FABRICATION OF POLY(LACTIC-CO-GLYCOLIC ACID)_CALCIUM PHOSPHATE

by A Kafrawi Nasution

Submission date: 17-Nov-2020 04:22PM (UTC+0800)

Submission ID: 1448799529

File name: POSITE__SYNTHESIZATION_OF_CALCIUM_PHOSPHATE_FROM_CRAB_SHELLS.pdf (550.34K)

Word count: 4296

Character count: 23200

FABRICATION OF POLY(LACTIC-CO-GLYCOLIC ACID)/CALCIUM PHOSPHATE BONE CEMENT COMPOSITE: SYNTHESIZATION OF CALCIUM PHOSPHATE FROM CRAB SHELLS

Mohammad Redzuan Abdul Hanan^a, Ahmad Kafrawi Nasution^b, Rafaqat Hussain^c, Syafiqah Saidin^{a,d*}

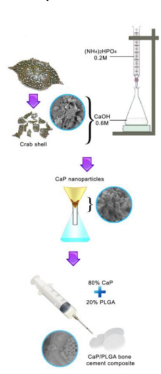
^aFaculty of Biosciences and Medical Engineering, Universiti Teknologi Malaysia, 81310 UTM Johor Bahru, Johor, Malaysia ^bDept. of Mechanical Engineering, Faculty of Engineering, Muhammadiyah University of Riau, Pekanbaru, Indonesia ^cDepartment of Physics, COMSATS Institute of Information Technology, Islamabad, Pakistan

JN-UTM Cardio Centre, Universiti Teknologi Malaysia, 81310 UTM Johor Bahru, Johor, Malaysia

Article history
Received
24 August 2017
Received in revised form
6 February 2018
Accepted
15 February 2018
Published online
3 June 2018

*Corresponding author syafiqahs@utm.my

Graphical abstract



Abstract

Crab shells are waste product, rich with calcium compound. Calcium element is often used as a naterial for bone scaffold due to its bioactive and biodegradation properties. In this tudy, calcium phosphate (CaP) nanoparticles were synthesized from crab shells through a wet chemical route. The CaP nanoparticles were then Intered and mixed with poly(lactic-co-glycolic acid) (PLGA) to form a bone cement composite. The mixture was casted in a cylinder shape and it was characterized through ATR-FTIR, XRD, FESEM, contact angle and DSC analyses. The CaP pellet and the CaP/PLGA bone cement composite were then subjected to in vitro simulated body fluid (SBF) bioactivity test. The CaP/PLGA bone cement composite was found to have <mark>a composition of</mark> crystal <mark>CaP and PLGA</mark> with a tolerable glass transition state, suitable to be used in a physiological environment. The CaP nanoparticles were agglomerated on the 3D interconnected surface of PLGA. The hydrophobicity of the CaP was increased (66.94%) with the addition of PLGA as a binder matrix where this composite has induced the formation of apatite layer. This bioactive property is crucial in fabricating a bone substitute material as it can promotes cell penetration, attachment and proliferation.

Keywords: Bone cement, calcium phosphate, poly(lactic-co-glycolic acid), crab shells, bioactive

Abstrak

Cangkerang ketam ialah produk buangan, kaya dengan kompaun kalsium. Elemen kalsium sering digunakan sebagai bahan perencah tulang kerana sifat bioaktif dan bioterurai. Dalam kajian ini, nanopartikel kalsium fosfat (CaP) disintesis daripada cangkerang ketam melalui jalan kimia basah. Nanopartikel CaP kemudiannya dibakar dan dicampur dengan asid poli(laktik-ko-glikolik) (PLGA) untuk menghasilkan komposit simen tulang. Campuran dihasilkan dalam bentuk silinder dan ia dicirikan melalui analisis ATR-FTIR, XRD, FESEM, sudut sentuhan dan DSC. Pelet Ca/P dan komposit simen tulang CaP/PLGA kemudiannya melalui ujian bioaktif cecair simulasi badan secara in vitro. Komposit simen tulang CaP/PLGA mempunyai komposisi CaP kristal dan PLGA dengan toleransi tahap perubahan kaca, sesuai untuk digunakan

