SYNTHESIS AND CHARACTERIZATION OF MAGNETIC-HYDROXYAPATITE COMPOSITE USING ONE-SPOT COPRECIPITATION METHOD

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Received: 23 September 2016 Revised: 15 December 2016 Accepted: 28 December 2016

ABSTRACT

SYNTHESIS AND CHARACTERIZATION OF MAGNETIC-HYDROXYAPATITE COMPOSITE USING ONE-SPOT COPRECIPITATION METHOD. Nanocomposite of magnetic-hydroxyapatite (MHAP) has the potential to be developed as a biocompatible-biodegradable material for bone cancer diagnosis and therapy. In this study, such composite materials have been successfully synthesized by one-spot coprecipitation method and ultrasonic dispersion. The mass ratio between magnetic and hydroxyapatite fraction were varied to 30:70 (K30), 40:60 (K40) and 50:50 (K50). X-Ray Diffractometer, Transmission Electron Microscope, Fourier Transform-Infrared Spectrometer and Vibrating Sample Magnetometer were used for characterizing the properties of MHAP composite. X-Ray Diffraction pattern reveals the presence of magnetic and HAP phases, confirm the establishment of MHAP composite system. TEM image and FT-IR spectra shows the spherical morphology of magnetic nanoparticles with a size of ~ 10 nm entrapped within HAP nanorod structure without any chemical bonding between the two phase. With such physical composite mechanism and higher energy induce by ultrasonic dispersion process, magnetic fraction of even 50% mass fraction could still be loaded into HAP matrix and provide maximum magnetisation value of 34.6 emu/g. This magnetization value is higher than the result of another study of MHAP synthesis, giving better prospect for bone cancer diagnosis and therapy.

Keywords: Composite, Magnetite, Maghemite, Hydroxyapatite, One-spot coprecipitation

SYNTESIS KOMPOSIT MAGNETIK-HIDROKSIAPATIT DENGAN METODE KO-PRESIPITASI SATU TAHAP DAN KARAKTERISASINYA. Nanokomposit magnetik-hidroksiapatit (MHAP) berpotensi dikembangkan sebagai bahan biocompatible-biodegradable untuk diagnosis dan terapi kanker tulang. Dalam studi ini, bahan komposit tersebut telah disintesis dengan metode ko-presipitasi satu tahap diikuti dengan proses dispersi ultrasonik. Perbandingan fraksi massa antara bahan magnetik dan hidroksiapatit diariaskan dengan komposisi 30:70 (K30), 40:60 (K40) dan 50:50 (K50). X-Ray Diffractometer (XRD), Transmission Electron Microscope (TEM), Fourier Transform-InfraRed (FT-IR) Spectrometer dan Vibrating Sample Magnetometer (VSM) digunakan untuk mengkarakterisasi sifat-sifat komposit MHAP. Pola difraksi sinar-X menunjukkan hadirnya fase magnetik dan HAP, mengkonfirmasi terbentuknya sistem komposit MHAP. Gambar TEM dan spektra FT-IR menunjukkan adanya nanopartikel berbentuk bulat dengan ukuran ~ 10 nm yang terperangkap di dalam struktur nanorod HAP tanpa adanya ikatan kimia di antara kedua fase tersebut. Dengan mekanik komposit fisis tersebut serta proses dispersi ultrasonik, 50% fraksi massa magnetik masih dapat dimuatan ke dalam matriks HAP dan menghasilkan nilai magnetisasi komposit hingga 34.6 emu/g. Nilai magnetisasi ini lebih tinggi dari hasil studi sintesis MHAP lainnya, memberi prospek yang lebih baik untuk diagnosis dan terapi kanker tulang.

Kata kunci: Komposit, Magnetit, Maghemit, Hidroksiapatit, One-spot coprecipitation

ABSTRAK

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INTRODUCTION

Hydroxyapatite (HAP) is widely known as one of material for bone regeneration. Having a chemical composition of Ca_{10}(PO_4)_{6}(OH)_{2}, which is similar to the composition of human bones, synthetic HAP has been used in healing or recovery of broken bones [1]. Recently, considerable research were carried out on combining HAP with magnetic nanoparticle [2-5]. Magnetic nanoparticle has been developed to be applied in some biomedical field due to its magnetic properties that can support diagnostic as well as therapeutic process. Usually the surface of nanoparticles must be modified first with organic or inorganic coating to get biocompatible system fit for such application [6]. Composite between HAP and magnetic nanoparticle (MHAP) is expected to give biocompatible ceramic magnetic materials which would be applied as an MRI contrast agent for diagnostic application as well as hyperthermia agent for therapy of bone cancer [4-5,7]. There are two general routes for synthesizing this composite, two-steps and one-step method. Two-steps method is carried out in which magnetic nanoparticle and HAP separately synthesized before mixing them to form composite. This synthesis route is performed using microemulsion [2], hydrothermal [4], mechanochemical process [5], and spray drying or spray pyrolysis technique [7,8]. All these processes usually result in the formation of hundreds size of composite particles having low magnetization value due to high fraction of HAP within the composite. Other route employing one-step precipitation process is accomplished by mixing all the precursors of magnetite and HAP at once followed by addition of strong base solution and some sequential treatment to establish the formation of MHAP precipitate [3]. This process, indeed, give higher magnetic saturation but still too low comparing to theoretical calculation due to low magnetization value of magnetic nanoparticle being precipitate within HAP matrix.

