Synthesis of bone cement from a natural mineral for biomedical industry


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Abstract- Study carried to find out chemical and structural suitability of newly synthesized Eppawala Hydroxyapatite composite as bone cement, by comparing and contrasting it with human bone as well as commercially available bone cement, which is currently used in orthopedic surgeries. Therefore, a mixture of commercially available bone cement and its liquid monomer, commercially available Methyl Methacrylate (MMA) and a mixture of Solid State synthesized Eppawala Hydroxyapatite powder with commercially available MMA were prepared as the direct substitution for bone cement. Then physical and chemical properties including composition, crystallinity, presence of functional groups, thermal stability, surface morphology, and microstructural features were examined compared to human bone. Results show there is a close similarity between synthesized product and human bone while credenting high thermal stability, good crystalline, and porous properties than the commercial product. Finally, study concluded newly synthesized composite can be applied directly as a substitution for commercial bone cement.

Index Terms- Bone cement, Human bone, Hydroxyapatite, Methyl Methacrylate, Orthopedics,

I. INTRODUCTION

Hydroxyapatite is widely used as a bioceramic due to its close chemical and structural similarity with human hard tissues. It performs several outstanding properties biocompatibility, non-inflammatory in nature, osteoconductivity, non-toxicity, bioactivity etc.[1-6] As a result it has a range of biomedical applications mainly in the fields of orthopedics and dentistry.[1-20]

Here in this study we have synthesized Hydroxyapatite by converting Chloroapatite using a Solid State Sintering method considering its ability to replace chlorine with other groups at high temperature due to the increase of reactivity as its chlorine positions are under strain in the structural framework.[21,22] Chloroapatite were collected from Sri Lankan Eppawala apatite deposit ,which usually contains 34-40% total phosphorus expressed as percentage of Phosphorus pentoxide (P2O5).[21-25] Further synthesized Hydroxyapatite is reinforced with a reactive resin, Methyl Methacrylate. It is a methyl ester of methacrylic acid. Polymerized forms of synthetic methacrylate resins used as cements in Orthopedic and dentistry applications. Also it is able fix prosthetic devices to bones and to cement bone to bone in difficult fractures as adhesives. [26]

Selected commercial product containing two main parts as bone cement majorly consists of Zirconium Dioxide and a liquid monomer Methyl Methacrylate (MMA), is currently used as fast curing bone cement in Sri Lankan government hospitals which indicates for stable attachment of total or partial joint endoprotheses in bone, filling and stabilization bone defects within the scope of internal fixation treatment or for endoprosthesis revision surgery and primary and secondary coverage of skull bone defects. It is prepared directly before use by mixing its powder component with liquid monomer component clinically. As a result ductile dough forms which cure within a few minutes. [27]

As this study designed only to find out the possibility of substituting newly synthesized ceramic composite into human bone, we are only focusing to find out structural suitability of ceramic composite as bone cement.

II. EXPERIMENTAL PROCEDURE

A. Sample preparation

Natural raw apatite mineral were collected from the Eppawala Apatite site. Then they were sorted as High Grade Rock Phosphate by the visual appearance of less coated apatite. After removing mud, collection of Apatite rocks were dried under sunlight, crushed using a jaw Crusher (Serial no: 1720011, China) into small crystals /powder, grind further into micron/Nano level HERP powder using a planetary Ball Mill (XQM – 4.0A) and sieved using sieve set (A060_01AC/0219, Scotland). Less than 63 micron range particle size powder were collected and oven dried at a temperature less than 150 °C for 5 hrs to prepare Moisture Removed HERP powder (MHERP). MHERP was taken as the raw material for synthesizing Hydroxyapatite. Samples were prepared using Solid state sintering technique as mentioned bellow. MHERP powder was added with needed weight of Ca(OH)2 powder, after well mixing, sieving and high temperature heat treating Solid State Sintered Eppawala Hydroxyapatite powder (SSHAp) was synthesized according to equation (1) given below.[12,21]
Then the synthesized ceramic powder was mixed with commercial Methyl Methacrylate (MMA) liquid monomer, until the ductile dough forms. As the second step, commercial cement powder and liquid monomer were mixed together until the ductile dough forms.

### B. Sample characterization

Before mixing with the liquid monomer, Commercial bone cement, and raw Eppawala Hydroxyapatite was examined under X-ray fluorescence Spectroscopy (Rigaku XRF Spectrometer) to find out its elementary composition and presence of impurities. Liquid monomer was examined with Fourier Transform Infrared Spectroscopy (Bruker – Alpha FTIR Spectroscopy) ATR mode to confirm its composition. Then Sample mixture of newly prepared bone cement and the sample mixture of commercial bone cement were characterized using XRD, FTIR, TGA, and SEM with EDS techniques together with the human bone sample. The crystallographic phases of samples were determined by X-ray diffractometer (Rigaku – Ultima IV diffractometer) in reflection mode with Cu Kα1: 0.154 nm radiation. 1.50 min⁻¹ scanned speed was used to collect data within a 2θ range from 10° to 80°. The presence of functional groups was confirmed by scanne