dalam persekitaran fisiologi. Nanopartikel CaP berkumpul pada permukaan tiga dimensi hubungan antara PLGA. Sifat fobia air CaP bertambah (66.94%) dengan penambahan PLGA sebagai pengikat matrik di mana komposit ini telah menumbuhkan lapisan apatit. Sifat bioaktif ini penting dalam pembuatan bahan tulang di mana ia dapat menggalakkan penembusan, pelekatan dan pertambhan sel.

Kata kunci: Simen tulang, kalsium fosfat, asid poli(laktik-ko-glikolik), cangkerang ketam, bioaktif.

© 2018 Penerbit UTM Press. All rights reserved

1.0 INTRODUCTION

Crab is classified as crustacean group which has a thick exoskeleton shell [1]. This creature lives in seawater, freshwater and partly on land [1]. Portunus pelagicus or also called "flower crab" in East Asian countries [2], is commonly available at seafood markets and restaurants to be served as one of the main seafood dishes [3]. Crab shell is a waste product as it cannot be eaten and does not give advantage to its collector. Thus, it is abundantly accessible and available without any specific use.

Crab shell is one of the natural resources, rich with calcium carbonate (CaCO₃) [4]. Extraction of CaCO₃ from natural resources is beneficial as an alternative option to synthesize bioceramic material [5]. Bioceramic material derived from natural resources could provide better metabolic activity and active response of physiological environment compared to a synthetic CaP [6]. Several CaP derivatives such as hydroxyapatite $Ca_{10}(PO_4)_6(OH)_2$, biphasic calcium phosphate (BCP) and tricalcium phosphate (TCP, Ca₃(PO₄)₂) which obtained from natural resources, are actively being used in dentistry and orthopedic applications [5, 7-8] as well as in bone reconstructive surgery [9].

Calcium phosphates are also being utilized as synthetic bone graft substitutes [10] and being incorporated in a bone implant coating [11] due to its biocompatible, bioactive, biodegradable and osteoconductive properties. The emergence of CaP from natural resources that has similar structure and morphology with human bones [12] provides benefit in the treatment of bone disease and for regeneration of damaged organs and tissues. Besides, CaP can be produced in large quantities at relatively low cost [5]. The synthesization of CaP from waste product will further cut the cost of CaP production.

From the last four decades, several efforts have been conducted in developing CaP bone cement including the study on various size of CaP granules to cover the size of bone defect [13]. Furthermore, the viscosity of injectable CaP bone cement has also been optimized to tolerate the grafting and filling of missing bone [14-16], either in the form of cement or paste by adjusting the wettability property of the

materials [17-19]. Another strategy to enhance the property of CaP bone cement is, by incorporating a degradable polymer such as poly(lactic-co-glycolic acid) (PLGA), polylactic acid (PLA) and polyetherketoneketone (PEKK) which acts as a binder to the cement [20].

Therefore, the aim of this study was to synthesize CaP from crab shells and to incorporate the synthesized CaP into PLGA to construct a bone cement composite. The samples were characterized through several characterization analyses including attenuated total reflectance-Fourier transform infrared spectroscopy (ATR-FTIR), X-ray diffractometer (XRD), field emission scanning electron microscopy (FESEM), wettability and differential scanning calorimetry (DSC) analyses. Finally, the bioactivity of the CaP and the CaP/PLGA bone cement composite were investigated through in vitro simulated body fluid (SBF) bioactivity test by visualizing the formation of apatite layer under SEM.

