Injectable Bone Substitute Paste Based on Hydroxyapatite, Gelatin and Streptomycin for Spinal Tuberculosis

Hendita Nur Maulida MD, Dyah Hikmawati and Aniek Setiya Budiani MD

1 Biomedical Engineering Study Program, Faculty of Science and Technology, The University of Airlangga, Surabaya, Indonesia
2 Department of Physics, Faculty of Science and Technology, The University of Airlangga, Surabaya, Indonesia
3 Department of Clinical Pharmacy, Faculty of Pharmacy, The University of Airlangga, Surabaya, Indonesia

Abstract

World Health Organization (WHO) in 2005 reported that cases of tuberculosis (TB) in the world occur more than 8 million annually and 5-10% was attacked in spine. The most effective treatment of spinal TB is evacuation of infected bone segments and fills with bone graft. It has been synthesized and characterized of Injectable Bone Substitute (IBS) paste based on hydroxyapatite, gelatin and streptomycin. IBS paste synthesized by mixing hydroxyapatite and gelatin 20% w/v with 75:25, 70:30, 65:35 and 60:40 ratio and streptomycin 10 w%. The mixture was then added with hydroxypropyl methylcellulose (HPMC) 4% w/v as suspending agent. In vitro characterization performed includes acidity (pH), injectability test, setting time, cytotoxicity (MTT assay) and microbacterium test. Acidity test results indicate a fourth variation of the samples had pH values approaching normal body pH (7.3 to 7.6) and is able to maintain stability when measured in 7 days. Injectability test results indicate IBS paste is injectable with the highest percentage of the injectability value at 97.74% ± 0.19%. IBS paste has been setting within 30 minutes to 1 hour when injected on hydroxyapatite scaffold that resembles the bone cavity and is able to cover the pore scaffold seen from the Scanning Electron Microscope (SEM). Scaffold pore size is smaller from range of 780.8 to 835.4 μm into 225.2 μm. MTT assay results showed that IBS paste is not toxic and experiencing proliferation (viability >100%) that are expected to trigger osteoblast cell growth when applied. Microbacterium test results showed that IBS paste is an antibacterial seen from inhibition zone diameter of Staphylococcus aureus and has a high strength-sensitive antibacterial. Thus, hydroxyapatite, gelatin and streptomycin composites had qualified as injectable bone substitute which applied in cases of spinal tuberculosis.

Keywords: Spinal tuberculosis; Injectable bone substitute; Hydroxyapatite; Gelatin; Streptomycin; Hydroxypropyl methylcellulose

Introduction

Tuberculosis (TB) is an infectious disease caused by Mycobacterium tuberculosis which are systemic and can manifest in almost all organs of the body with the lungs as the primary infection site [1]. World Health Organization (WHO) in 2005 reported that TB cases in the world occur more than 8 million per year and 3% of them occur in Southeast Asia, including Indonesia [2,3]. Indonesia is the 3rd largest contributor to cases of TB after India and China with every 4 minutes there was one patient who died and transmission occurs every 2 seconds [1]. The incidence of extra-pulmonary TB occurs in 25-30% and 5-10% of which occur in the bones and joints and the most attacking in the spine. Bone TB cases is diagnosed when the symptoms have started after one year [4].

Spinal infection by TB could potentially cause serious morbidity including neurological deficits and permanent spinal deformity. Handling of spinal TB in general is giving antituberculous drugs, immobilization using gaps also orthopedic and neurological surgical intervention [5]. Surgical intervention will be taken when after 3-4 weeks of antituberculosis drug administration and conservative therapy does not give a good response. In this case, the most effective spinal lesions treated with immediate surgery to evacuate TB bacteria, take the infected bone and fill the spinal segment involved with bone graft to promote healing and achieve spinal stability [6].