In this article, we report the synthesis of an MHAP nano-composite which has higher magnetization value. The synthesis was conducted in one-spot co-precipitation of magnetic and HAP fraction with different mass ratio followed by ultrasonic irradiation. Chemical and physical characteristics of the composite were thoroughly evaluated for understanding the mechanism inducing composite characteristics.

EXPERIMENTAL METHOD

Materials and Equipment

All chemicals are used as received without special treatment. FeCl_3·6H_2O, FeCl_2·4H_2O, Ca(NO_3)_2·4H_2O, (NH_4)_2HPO_4, and glacial acetic acid was obtained from Merck (Germany). Another materials used are NH_4OH, nitrogen gas (UHP), NaOH and Deionized Water. The equipment for mixing and homogenizing are ultrasonic probe “Sonics” (20 kHz, 40 watts) and IKA Eurostar Mixer. Other equipment used is glassware, stirrer and ovens.

MHAP Composite Preparation

MHAP composite was synthesized by following the method of Nguyen VC et al [3] with slight modification using ultrasonic irradiation instead of mechanical stirring. The samples were prepared with magnetic to HAP mass ratio of 30:70, 40:60 and 50:50 which denoted as K30, K40 and K50 samples respectively. FeCl_3·6H_2O and FeCl_2·4H_2O were dissolved in deionized water which has been aerated with nitrogen gas. Some volume of 25% NH_4OH solution was added to these salts solution and dispersed with ultrasonic probe for 5 minutes. Ca(NO_3)_2·4H_2O and (NH_4)_2HPO_4 were dissolved in deionized water and added slowly to the previous solution. The pH value of the mixture was adjusted to 11 and continued with ultrasonic irradiating process for another 5 minutes. The solution was then heated at 90 °C for 2 hours while stirring using a mechanical mixer. The black solution was allowed to cool to room temperature, and then allowed to further precipitate for 24 hours without stirring. The precipitate was then separated using permanent magnet and washed with deionized water until it reached a neutral pH. After drying in an oven at a temperature of 90 °C for 1 hour, the precipitate was pulverized using a mortar. MHAP fine powder obtained were then characterized to determine its properties.

Characterization

Characterizations were carried out to determine the phases, magnetic properties, chemical bonding formed between the components of MHAP composites and their morphology. The phases were analyzed from diffraction pattern collected using PANalytical X-Ray Diffractometer with Cu radiation source (λ = 0.154056 nm) in the range of diffraction angles, 2θ 10°- 90°. Detail phase and structure analysis were carried out using Rietan 2000 program [9]. Magnetic properties were analyzed from the hysteresis curve measured using Vibrating Sample Magnetometer (VSM) OXFORD 1.2H. Measurements were performed at room temperature with an external magnetic field up to 1 Tesla. Analysis of chemical interactions between the components of the composite was performed using Bruker FTIR spectrophotometer Tensor 27 equipped with ATR system. These three facilities are installed at Centre for Science and Technology of Advanced Material (PSTBM), BATAN. Morphology of MHAP composite was observed and analyzed using Transmission Electron Microscope (TEM) JEOL JEM 1400 at Chemistry Departement of Universitas Gadjah Mada.
RESULT AND DISCUSSION

XRD Analysis

MHAP sample diffraction pattern in Figure 1, shows the formation of Fe₃O₄ (magnetite) phase, identified from the presence of peaks at an angle of 30.30°, 35.70°, 43.40°, 57.40°, and 63.10°, fit with the data on crystallography database (JCPDS ICDD 19-629) for Fe₃O₄[10].