2.0 METHODOLOGY

2.1 Sample Preparation

Crab (Portunus pelagicus) shells were collected from local seafood restaurants in the area of Johor Bahru, Johor, Malaysia. The shells were cleaned and washed to remove macro residues. They were crushed into small pieces using mortar and pestle, treated in an ultrasonic bath (LIR Biotech™, Malaysia) for 15 minutes at room temperature to remove micro residues and dried in an oven (UN30, Memmert Gmbh, Germany), overnight. The treated shells were then immersed in 1.0 M NaOH at 80°C for 1 hour under stirring condition to denature the protein and to deacetylation the chitin and chitosan [7]. The shells were rinsed twice with deionized water and dried in an oven (UN30, Memmert Gmbh, Germany) for 6 hours at 110°C [12]. It was denoted as CaCO₃ powders. The CaCO₃ powders were then calcined at 1000°C in a muffle furnace (BSK-1700-X-S, MTI Corp, USA) for 5 hours to form CaO [12] and further crushed with mortar and pestle into a powder form.

The procedures to synthesize CaP nanoparticles were conducted according to Unabia et al. [21] with

slight modification. The CaO powders were dissolved in distilled water at 40°C under stirring condition to form calcium hydroxide (CaOH). A total of 100 mL of 0.2 M diaremonium phosphate ((NH4)2HPO4) was titrated into 100 mL of 0.6 M CaOH where the titration parameters were set at 40°C and pH 9, under stirring condition. The CaP mixture was then continuously stirred at 40°C for another 4 hours. The homogenized CaP mixture was leaved at room temperature for 24 hours to allow a complete reaction and precipitation. The filtration of CaP was done using a filter paper, followed by a drying process at 110°C for 5 hours and sintering process in a muffle fumace (BSK-1700-X-5, MTI Corp, USA) at 1000°C for 4 hours.

Finally, a bone cement composite was prepared with a composition of 20 wt% PLGA and 80 wt% CaP. A total of 0.2 g of PLGA was dissolved in 0.5 mL of dimethyl sulfoxide (DMSO). The PLGA solution was then added into 0.8 g of CaP nanoparticles under vigorous agitation. Finally, the CaP/PLGA mixture was injected, molded and left to solidify at room temperature.

2.2 Surface Characterization

The ATR-FTIR (Nicolet iS5-IR Spectrometer, Thermo Scientific, USA) was used overify the chemical composition of the samples at a frequency interval between 500 cm⁻¹ and 1000 cm⁻¹. The number of scanning and resolution of 36 and 4 were used, respectively, to analyze the samples. The chemical composition of the samples was further clarified by XRD instrument (D5000, Siemens, Germany). The analysis was performed at 40 kV and 40 mA to recognize the crystallographic phases of the samples and the size of crystal particles. The data were recorded in 20 range from 5° to 60° using CuKa radiation of 1.5406 Å.

The structural morphology of the samples was examined by FESEM (SU8020, HITACHI, Japan) at 5000×, 10000× and 30000× magnifications. Each sample was sputter coated with an ultrathin platinum film (Q150R, Quorum Technologies, England) for better image visualization. The FESEM images were then subjected to a mapping analysis using Image J software (NIH, MD, USA) to determine the particle size

For the water contact angle measurement, a pellet compressor (E-Z Press, International Crystal Laboratories, USA) was used to compress 1.0 g of CaP powders into a pellet form (diameter = 1.4 cm; thickness = 0.3 cm). The compression was run for 60 seconds at 1000 Psi. While, a sample of CaP/PLGA bone cement composite with a diameter of 1.4 cm and a thickness of 0.3 cm was also prepared for the same purpose. The surface wettability of the samples was then investigated by using a video contact angle instrument (VCA, Optima, AST products, Inc., USA) by dropping 0.5 mL deionized water on the sample surfaces. The angle of water drop was recorded within 2 seconds at three difference spots to obtain an average value.

2.3 Thermal Characterization

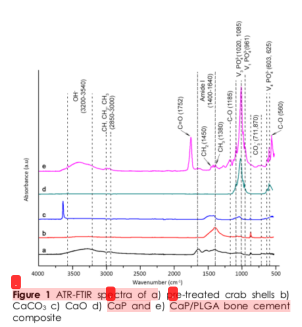
The thermal characteristic of the CaP/PLGA bone cement composite was analyzed using DSC (DSC 2000, Perkin Elmer, US) to determine the transition glass temperature, Tg, of the bone cement composite. A heat/cool/reheat method was used to remove the thermal history. The heating temperature was set between 24°C and 180°C with a rate of 5°C/min under nitrogen atmosphere. The reference pan with a cover was let empty during the scanning process.