Commericially, the material that can be used as bone graft is Hydroxyapatite (HA). Hydroxyapatite is a bioceramic materials that is biocompatible and bioactive because of its mineral content, both physically and chemically very similar to bone [7]. Hydroxyapatite can be obtained from the bones of mammals, fish bones, shells and other materials based on calcium phosphate. However, hydroxyapatite is brittle so that hard formed according to the required implant materials [8]. Some researchers then developed a composite material of hydroxyapatite with polymers to improve the mechanical properties, one of which gelatin. Gelatin as a polymer derived from natural materials is biodegradable, biocompatible and non-toxic. Gelatin is widely used as space filler because it is easy to set up [9].

Some use in orthopedics requires injectable biomaterial graft which serves as a filler. Injectable system is ready to use, can follow the shape of bone cavity to be filled and polymerized in-situ after being injected. To produce composite injectable, it is required a gel as matrix material maker. Polymers are widely used came from the class of cellulose such as Hydroxypropyl Methylcellulose (HPMC) [8]. HPMC is a water-soluble polymer and is widely used in the food industry as a thickening maker. Polymers are widely used came from the class of cellulose such as Hydroxypropyl Methylcellulose (HPMC) [8]. HPMC is a water-soluble polymer and is widely used in the food industry as a thickening due to high viscosity [10]. Manufacture of injectable bone substitute (IBS) has been done by several researchers including Weiss et al. [11] with IBS based on calcium phosphate and Shen et al. [12] with IBS based on alendronate and calcium phosphate.

In this study, we have been synthesized and characterized of IBS paste based on nano-hydroxyapatite, gelatin and streptomycin for spinal TB cases. The use of nano-sized hydroxyapatite is expected to easily fit into the pores of the bones thoroughly. While the streptomycin is antibiotic, the primary antituberculosis drugs produced by the soil fungus Streptomyces griseus. This drug is readily soluble in water [13].

*Corresponding author: Hendita Nur Maulida, Biomedical Engineering Study Program, Faculty of Science and Technology, The University of Airlangga, Surabaya, Indonesia, Tel: +62 856-5524-2442; E-mail: henditalamaulida@gmail.com

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The main target of streptomycin is working mechanism at the level of ribosome [14].

Characterization in this study include acidity test to determine degree of acidity of IBS to reach the body’s normal pH and its stability as paste, injectability test to measure the ability of IBS can come out of injections, setting time to determine the time of IBS experience setting when injected, MTT Assay test to determine the citotoxicity of IBS when tested in fibroblast BHK-21 cells and microbacterium test to determine the IBS resistance against bacterial pathogens using *S. aureus*. These characterizations are to evaluate the ability of hydroxyapatite, gelatin, and streptomycin as injectable bone substitute for spinal TB cases.

Materials and Methods

Materials

Tools used include digital scales Mettler Toledo, magnetic stirrer Yellow MAG HS 7, deep freezer, freeze dryer, thermometer, Terumo Syringe 12 ml and 5 ml, glassware (vial bottles, beaker glass and petri dish), micrometers couplers, stopwatch, viscoseter VT-04F RION, pH meter Benchtop OAKTON, Scanning Electron Microscope (SEM) FEI Inspect S50 Japan, a set of MTT Assay and microbacterium test.

Materials used include nano-hydroxyapatite of Barramundi fish from the National Nuclear Energy Agency (BATAN) Jakarta, cow skin gelatin from 150 bloom Rousselot (Guangdong, China), streptomycin sulfate from PT. Meiji Indonesian Pharmaceutical Industries, Hydroxypropyl Methylcellulose (HPMC) from Sigma Aldrich, hydroxyapatite scaffold from Tissue Bank of Dr. Soetomo hospital Surabaya, *Staphylococcus aureus*, Trypticase Soy Broth (TSB) and microbiology nutrient agar from Merck.

Synthesis of injectable bone substitute paste

IBS paste made in 4 variations in the composition of hydroxyapatite-gelatin that is 75:25, 70:30, 65:35 and 60:40 (w/w). Gelatin 20% w/v is dissolved in distilled water at 40°C. Hydroxyapatite with variations in composition put into the gelatin solution then added streptomycin 10 wt% as a local dose [15]. Meanwhile, HPMC 4% w/v is dissolved in distilled water at 90°C. Furthermore HPMC solution is added to a solution of gelatin, hydroxyapatite and streptomycin at 40°C and the mixture was stirred for six hours to produce a white IBS paste.