### III. RESULTS AND DISCUSSION

Synthesized Solid State Sintered Eppawala Hydroxyapatite (SSHAp) powder contains Ca, P and O include in higher weight percentages and Fe, Al and Si as the impurities with hexagonal crystal structure showing a close similarity with mammalian bones and consists of many correlated, microcrystalline structures/particles/ spherulites with micro pores while credenting good thermal stability. The presence of functional groups was confirmed by

#### Table 1. XRF results for Commercial Bone cement

<table>
<thead>
<tr>
<th>Element</th>
<th>Spot 1</th>
<th>Spot 2</th>
<th>Spot 3</th>
<th>Spot 4</th>
<th>Spot 5</th>
<th>Spot 6</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mass %</td>
<td>Wt %</td>
<td>Wt %</td>
<td>Wt %</td>
<td>Wt %</td>
<td>Wt %</td>
<td>Wt %</td>
</tr>
<tr>
<td>Ca</td>
<td>20.3</td>
<td>19.8</td>
<td>28.4</td>
<td>0.2</td>
<td>0.0</td>
<td>0.1</td>
</tr>
<tr>
<td>O</td>
<td>56.8</td>
<td>56.8</td>
<td>49.8</td>
<td>72.5</td>
<td>72.2</td>
<td>71.4</td>
</tr>
<tr>
<td>P</td>
<td>8.1</td>
<td>8.1</td>
<td>8.7</td>
<td>0.1</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>Cl</td>
<td>0.8</td>
<td>0.8</td>
<td>1.2</td>
<td>0.0</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>Zr</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>0.7</td>
<td>0.7</td>
<td>1.9</td>
</tr>
<tr>
<td>Fe</td>
<td>0.1</td>
<td>0.1</td>
<td>0.1</td>
<td>-</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>S</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>0.2</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>Na</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>0.4</td>
<td>0.4</td>
</tr>
<tr>
<td>Mg</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>0.2</td>
<td>0.2</td>
</tr>
</tbody>
</table>

Table 2. SEM with EDS results for SSHAp with MMA mixture, Commercial bone cement with MMA mixture and Human bone

<table>
<thead>
<tr>
<th>Element</th>
<th>SSHAp with mixture</th>
<th>MMA with mixture</th>
<th>Commercial bone cement with MMA mixture</th>
<th>Human bone</th>
</tr>
</thead>
<tbody>
<tr>
<td>Spot 1</td>
<td>Spot 2</td>
<td>Spot 3</td>
<td>Spot 1</td>
<td>Spot 2</td>
</tr>
<tr>
<td>Wt %</td>
<td>Wt %</td>
<td>Wt %</td>
<td>Wt %</td>
<td>Wt %</td>
</tr>
<tr>
<td>O</td>
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<tr>
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<td>0.2</td>
</tr>
<tr>
<td>P</td>
<td>8.1</td>
<td>8.1</td>
<td>8.7</td>
<td>0.1</td>
</tr>
<tr>
<td>Mg</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>0.2</td>
</tr>
</tbody>
</table>

Results OF SEM with EDS Analysis for SSHAP with MMA mixture, Commercial bone cement with MMA mixture and Human bone are indicated in the Table 2. According to those results; SSHAp with MMA mixture sample contains O, Ca in higher amounts and then P, C, Cl in order. Fe also found in very

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less amount as an impurity. In the Commercial product mixture O, and C carried in higher amounts and then Ca, Zr, S, and P in order. When consider human bone O presence as the highest amount and then C, Ca, P and Na presence in descending order. Na, Mg, Al, Si, S, Fe presence in very fewer amounts. It performs composition similarity with the mixture of SSHAp and MMA.

![SEM images for SSHAp ceramic+ MMA mixture, 10.0 kv, a) 10K b) 5KX c) 2KX d) 500X](image1)

![SEM images for Commercial bone cement + MMA mixture, 10.0 kv, e) 10K f) 5KX g) 2.2KX h) 500X](image2)

![SEM images for Human bone, 10.0 kv, i) 30KX j) 5KX k) 2KX l) 500X](image3)

Considering Figure 1-3; SEM images of all mixtures and human bone show that there are good correlations of particles. SSHAp with MMA mixture and human bone only carried out micropores as shown in the Figure 1. Porosity would be helpful for bone ingrowth as well as for good blood circulation. Presence of some different particles shaped in ball with some rough surface, was found in Commercial bone cement mixture which may lead to have higher surface area as mentioned in Figure 2. According to Figure 3, some crystalline property can be found in both human bone and SSHAp with MMA mixture.