2.4 In Vitro SBF Bioactivity Test

The bioactivity of the CaP/PLGA bone cement composite and the CaP pellet was identified through in vitro simulated body fluid (SBF) bioactivity test to simulate an apatite formation. The preparation of SBF solution was referred based on the work of Kokubo and Takadama [22]. The CaP/PLGA bone cement composite and the CaP pellet were immersed in the SBF solution at pH 7.4 for 1, 3 and 7 days by maintaining the environment temperature at 36.5°C in an ambient shaker incubator (SI-50D, Protech®, Malaysia). The SBF solution was refreshed every 24 hours to provide adequate ions for apatite nucleation. The samples were collected following the time-points, dried and coated with a thin gold film using a gold sputter coater (EM CPD300, LEICA, Germany) for the visualization under SEM (SEM TM3000, HITACHI, Japan). The images were captured at 100×, 500× and 1000× magnifications.

3.0 RESULTS AND DISCUSSION

Calcium phosphate is biologically active to enhance osseointegration. It has been used in the fabrication of clinical total hip implants for cell survivability [23], to overcome the loosening of pin resident of an external fixator [24], to allow earlier weight bearing after tibial plateau fractures [25], etc. In this study, the CaP nanoparticles were synthesized through a chemical route by utilizing the extraction of CaCO₃ from crab shells. Most CaPs are demonstrating burst degradation. Therefore, this property was improved through the incorporation of PLGA polymer. Figure 1 shows the ATR-FTTR spectra of pre-treated crab shells, CaCO₃, CaO, CaP and CaP/PLGA bone cement composite. The pre-treated crab shells demonstrated the existence of several peaks of primary amine at $1\,400$ - 1640 cm $^{\!-1}.$ The process to produce CaCO $_3$ has removed one the amine peaks at 1640 cm⁻¹, signifying the detachment of some proteins from the shells. The CO₃²⁻ peaks were clearly visible at 711 and 870 cm⁻¹ in the CaCO₃ spectrum, presenting the component of carbonate in the powders [26]. While, high intensity peak of crystal OH- in the CaO spectrum showed the calcification of hydroxyl compound after the sintering process.



The synthesization of CaP led to the removal of amine groups and to the appearance of phosphate peaks, V_1 PO₄3- (961 cm⁻¹), V_3 PO₄3- (1010 cm⁻¹ and 1085 cm⁻¹) and V_4 PO₄³⁻ (603 cm⁻¹ and 625 cm⁻¹). Those peaks were accessible in the CaP powders and the CaP/PLGA bone cement composite. The sharp peaks of V_3 PO₄³⁻ at 1010 cm⁻¹ and 1085 cm⁻¹ were pre-identified as TTCP and β-TCP, respectively [27]. While, the V_1 PO_4^{3-} and the V_4 PO_4^{3-} peaks resembled HA formation [27]. Those CaP compounds were further clarified by the XRD analyses. The incorporation of PLGA inside the bone cement composite also led to the detection of several PLGA functional groups including CH, CH2, CH3 stretching vibrations in between of 2850 and 3000 cm⁻¹ [28]. Besides, weak peaks of CH2 and CH3 were observed at 1450 cm⁻¹ and 1380 cm⁻¹, respectively. Another two peaks at 560 cm⁻¹ and 1185 cm⁻¹ representing the C-O functional groups and one peak at 1750 cm⁻¹ representing the carbonyl C=O functional group were also noticed in the spectra [29].