Acidity (pH) test

Acidity test is used to determine the degree of acidity and the stability of IBS as paste when measured in 7 days. Measurement performed with placing electrodes into the sample and pH will automatically appear on the screen of pH meter and make sure that the samples are measured at room temperature.

Injectability test

Injectability conducted to evaluate the ability of IBS paste can come out of the injection within a certain time. IBS first measured the viscosity to ensure the applicable value as injectable system. Measurement results showing viscosity IBS in units dPa.s. Furthermore, injectability test performed using the reference method reported by Shen et al. [12]. In this test, we used the syringe 12 ml with inner diameter of 1.5 cm and needle with inner diameter of 12 mm. Mass of IBS paste before and after injection within 2 minutes measured and its injectability calculated using the following equation.

$$\text{Injectability} \% = \frac{\text{mass expelled from the syringe}}{\text{total mass of the paste before injecting}} \times 100 \% \quad (1)$$

Each test was repeated five times and the average value was calculated as a result of percentage of injectability.

Setting time

Setting time testing was performed using hydroxyapatite scaffold that has been freeze-dried as a substrate [16]. Hydroxyapatite scaffold mass were measured to observe the changes that occur after setting. Testing was performed by injecting IBS samples into the scaffold vertically. Time counting begins when IBS penetrated into the scaffold pores and stopped when the surface of the scaffold was completely dry and covered by the IBS. The test results also were observed using SEM to determine the surface morphology of microscopic scaffold in sizes below 200 nm.

Cytotoxicity test with MTT assay

Cytotoxicity assay was performed using reagents MTT [3-(4,5-dimethyl-2-thiazoli) -2.5-diphenil-2H-tetrazolium bromide] which comprises the step of culturing fibroblast Baby Hamster Kidney (BHK-21) cells, placing samples and reading the results. Optical cell density can be determined using Elisa Reader and use equation (2) to calculate the percentage of living cells. Material is not-toxic if the percentage of living cells is more than 50% [17].

$$\% \text{ Living Cells} = \frac{OD \text{ Treatment} - OD \text{ Media Control}}{OD \text{ Cells Control} + OD \text{ Media Control}} \times 100 \% \quad (2)$$

where : OD Treatment = optical density value of the sample after treatment

OD Cells Control = optical density value of the control cells

OD Media Control = optical density value of media control

% Living Cells = the percentage of the number of cells after treatment

Microbacterium test

Microbacterium test is used to determine the IBS resistance against bacterial pathogens by looking at the profile of bacterial inhibition zone diameter for several days. Culture of one ose *Staphylococcus aureus* (SA) suspended in 9 ml Trypticase Soy Broth (TSB) then cultured in incubator 60°C for 24 hours. Absorbance values of SA bacteria are calculated using UV-Visible Spectrometer. Nutrient agar was prepared as a medium for bacteria. IBS paste put in wells agar medium with four repetitions and observed inhibition zone diameter after incubation for 24 hours. IBS strength levels against bacterial resistance were evaluated by comparing clear zone diameter according to Table 1 [18].

<table>
<thead>
<tr>
<th>Clear Zone Diameter</th>
<th>Microorganism Resistance</th>
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<tbody>
<tr>
<td>&gt;20 mm</td>
<td>Very Sensitive</td>
</tr>
<tr>
<td>10-20 mm</td>
<td>Sensitive</td>
</tr>
<tr>
<td>5-10 mm</td>
<td>Less Sensitive</td>
</tr>
<tr>
<td>&lt;5 mm</td>
<td>Resistant</td>
</tr>
</tbody>
</table>

Table 1: Strength rate of bacteria.