![FTIR graph for liquid monomer](image4)

![FTIR comparison for Commercial Bone cement with MMA mixture, SSHAp with MMA mixture and Human bone](image5)

Figure 4 shows the resulted graph for liquid monomer it has coincided with the FTIR characteristic graph for Methyl Methacrylate monomer. It interprets several peaks related to stretching vibrations including a sharp intense peak at 1731 cm\(^{-1}\) related to the presence of ester carbonyl group, broad peak nearly 1150 cm\(^{-1}\) due to the C-O (ester bond) and a peak nearly 800 cm\(^{-1}\) is due to the bending of C-H. Also literature shows the broad peak ranging from 3100-2900 cm\(^{-1}\) is owing to the presence of stretching vibration. As shown in the Figure 5, all peaks for phosphate groups in the 560 cm\(^{-1}\) - 640 cm\(^{-1}\), 963 cm\(^{-1}\), 1028 cm\(^{-1}\) and 1110 cm\(^{-1}\) wave no range and Characteristic peak for OH/ Hydroxyapatite nearly 3572 cm\(^{-1}\) wave no appeared in the Human bone as well as the SSHAp with MMA mixture. It confirms that even after the mixing with a monomer, the presence of Hydroxyapatite in the SSHAp product. When considering Commercial product mixture, it shows peak nearly 3572 cm\(^{-1}\) wave no range, which may due to the presence of OH\(^{-}\) group, but that couldn’t be identified as Hydroxyapatite characteristic peak, as no peaks found related to phosphate groups. Apart from that, some peaks can be found commonly in both commercial bone cement+ liquid monomer mixture and SSHAp with liquid monomer mixture except in human bone nearly (750 cm\(^{-1}\)- 2000 cm\(^{-1}\)) wave no range and 3000 cm\(^{-1}\) wave no which is also appeared in Figure 4, they are the related peaks for Methyl Methacrylate monomer. As a result it can be
concluded that both commercial bone cement and SSHAp mixed well with the liquid monomer.

Figure 6. XRD pattern for (a) SSHAp with MMA mixture (b) Commercial bone cement with MMA mixture

Figure 6(a), explains even after mixing MMA monomer, XRD results of SSHAp mixture all characteristic peaks related to the crystallographic phases 002, 210, 211, 112, 300, 202, 310, 222, 213 and 004 of hexagonal Hydroxyapatite, which shows similarity to Human bone. Also, it interprets 96% crystallinity. Figure 6(b), carries the XRD results for Commercial bone cement and it has interpreted all the peaks related to 110, 210, 111, 002, 200, T02, 211, 022, 122, 300, 013, 302, T13 and 222 crystallographic phases of monoclinic Zirconium Dioxide crystal structure with 84.1% crystallinity. Comparing those results with the literature for human it can be confirmed that both human bone and synthesized composite has structural similarity via consisting Hexagonal Hydroxyapatite except for commercial product.

12.9850 mg SSHAp with MMA sample mixture was subjected to TGA as shown in the Figure 7 First significant weight loss which occurs in between 100 °C- 400 °C (0.2449 mg) representing 1.886 %, may associate with the dehydration of the sample. Following that interval the sample reduced its weight nearly 0.1511 mg at 772.07 °C; it may occur due to the gas elimination. Then again from 772.07 °C to 1432.97 °C there is a weight loss indicating 0.45% (0.0584 mg) which has occurred due to the incipient transformation of produced HAp in β – TCP. Therefore, it indicates the formation of Hydroxyapatite in products. At 1432.97 °C 96.51 % of the original weight has remained.

As figured in Figure 8, 14.0710 mg of Commercial product mixture was subjected to TGA. Nearly 200 °C to 400 °C considerable amount of weight loss was occupied representing 85.14% from initial weight (11.96 mg). Basically up to 400 °C weight losses may occur due to the dehydration of sample/moisture removal. Therefore, it can be predicted first weight loss may occur due to moisture removal of the product, but there may be some other reasons also such as structural degradation or deformation. After that, weight loss rate has become slower and then a constant value. At the end 1435.98 °C, 15.23% of initial weight was remained.

According to the Figure 9, 10.115 mg human bone sample was subjected to TGA. As mentioned in the literature; first significant weight loss which occurs nearly at 200 °C (0.9112 mg) representing 9.008 %, may associate with the dehydration of the sample. Following that interval the sample reduced its weight nearly 5.6358 mg at 650 °C; it has occurred due to the bone structure collagen elimination. This reaction continues up to 936.88 °C, with a lowered rate. Above that temperature, a fine TGA curve descending slope is observed up to maximum
analyzed temperature of 1432.97 °C with the total weight loss of 54.79%, this being associated with the collagen remains removal & the incipient transformation of Hydroxyapatite in β – TCP. [12, 21, 29, 31]

When comparing Human bone with Commercial bone cement with MMA mixture and SSHAp with MMA mixture, according to Figure 10, Human bone and the SSHAp mixture have shown the same pattern of weight loss, which was slightly different from Commercial product mixture. That may happen due to the composition similarity of Human bone and SSHAp with MMA monomer mixture, as they were containing Hydroxyapatite. Also due to the least amount of weight loss in synthesized SSHAp mixture sample than bone and commercial product mixture, it can be concluded that the synthesized SSHAP with MMA monomer mixture perform high thermal stability and good material stability in nature and application.

IV. CONCLUSION

The study concludes that SSHAp with MMA composite has a chemical and structural similarity with human bone and performs high thermal stability and good material stability in nature. Therefore, it can be used as a direct substitution for bone cement.

REFERENCES


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