The XRD pattern in Figure 2 shows the mapping of HA and β -TCP following the Joint Committee on Powder Diffraction Standards (JCPDS) datasheets (HA: JCPDS 09-0432 and β -TCP: JCPDS 09-0169). Three main peaks of HA were found at 31.75°, 32.10° and 32.85° with the crystal lattice of (211), (112) and (300). While three main peaks of β -TCP were noticed at 25.90°, 28.00° and 31.25° with the crystal lattice of (1010), (214) and (0210). The crystallite size of the CaP powders was calculated to be 55.06 nm using Scherrer formula. The PLGA in the bone cement composite was not identified in the XRD pattern due to its non-crystallize property.

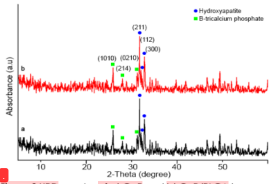


Figure 2 XRD spectra of a) CaP and b) CaP/PLGA bone cement composite

A 3D interconnected macro-porous structure of CaP is an ideal and essential structure for bone graft material [30]. A porous design is able to enhance mechanical interconnection between implanted scaffolds and adjacent tissues, thus providing better mechanical stability [31]. Figure 3 displays the morphology of pre-treated crab shell, CaCO₃, CaO, CaP and CaP/PLGA bone cement composite. A fiber-like shape was visible on the crab shell powders that hold the particles. After the NaOH treatment, the CaCO₃ powders demonstrated no fiber-like shape as the treatment process denatured most proteins that attached on the crab shells. This result was supported by the FTIR analysis in Figure 1 where the amine peak at 1640 cm⁻¹ was disappeared from the CaCO₃ spectrum.

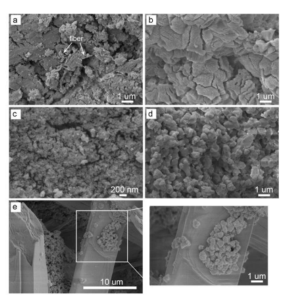


Figure 3 Surface morphologies of a) crab shells b) CaCO₃ c) CaO d) CaP and e) CaP/PLGA bone cement composite

The sintering process that transformed the CaCO₃ into the CaO has shrunk the CaO powders to be finer nanoparticles with an average diameter of 40.51±15.28 nm. While in Figure 3(d), the CaP nanoparticles were obtained with an average size of 298.99±123.05 nm. However, this range of size was bigger than the calculated value in the XRD analysis due to the gold sputtering process that united some of the CaP nanoparticles, cause the visualization of bigger compound with formation of tiny bridge between the nanoparticles. An interconnection of elongated rods-like shape of PLGA with an agglomeration of CaP nanoparticles was then noticed on the CaP/PLGA bone cement composite (Figure 3(e) and (f)). This structure is beneficial for the degradation of bone cement that allowing cell penetration and attachment.

The wettability measurement on the CaP pellet showed a classifying of hydrophilicity property (16.63±3.00°). The hydrophobicity of the CaP was increased to 50.30±4.78° after the incorporation of PLGA into the CaP to produce the CaP/PLGA bone cement composite. The increment in hydrophobicity property is important to control the degradation of CaP/PLGA bone cement composite by restricting the penetration of water through the bone cement composite.

The glass transition, T_g of the CaP/PLGA bone cement composite was measured above the physiological temperature as shown in Figure 4. The quantization of T_g is needed to ensure the T_g of PLGA is not shifted to a temperature lower than the temperature of human body. The CaP/PLGA bone cement composite which composed of PLGA, was soft and viscoelastic [32]. This property was steeply transformed into a glassy state when the temperature reached 45.15°C. The bone cement composite was then gradually become harder until it reached the T_g peak of 103.38°C. Heat treatment more than the T_g peak, drastically caused the bone cement composite to melt into a liquid phase.

