD characterization was aimed to determine the type of crystal of the synthesized material. Calcium carbonate crystals (CaCO$_3$, PCC) have three types: calcite, aragonite and vaterite (Ren et al., 2020; Zou et al., 2017). Each crystal identified by different diffraction angles and Miller’s indexed. The crystal form of PCC in Figure 3 is a mixture of calcite and aragonite. From the diffractogram, the calcite phase showed at 20: 29.55º (104), 39.56º (113), and 43.10º (202) (Zou et al., 2017). While the aragonite phase was showed at 20: 26.19º (111), 27.19º (102), 33.01º (201), 36.05º (020), 37.82º (211), 45.71º (122), and 48.48º (220) (Elfina et al., 2020). Comparison of the peak to the International Center Data Diffraction (ICDD) No. 01-083-0577 for calcite No. 01-070-9854 for aragonite phase was presented and Table 1. The symbol $D$ (nm) is the crystallite size that determines using Scherer equation:

$$D = \frac{k\lambda}{\beta \cos \theta}$$

Where $\beta$ (rad) is the Full Width at Half Maximum (FWHM), $k$ is the Scherer constant (0.9-1.0), $\lambda$ is the wavelength of X-ray source from CuK$\alpha$ radiation (0.15406 nm), and $\theta$ (rad) is the peak position of Bragg angle.

Figure 4 shows the IR spectra of CHAp that have been synthesized by the hydrothermal with the various of Ca/P ratios. The band observed at 570 cm$^{-1}$ corresponds to the $\nu_4$ bending mode of the phosphate group. Similarly, the O-P-O bending of phosphate ($\nu_4$)
group is observed at 606 cm\(^{-1}\) (Rajkumar et al., 2011). The asymmetric stretching \(\nu_3\) mode of vibration P-O in phosphate bonds was assigned to the band at 1049 cm\(^{-1}\). The observed bands at 872 and 1427 cm\(^{-1}\) denote the HPO\(_4^{2-}\) in characteristic HA with deficient calcium refers to non-stoichiometric CHAp and the symmetric-asymmetric stretching modes of C-O bonds of carbonates groups, respectively (Lü et al., 2007; Youness et al., 2017). However, the peak at 872 cm\(^{-1}\) was also attributed as a carbonate band according other scholars (Gieroba et al., 2021; Reyes-Gasga et al., 2013; Shaltout et al., 2011). The broad peaks spread over the range between 3387-3435 and 1635 cm\(^{-1}\) reveals the superposition of absorption because of the stretching mode of surface hydroxyl groups and adsorbed water molecules (Basuki et al., 2021; Lak et al., 2008).

Table 1. The comparison of characteristic peak of PCC with the calcite and aragonite database

| 20 (°) | D (nm) | β (rad) | 20 | D (nm) | 20 | D (nm) | Crystal Phase | Miller index |
|-------|--------|---------|----|--------|----|--------|---------------|-------------|
| 26.19 | 22     | 0.3788  | -  | -      | 26.22 | 23     | aragonite     | 111         |
| 27.19 | 16     | 0.5112  | -  | -      | 27.22 | 22     | aragonite     | 102         |
| 29.55 | 16     | 0.5232  | 29.41 | 20   | -    | -      | calcite       | 104         |
| 33.01 | 16     | 0.5105  | -  | -      | 33.15 | 28     | aragonite     | 201         |
| 36.05 | 25     | 0.3369  | -  | -      | 36.17 | 21     | aragonite     | 200         |
| 37.82 | 34     | 0.2460  | -  | -      | 37.90 | 25     | aragonite     | 211         |
| 39.56 | 21     | 0.4097  | 39.41 | 25   | -    | -      | calcite       | 113         |
| 43.10 | 11     | 0.8200  | 43.16 | 24   | -    | -      | calcite       | 202         |
| 45.71 | 27     | 0.3179  | -  | -      | 45.86 | 24     | aragonite     | 122         |
| 48.48 | 24     | 0.3670  | -  | -      | 48.46 | 22     | aragonite     | 220         |