Results and Discussion

Acidity (pH) test

Degree of acidity (pH) becomes one of the important factors to evaluate the performance of IBS material when applied. From the measurement results, we obtained an average pH of each sample with variation composition of HA-gelatin 75:25, 70:30, 65:35 and 60:40 (w/w) respectively is 7.62; 7.54; 7.41 and 7.35. Graph stability of IBS pH variation composition of HA-gelatin 75:25, 70:30, 65:35 and 60:40 (w/w) respectively is 7.62; 7.54; 7.41 and 7.35. Graph stability of IBS pH value can be seen in Figure 1.
IBS material requires pH more than 6 to be setting in bone. pH that is not too far from normal pH of around 7.8 (slightly alkaline conditions) can still be tolerated by the body [19]. pH IBS samples were approaching the body’s normal pH (about 6.8 to 7.4) are expected to have no pain effect in the bones when it is applied.

**Injectability test**

IBS paste viscosity values were measured using visoctester show number of 120 dPa.s and applicable in terms of injectability test results as shown in Figure 2. From the test results, it can be concluded that the four samples of IBS has a very good ability in terms of the percentage results approaching 100%. Best percentage of injectability owned by the sample C with HA-gelatin ratio at 65:35 (w/w) is equal to 97.74% ± 0.19%.

**Setting time**

Setting time performed using HA scaffold freeze-dried as a simulation of human bone parts. Media or substrates that have the same as the main constituent component of the sample will be able to trigger these components for neat and improve the crystallinity. From the test results, the final setting time raised about 30 minutes to 1 hour as shown in Figure 3.

Measurement results of mass scaffold before and after setting also showed an increase. This change is due to the scaffold HA synthesized by the method of freeze-dried is produce a pore that allows the IBS to infiltrate so that the crystallinity increases and pore seemed closed neatly. Changes in mass of the scaffold before and after setting surface microscopically analyzed using SEM as shown in Figure 4.

From the scanning results, it appears that HA scaffold surface is covered evenly by IBS paste and the pores is smaller. The pore size before tested shows the distribution of values in the range of 780.8-835.4 μm and after scaffold injected by IBS in the range of 225.2 μm. It might be concluded that the IBS paste is able to evenly into the pores of the scaffold and bind hydroxyapatite in the vicinity so that crystallinity increases. Thus, IBS paste is able as bone substitution to fill the infected bone segments and vulnerable to further trigger the growth of new bone cells.

**Cytotoxicity test with MTT assay**

Each various samples of IBS tested in four repetitions and cell viability was calculated according to the equation (2) and then averaged as shown in Figure 5. From the test results, all sample variation of IBS is not-toxic seen from the percentage of cell viability which exceeds 50% [17]. There are several samples that have a percentage of cell viability of more than 100%. This shows that fibroblast cells capable of undergoing proliferation in the sample so that it becomes greater than the control cells. Thus, it can be concluded that the IBS paste can become new cell growth media are also expected to be a medium for the growth of osteoblast cells in bone when applied.

**Microbacterium test**

Based on the test results, it is known that the IBS paste is antibacterial seen from the area of inhibition zone around IBS discs with a diameter.
of 28-33 mm (Figure 6). This inhibition zone diameter increased when incubated up to 3 days as shown in Figure 7.

The level of material strength against bacterial resistance can be evaluated by comparing the clear zone diameter by calculating the difference in bacterial inhibition zone diameter with a sample wells diameter. Obtained clear zone diameter test results in Table 2. In addition to a bone filling material, IBS paste also expected to act as an antibacterial drug delivery systems to evacuate the TB bacterium that encourages faster healing and achieve early spinal stability.

Conclusions

1. In the physical characteristics, IBS paste has a pH value close to normal body pH (7.3 to 7.6) and is able to maintain stability when measured in 7 days. IBS paste is injectable with the highest percentage of injectability value at 97.74% ± 0.19% and setting within 30 minutes to 1 hour when injected at HA scaffold that resembles the bone cavity and is able to cover the pore scaffold seen from the SEM results.

2. In the biological characteristics, IBS paste is not-toxic and it is antibacterial seen from inhibition zone diameter of *S. aureus* bacteria.

References