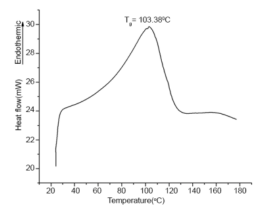


Figure 4 Glass transition, Tg, graph of CaP/PLGA bone cement composite

In spite of the physico-chemical properties, the prerequisite analysis of *in vitro* SBF bioactivity test is necessary to predict cells attachment and bone generation after the grafting and restoration surgery. Figure 5 shows the morphology of apatite formation on CaP pellet and CaP/PLGA bone cement composite at day 1, 3 and 7 of incubation. Within 7 days of incubation, abundant uniform apatite agglomeration were formed on the surfaces of CaP and CaP/PLGA bone cement composite, clearly visible at 1000× magnification.

A spider web-like apatite (Figure 5 (b, d, f, and h)) was observed on the CaP/PLGA bone cement composite. The degradation of the PLGA from the bone cement composite produced the formation of micro-pores that allowed the penetration of SBF ions through the pores for the nucleation of apatite. In a physiological environment, the pores will be useful for water and minerals intake, in need for cell attachment and nourishment [30]. At day 7, thicker apatite layer was formed on the CaP/PLGA bone cement composite as the apatite covered the hole of the micro-pores (Figure 5(f and h)). In comparison with the apatite formation on the CaP pellet, there was no appearance of micro-pores on the CaP surfaces due to non-incorporation of PLGA.

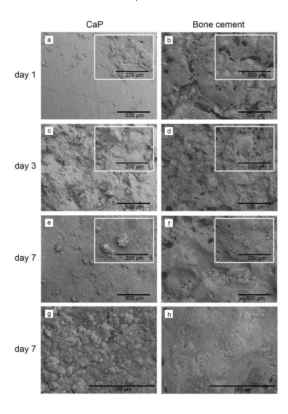


Figure 5 SEM images of apatite layer on CaP pellet and CaP/PLGA bone cement composite after *in vitro* SBF bioactivity test at day 1 (a and b), day 3 (c and d) and day 7 (e, f, g and h)

4.0 CONCLUSION

The recycling and reusable of crab shells to be used in the synthesization of CaP is one of the practical approaches to overcome the problem of seafood waste management. The CaP derived from crab shells has a bioactive property which is beneficial for bone cement formulation. The incorporation of PLGA into the CaP nanoparticles has displayed a morphology of 3D interconnected rod-like-structure with an agglomeration of CaP nanoparticles. It has also increased the hydrophobicity of the bone cement composite with a tolerate apatite formation.

Acknowledgement

The authors would like to declare that there is no conflict of interest and no ethical approval is required. This study was funded by Malaysian Ministry of Higher Education through Research University Grant (RUG) Tier 2 [Q.J130000.2645.11J68].

References

- Asphama, A. I., Amir, F., Malina, A. C., and Fujaya, Y. 2015. Habitat Preferences of Blue Swimming Crab (Portunus Pelagicus) Species Complex. Aquacultura Indonesiana. 16: 10-15.
- [2] Ikhwanuddin, M., Liyana, A. N., Azra, M. N., Bachok, Z., and Abol-Munafi, A. B. 2014. Natural Diet of Blue Swimming Crab, portunus pelagicus at Strait of Tebrau, Johor, Malaysia. Sains Malaysiana. 43: 37-44.
- [3] Talpur, A. D., Memon, A. J., Khan, M. I., Ikhwanuddin, M., Danish Daniel, M. M., and Abol-Munafi, A. B. 2011. A Novel of Gut Pathogenic Bacteria of Blue Swimming Crab Portunus pelagicus (Linneaus, 1758) and Pathogenicity of Vibrio harveyi a Transmission Agent In Larval Culture Under Hatchery Conditions. Res J Appl Sci. 6: 116-127.
- [4] Xu, Y., Gallert, C., and Winter, J. 2008. Chitin Purification from Shrimp Wastes by Microbial Deproteination and Decalcification. Appl Microbiol Biotechnol. 79: 687-697.
- [5] Habraken, W., Habibovic, P., Epple, M., and Bohner, M. 2016. Calcium Phosphates in Biomedical Applications: Materials for the Future? Mater Today. 19: 69-87.
- [6] Boutinguiza, M., Pou, J., Comesaña, R., Lusquiños, F., de Carlos A., and León, B. 2012. Biological Hydroxyapatite Obtained from Fish Bones. Mater Sci Eng C. 32: 478-486.
- [7] Ge, H., Zhao, B., Lai, Y., Hu, X., Zhang, D., and Hu, K. 2010. From Crabshell to Chitosan-Hydroxyapatite Composite Material via a Biomorphic Mineralization Synthesis Method. J Mater Sci Mater Med. 21: 1781-1787.
- [8] Metz, J. R., de Vrieze, E., Lock E-J, Schulten, I. E., and Flik G. 2012. Elasmoid Scales of Fishes as Model in Biomedical Bone Research. J Appl Ichthyol. 28: 382-387.
- [9] Amer Nordin, J., Prajitno, D. H., Saidin, S., Nur, H., and Hermawan, H. 2015. Structure-property Relationships of Iron–Hydroxyapatite Ceramic Matrix Nanocomposite Fabricated Using Mechanosynthesis Method. Mater Sci Eng C. 51: 294-299.
- [10] LeGeros, R. Z. 2002. Properties of Osteoconductive Biomaterials: Calcium Phosphates. Clin Orthop Relat Res. 395: 81-98.
- [11] Surmenev, R. A., Surmeneva, M. A., and Ivanova, A. A. 2014. Significance of Calcium Phosphate Coatings for the Enhancement of New Bone Osteogenesis - A Review. Acta Biomater. 10: 557-579.