The peaks are broadened which indicates the formation of nanocrystalline structure. These peaks could also be identified as the peak of maghemite, γ-Fe₂O₃ (JCPDS-ICDD 39-1346) [10] for these two phases are isomorphous. This second phase is very likely to occur because the magnetite can be oxidized to maghemite on MHAP synthesis process, especially during the heating process, following chemical reaction of:

$$4 \text{Fe}_3\text{O}_4 + \text{O}_2 \rightarrow 6 \gamma \text{Fe}_2\text{O}_3$$

Hydroxyapatite (HAP) phase is identified by the appearance of HAP typical peaks at an angle of 25.95°, 31.71°, 32.19°, 32.90°, 39.47°, 46.70°, 49.50° and 53.25° fit with the data on JCPDS (9-432). Peaks of HAP showed normal peak to peak ratio or did not show any preferred orientation, indicate the formation of HAP of granular or short-rod structure [11]. Figure 2 shows that HAP peaks also broadened and there is no clear splitting of (211), (112) and (300) peaks in the range of 31° to 33° diffraction angle which indicates a low degree of HAP crystallinity. This low crystallinity is analyzed due to the HAP precipitation is accomplish only by strong base solution environment [12], followed by low temperature heating at 90 °C, simultaneously with the precipitation of magnetic nanoparticles. Increasing magnetic fraction results in lowering intensity and more broaden HAP peaks, hence more amorphous HAP. This condition is related once again with the fact of only strong base induce precipitation. Higher magnetic fraction will consume more base solution and leaving only a small volume of base for HAP precipitation and consequently will lower HAP degree of crystallinity.

For magnetic phase (see Figure 2), more detail observation on (311) principal peak of magnetite phase at 35.70°, it can be seen that no significant change of peak broadening with increasing magnetic fraction, ensuring nearly the same crystallite size of magnetite nanoparticle for all composition. The crystallite size is about 5 nm, estimated using Debye-Scherrer equation.

These magnetic and HAP phases are bound to each other forming a composite of MHAP. This conclusion comes from the fact that the sample was obtained as a result of washing and separation process using permanent magnets.

If these phases are not bound to each other, then HAP will not be attracted by magnet and will be left at the solution during the washing step, leaving only magnetic phase powder as a synthesis product.

Further analysis on the patterns were carried out using Rietveld analysis method implemented on Rietveld analysis method implemented on Rietveld analysis method implemented on Rietveld analysis method implemented on RIETAN2000 program [9] and HAP base structure parameter from article written by Petr Ptacek [13]. Figure 3 shows refined pattern of each composite sample, assuming the existence of three phases of Fe₃O₄, γ-Fe₂O₃ and HAP. The refined patterns were well fitted with the observation patterns and give the reliability factor, $R_{wp}$ ~ 2 and goodness of fit, $S$ ~ 1.5. Detail composition and refined parameter results are presented in Table 1.

There is slight changing of lattice parameter of Fe₃O₄/γ-Fe₂O₃ to smaller value compared to their initial lattice parameter of ~ 8.4 Å while HAP lattice parameter show no changing with varying composition. It could be analyzed that magnetic nanoparticles growth were slightly inhibited by surrounding HAP phase and result in suppressing magnetic lattice. Refined composition tend to deviate from their initial value give non-linear relationships with increasing magnetic fraction analyzed due to unhomogenized mixing process.

![Image](image1.png)

**Figure 1.** X-ray diffraction pattern of magnetic-hydroxyapatite composites at different composition: (a). K30; (b). K40 and (c). K50.

![Image](image2.png)

**Figure 2.** Detail diffraction pattern of HAP (30.5° - 33.5°) and Fe₃O₄/γ-Fe₂O₃ (34° - 37°) principal peaks for (a). K30; (b). K40 and (c). K50 samples.
Sample of K40 shows lower magnetic fraction with high maghemite fraction which is unfavourable for high magnetic properties establishment.

**TEM imaging and Particle Size Analysis**

Morphology of K50 composite observed by TEM shown in Figure 4 confirm analyses result of X-ray diffraction pattern. Figure 4(a) displays the morphology of irregular shape composite having ~ 100 nm - 200 nm in size. More detail picture in Figure 4(b) shows the spherical morphology of magnetic nanoparticles (dark image) with a size of ~ 10 nm trapped within HAP needle like structure (gray image) and surrounding by the short rod of HAP structure (Figure 4(c) and 4(d)). This morphological picture support the synergetic growth mechanisms between magnetic nanoparticle and HAP nanorod structure, which are bound to one another forming nano-composite system. This result is in agreement with other study of MHAP structures using energy filter imaging and electron tomography which showed that HAP nanoparticles tend to have a needle structure for the one interact directly with the surface of magnetic nanoparticle, while the surrounding HAP is more likely to form a rod structure [14].

The images also showed the more separate nanoparticles dispersed within HAP matrix, which is analyzed due to higher energy induce by ultrasonic process compared to the mechanical mixing.

This high energy and more monodisperse magnetic nanoparticle is assumed being responsible for higher fraction of magnetic nanoparticle as high as 50% that could be incorporated within HAP matrix. Other experiment using hydrothermal process stated that mass fraction of only 30% could maximally be entrapped [4].