Table 2. The comparison of synthesized CHAp in this work with the CHAp in literature

| Material                        | Assignment (cm\(^{-1}\)) | References          |
|--------------------------------|--------------------------|---------------------|
| Standard HAp                   | 3570, 1620-1635, 1440-1470, 1040-1090, 960, 874, 603, 565 | Lü et al. (2007)    |
| HAp from human teeth           | 3569, 1633, 1089, 1045, 960, - | Lü et al. (2007)    |
| HAp powder                     | 3570, 633, 1043, - | Safari Gezaz et al. (2019) |
| HAp coated Ge crystal          | 3570, 1656, 1446, 1098, 962, - | Gieroba et al. (2021) |
| HAp from dental enamel         | 633, 1640, 1445, 1090-1032, 960, 870, 600-500 | Reyes-Gasga et al. (2013) |
| HAp from natural source        | 3567, 1645, 1446, 1035, 871, 617 | Shaltout et al. (2011) |
| HAp by mechano-chemical synthesis | 3570, 1620, 1470, 1055, 875, 605, 560 | Youness et al. (2017) |
| HAp from Rebon shrimp          | 3448, 1635, 1427, 1049, 872, 606, 570 | This work          |
Compared to the IR spectrum of CHAp, the carbonate band at 1489 cm\(^{-1}\) in PCC spectrum was shifted into 1427 cm\(^{-1}\) in CHAp spectra of all Ca/P ratios (Figure 4). In addition, the intensity at 1427 cm\(^{-1}\) increased with the increasing of Ca/P ratio. This might be attributed to substituting of CO\(_3^{2-}\) with PO\(_4^{3-}\) or OH in the apatite for charge compensation. In present work, the CO\(_3^{2-}\) peak can be observed in the spectra of synthesized CHAp samples in all Ca/P ratios. This is due to CO\(_3^{2-}\) caused from the rest of carbonate in PCC or from CO\(_2\) in air by calcination during synthesis process. It was also observed that the intensities of the phosphate band are increase with the increasing of the Ca/P ratio. The comparison of synthesized CHAp in this work with the CHAp in literature was listed in Table 2.

Figure 5. X-Ray diffraction pattern of the synthesized CHAp with 1.61; 1.67; and 1.73 Ca/P ratio

The XRD patterns of CHAp with the 1.61; 1.67; and 1.73 Ca/P ratios are shown in Figure 5. All the observed peak are readily indexed and compared with the reported value of hexagonal CHAp database (JCPDS file no. 09-0432). The main peaks indexed to (002) at about 26, broad and overlapped peaks of (210), (211), (300), (202) at 31-32, (310) and (311) at approximately 40, (222) at about 47, (320) at 50 and (004) at 54 indicate that the synthesized materials are CHAp (Aneem et al., 2019). After comparing with biological apatites from Aneem et al. (2019), it was concluded that the synthesized apatites resembled bone like apatite and indicates carbonate incorporation in the apatite crystal. There are no obvious peaks for the presence of other phases of CHAp that indicates the purity of the synthesized CHAp. The high intensity and broad width of the bands showed that the particles were highly crystalline and nano scale size, respectively.

The calculated lattice parameters are listed in Table 3. Average crystallite sizes (\(D\)) of samples with Ca/P ratios 1.61; 1.67; and 1.73 were calculated from XRD pattern by Scherer equation to be 26.3, 18.6, and 25.3, respectively. This shows that the higher Ca/P will relatively produce the smaller crystallite size. The study of effect of Ca/P ratio on the crystallite size by Syafaat & Yusuf (2018) reveal the similar result. On the other hand, the crystallinity of the samples was decrease with the increasing of Ca/P ratios (Table 3). Further, it also decreases in the micro strain (\(\varepsilon\)) and dislocation density (\(\delta\)). At constant pH, the crystal analyses showed that the higher Ca/P ratio will decrease the crystallinity (Mekmene et al., 2009).