- [12] Akram, M., Ahmed, R., Shakir, I., Wan Ibrahim, W. I., and Hussain, R. 2014. Extracting Hydroxyapatite and Its Precursors from Natural Resources. J Mater Sci. 49: 1461-1475.
- [13] Tadic, D., and Epple, M. 2004. A Thorough Physicochemical Characterisation of 14 Calcium Phosphate-based Bone Substitution Materials In Comparison to Natural Bone. Biomater. 25: 987-994.
- [14] Andrianjatovo, H., and Lemaitre, J. M. 1995. Effects of Polysaccharides on the Cement Properties in the Monocalcium Phosphate/B-Tricalcium Phosphate System. Innovation et technologie en biologie et médecine. 16: 140-147
- [15] Apelt, D., Theiss, F., El-Warrak, A. O., Zlinszky, K., Bettschart-Wolfisberger, R., Bohner, M., Matter, S., Auer, J. A., and Von Rechenberg, B. 2004. In Vivo Behavior of Three Different Injectable Hydraulic Calcium Phosphate Cements. Biomater. 25: 1439-1451.
- [16] Bohner, M. 2010. Design of Ceramic-based Cements and Putties for Bone Graft Substitution. Eur Cell Mater. 20: 1-12.
- [17] Takagi, S., Chow, L. C., Hirayama, S., and Sugawara, A. 2003. Premixed Calcium-phosphate Cement Pastes. J Biomed Mater Res B Appl Biomater. 67B: 689-696.
- [18] Aberg, J., Brisby, H., Henriksson, H. B., Lindahl, A., Thomsen, P., and Engqvist, H. 2010. Premixed acidic calcium phosphate cement: Characterization of strength and microstructure. J Biomed Mater Res B Appl Biomater. 938: 436-441.
- [19] Heinemann, S., Rössler, S., Lemm, M., Ruhnow, M., and Nies, B. 2013. Properties of Injectable Ready-to-Use Calcium Phosphate Cement Based on Water-Immiscible liquid. Acta Biomater. 9: 6199-6207.
- [20] Wagoner Johnson, A. J., and Herschler, B. A. 2011. A Review of the Mechanical Behavior of CaP and CaP/polymer Composites for Applications in Bone Replacement and Repair. Acta Biomater. 7:16-30.
- [21] Unabia, R., Piagola, J. C., Guerrero, J. R., Vequizo, R., Gambe, J., Odarve, M. K., and Sambo, B. R. 2015. Synthesis and Characterization of Nanocrystalline Hydroxyapatite and Biphasic Calcum Phosphate using Ca(OH)₂ and (NH₄)H₂PO4. Phys Status Solidi C. 12: 572-575.
- [22] Kokubo, T., and Takadama, H. 2006. How Useful is SBF in Predicting in Vivo Bone Bioactivity? Biomater. 27: 2907-2915.
- [23] Havelin, L. I., Engesaeter, L. B., Espehaug, B., Furnes, O., Lie, S. A., and Vollset, S. E. 2000. The Norwegian Arthroplasty Register: 11 years and 73,000 Arthroplasties. Acta Orthop Scand. 71: 337-353.
- [24] Moroni, A., Faldini, C., Rocca, M., Stea, S., and Giannini, S. 2002. Improvement of the Bone-screw Interface Strength with Hydroxyapatite-Coated and Titanium-Coated AO/ASIF Cortical Screws. J Orthop Trauma. 16: 257–263.
- [25] Russell, T. A., and Leighton, R. K. 2008. Alpha-BSM Tibial Plateau Fracture Study Group, Comparison of Autogenous Bone Graft and Endothermic Calcium Phosphate Cement for Defect Augmentation in Tibial Plateau Fractures, A Multicenter, Prospective, Randomized Study. J Bone Joint Surg. 90: 2057-2061.
- [26] Reig, F. B., Gimeno Adelantado, J. V., and Moya Moreno, M. C. M. 2002. FTIR Quantitative Analysis of Calcium Carbonate (Calcite) and Silica (Quartz) Mixtures Using the Constant Ratio Method. Application to Geological Samples. *Talanta*. 58: 811-821.
- [27] Ducheyne, P., Healy, K., Hutmacher, D. E., Grainger, D. W., and James Kirkpatrick, C. 2011. Comprehensive Biomaterials. 1st ed. Amsterdam: Elsevier. 1: 1-705.
- [28] Marques, D. R., dos Santos, L. A., Schopf, L. F., and de Fraga, J. C. S. 2013. Analysis of Poly (lactic-co-glycolic acid)/poly (isoprene) Polymeric Blend for Application as Biomaterial. Polímeros. 23: 579-584.
- [29] Nath, S. D., Son, S., Sadiasa, A., Min, Y. K., and Lee, B. T. 2013. Preparation and Characterization of PLGA Microspheres by the Electrospraying Method for