**FT-IR Spectrum Analysis**

Chemical bond presence in MHAP samples are analysed from their infrared spectrum as shown in Figure 5 measured using ATR technique. In MHAP spectrum there are absorption bands at 1452, 1410, which characterize the carbonate functional group. An
absorption band at 3525 cm\(^{-1}\) is produced by hydroxyl functional groups on MHAP and wave numbers 1033, 572, and 571 cm\(^{-1}\) in accordance with the functional groups on the HAP PO\(_4^{3-}\) [15]. Absorption band typical for Fe-O bond also appears at wave number 570 cm\(^{-1}\). The existence of this typical band of HAP and magnetite at MHAP spectra, revealed the formation of composite of magnetite-HAP. However, there is no chemical bond between magnetite and HAP phase, which assuring the more physical interaction between the phases.

**Magnetic Properties Analysis**

The most important properties of these composites are their magnetic properties. These properties will be related to the ability of the composite to become an agents of hyperthermia therapy as well as an MRI contrast agent. Figure 6 displays magnetic hysteresis curve of MHAP composites measured at room temperature with a maximum external magnetic field of 1 Tesla. In general, MHAP samples show superparamagnetic properties, which usually obtained for Fe\(_3\)O\(_4\) nanoparticle with crystallite size below 20 nm [16].

The saturation magnetization, \(M_s\), is the maximum value of magnetization that can be achieved when all the magnetic moments aligned in one specific direction and will have the value of 92 emu/gram for bulk and pure Fe\(_3\)O\(_4\) and 76 emu/gram for \(\gamma\)-Fe\(_2\)O\(_3\) [17]. For MHAP K50, with real mass composition of 53.3% for magnetic fraction as discussed in (3.1) section, theoretically the \(M_s\) value should be in the range of (40.5 – 49.04) emu/gram. However, the measurement result give only 34.6 emu/gram. Referring to the discussion of phase and morphology above, this decrease could be explain due to the existence of two-phase mixture of Fe\(_3\)O\(_4\) and \(\gamma\)-Fe\(_2\)O\(_3\) phase and low degree of crystallinity of magnetic nanoparticle produce in this process. This fact also could be used to explain non-linear behaviour of magnetization for K40 sample as displayed in Figure 6.

The presence of HAP phase between the magnetic nano powders could also reduce magnetic interaction between iron oxide. When the magnetic nanoparticles bound firmly in the HAP rod system, it will reduce the freedom of magnetic nanoparticles to rotate and respond to any external magnetic field given to reach saturation value [18].

This K50 MHAP \(M_s\) values are still higher than the results of the same process but using mechanical stirrer [3], which only reach ~ 40 emu/gram for pure magnetic powder and would provide much lower magnetic value for MHAP. It can be assumed that ultrasonic irradiation used in this work provide faster precipitation of Fe\(_3\)O\(_4\) nanoparticles and inhibit further oxidation to \(\gamma\)-Fe\(_2\)O\(_3\). Consequently, this will result, not only magnetic nanoparticles with smaller size and homogeneous, but also enhancement of magnetic phase formation. However, other research result show that too high ultrasonic frequency and power (35 kHz, 320 W) will significantly decrease the magnetization to the value of only 7.40 emu/gram which is analyzed due to the magnetic domain of the particle hindered by molecular binding with HAP [18]. Similarly, spray pyrolysis process, producing only a composite MHAP with a value of magnetization of 11.8 emu/gram for 36.4% magnetic mass fraction [7] or about 35 emu/gram of pure magnetite sample. Even for higher ratio between magnetite and HAP of 3.2, the only maximum of 30 emu/gram could be obtained by other spray pyrolysis process [8].

**CONCLUSION**

High magnetization nanocomposite of magnetic-hydroxyapatite (MHAP) has been successfully synthesized by one-spot co-precipitation method and ultrasonic dispersion. Fe\(_3\)O\(_4\) /\(\gamma\)-Fe\(_2\)O\(_3\) magnetic nanoparticles with a size of ~ 10 nm entrapped within HAP nanorod structure without any chemical bonding between the two phases. With such physical composite mechanism and higher energy induce by ultrasonic dispersion process, magnetic fraction of even 50% mass fraction could still be loaded into HAP matrix and provide maximum magnetization value of 34.6 emu/gram. It can
be concluded that MHAP produced in this study has a magnetization value higher than the synthesis of other similar studies and will have a prospect to produce better contrast in MRI image and desired heat for hyperthermia therapy.

ACKNOWLEDGEMENTS

This work was supported by the DIPA of PSTBM-BATAN at fiscal year of 2015. The authors would like to thank PSTBM-BATAN management for providing this fund.

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