Figure 5 shows the as-prepared CHAp micrograph is a spherical type of characteristic crystal carbonated apatites synthesized by hydrothermal procedures (Garskaite et al., 2014). Morphological results of as-prepared CHAp particles were consistent with the lattice parameter of XRD pattern in Table 3 (18.6-26.3 nm). All powder samples, Fig. 6(a–c), consist of agglomerates form which the shapes are identical. It is observed from particle size analysis by Origin® Software that the size of agglomerates is between 5 and 20 μm, and they were composed from fine particles 18-26 nm in size. It is evident that the microstructure of hydrothermally synthesized CHAp was formed through several stages, starting from crystallite structure of particles of 18–26 nm to fine sub-agglomerate particles up to 200 nm, which further cluster to give agglomerates 5–20 μm in size.

The elemental analysis of synthesized CHAp in all ratios was listed in Table 4. Based on the table, the composition of Ca/P ratios of synthesized CHAp was similar to the Ca/P ratio in early stages of CHAp synthesis. This result indicated that the all of carbonate ion in PCC was relatively react with the phosphate ion to form CHAp. Even though, the excess calcium (carbonate) compose the higher agglomerated CHAp.
Table 3. The calculated crystal size (D), crystallinity, micro strain (ε), and dislocation density (δ) of synthesized CHAp for different Ca/P ratios

| Ca/P ratio | D (nm) | Crystallinity (%) | ε (×10⁻³) | δ (×10⁻³ nm⁻²) |
|------------|--------|-------------------|------------|-----------------|
| 1.61       | 26.3   | 72.1              | 11.2       | 11.6            |
| 1.67       | 18.6   | 70.5              | 5.0        | 2.1             |
| 1.73       | 25.3   | 68.0              | 4.1        | 2.2             |

Figure 6. Characteristic microstructure by SEM of synthesized CHAp with Ca/P ratio 1.61 (a), 1.67 (b), and 1.73 (c)

Table 4. Elemental analysis of synthesized CHAp

| Elements         | Precursor CHAp ratios 1/1.61 (wt %) | Precursor CHAp ratios 1/1.67 (wt %) | Precursor CHAp ratios 1/1.73 (wt %) |
|------------------|------------------------------------|------------------------------------|------------------------------------|
| Calcium          | 8.58                               | 14.91                              | 6.35                               |
| Phosphorus       | 5.78                               | 8.99                               | 3.73                               |
| Oxygen           | 58.97                              | 53.35                              | 39.05                              |
| Carbon           | 25.86                              | 21.43                              | 50.18                              |
| Others           | 0.81                               | 1.32                               | 0.69                               |
| Synthesized CHAp | 1.49                               | 1.66                               | 1.71                               |

4. CONCLUSION
Calcium was found as the dominant element of Rebon shrimp (Acetes erythraeus) which are highly potential as the basic material of carbonated calcium hydroxyapatites. The present work investigates a detailed characterization of CHAp from Rebon shrimp (Acetes erythraeus). Spectroscopic characterization by FT-IR was assigned the characteristic CHAp band of hydroxide (3448, 1635 cm⁻¹), carbonate (872, 1427 cm⁻¹), and phosphate (1049, 606, 570 cm⁻¹). The XRD pattern assigned the all observed peak of CHAp are in good agreement compared to CHAp database (JCPDS file no. 09-0432) with the nano-scale size obtained by Scherer equation. The main peaks indexed to (002) at about 26, broad and overlapped peaks of (210), (211), (300), (202) at 31-32, (310) and (311) at approximately 40, (222) at about 47, (320) at 50 and (004) at 54 indicate that the synthesized materials are CHAp. It also observed that the high Ca/P ratio will decrease the crystallinity of CHAp. The as-prepared CHAp micrograph is agglomerates spherical form with size between 5-20 nm which build up from 18–26 nm crystallite particles. It is obviously evidenced that the Rebon shrimp can be used as a source of calcium for CHAp production which is a candidate material of bone or dental restoring candidate.

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