- Delivering Simvastatin for Bone Regeneration. *Int J Pharm.* 443: 87-94.
- [30] Félix Lanao Rosa P., Kemal S., Huanan W., Wolke Joop G. C., Jansen John A., and Leeuwenburgh Sander C. G. 2013. Accelerated Calcium Phosphate Cement Degradation Due to Incorporation of Glucono-Delta-Lactone Microparticles. Tissue Eng A. 20: 378–388.
- [31] Chang, H. I., and Wang, Y. 2011. Cell Responses to Surface and Architecture of Tissue Engineering Scaffolds. In: Eberli D, Editor. Regenerative Medicine and Tissue Engineering-Cells and Biomaterials, Croatia: InTech. 27: 569-588
- [32] D'Souza, S., Dorati, R., and DeLuca, P. P. 2014. Effect of Hydration on Physicochemical Properties of End-capped PLGA. Adv Biomater. 1-9.

FABRICATION OF POLY(LACTIC-CO-GLYCOLIC ACID)_CALCIUM PHOSPHATE

ORIGINALITY REPORT

9%

0%

9%

0%

SIMILARITY INDEX

INTERNET SOURCES

PUBLICATIONS

STUDENT PAPERS

MATCH ALL SOURCES (ONLY SELECTED SOURCE PRINTED)

10%

- ★ N M Daud, Z Tapsir, A Idris, A K Nasution, S Saidin.
- "Synthesization and characterization of chlorhexidine/calcium phosphate/poly(lactic--glycolic acid) bone cement ", IOP Conference Series: Materials Science and Engineering, 2019

Publication

Exclude quotes

On

Exclude matches

< 3%

Exclude bibliography

On