SYNTHESIS OF HA AND BETA-TCP USING SOL-GEL PROCESS AND ANALYSIS WITH FTIR

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Abstract. Nanotechnology has been attracting researchers’ attention due to its innovative concepts and associated technology development. Nowadays, one approach to control the surface’s chemistry and in vivo solubility of calcium phosphate compounds are associated to the employment of nanoparticles. The sol-gel process is used for synthesis of many materials in nanoscale, because your process starts on a colloidal suspension. The hydroxyapatite \( \text{Ca}_{10}(\text{PO}_4)_{6}(\text{OH})_2 \) is the mineral phase of bone tissue. The hydroxyapatite bioceramic is a bioactive material and because of this reason its application with bone substitute and also as a coating or film in materials with a biological response less then hydroxyapatite. \( \beta \)-TCP, \( [\beta\text{-Ca}_3(\text{PO}_4)_2] \), is widely used on the development of bone defects fulfilling since its chemical composition leads to implants that exhibit excellent biocompatibility, bioactivity and resorbable. This material could be employed as bone cement, as porous or dense blocks, as granulated material and in all cases the particle size and morphology are key factors to achieve appropriate chemical, biological and mechanical properties. In this context, the goal of this work is to synthesize nanostructured HA and \( \beta \)-TCP using sol-gel method which has proved to be very efficient in obtaining nanomaterials with high homogeneity. The DSC showed the optimal temperature range of gel work and the best temperature range from 50 to 90ºC. For characterization was used scanning electron microscopy (MEV), X-ray fluorescence (XFR), and Scherrer equation associated with X-ray diffraction (XRD). Surfaces good bioactivity have investigated using bands derived from \( \text{PO}_3^-\text{4} \) by Fourier transform infrared spectroscopy (FTIR).

Keywords: Hydroxyapatite, beta-tricalcium phosphate, Fourier transform infrared spectroscopy, sol-gel process with sucrose, nanotechnology.

1. INTRODUCTION

The calcium phosphates are the most important inorganic components of bone tissue “Abdel Fattah et al. (2008)”. The common calcium phosphate laboratory syntheses are: hydroxyapatite, beta-tricalcium phosphate and alpha-tricalcium phosphate. The hydroxyapatite (HA) is a bioactive calcium phosphate bioceramic, that in biological environment is not stoichiometric as the synthetic material (Ca/P: 1.67). The mineral phase of synthetic material is similar to bone tissue and it is used for bone replacement. The bioactivity of HA induces bone growth and facilitates the regeneration process “Guha et al. (2009)”.

Beta-tricalcium phosphate \( \beta \)-TCP is a bioceramic with Ca/P of 1.5. This material is used as bone cement for filling small facial defects, and also be possible used for the fabrication of dense or porous ceramic composites. Beta-TCP is a biodegradable material that can be replaced by bone tissue “Ohura et al. (1996)”.

Calcium phosphates are being applied on scaffolds construction for tissue engineering “Kawachi et al. (2000)”.

The nanotechnology is an innovative concept with numerous technological advances, therefore became the object of study by many researchers worldwide “Souza et al. (2007)”. The nanotechnology in biomaterials enable functional regulation of cells, which plays a central role in nanomedicine targeting at regenerative medicine and intractable diseases. It is possible design the nanostructure of materials that induces cell differentiation at specific positions of the living body. The nanostructures are based on design bioinspired performing necessary function at the right time and the right place by manipulating assembly of molecules and atoms forming the natural tissue, for example bone tissue.

During the development of new biomaterials, nanoparticles play an important role because their greater superficial are often leads to distinct values of reactivity, mechanical, chemical and biological properties in vitro and in vivo when compared to micro and macroscopic particles “Quina et al. (2004)”.

The sol-gel product is characterized by primary nanoparticles, and this is a crucial parameter to improve the surface reaction and the stability at implant/natural bone interface. Another approach to the sol-gel process is the sucrose addition, which allows the synthesis of particles smaller than the conventional process “Rodrigues et al. (2009)”.

In this work, two types of nanostructured calcium phosphate were synthetized by sol-gel process with sucrose, were: hydroxyapatite and beta-tricalcium phosphate.
For characterization of this bioceramics were used: X-ray diffraction (XRD), X-ray fluorescence (XRF), scanning electron microscope (SEM) and Fourier transform infrared spectroscopy (FTIR).

In XRD were analyzed for crystalline materials. The standards used for the comparison and confirmation of results it was obtained in the JCPDS library.

The XRF is used to characterize the materials by semi-quantitative analysis of each chemical element present in the sample.

The SEM is used to visualize the nanoparticles size and identify the morphology of material.

The FTIR analysis indicated characteristic spectra bands for all the studied materials and identifies a composition of sample “Laranjeira et al. (2010)”.

2. MATERIALS AND METHODS

2.1. Materials

The reagents used were: phosphoric acid - LAFAN 85% (H₃PO₄), calcium nitrate tetrahydrate – Synth 99% [Ca(NO₃)₂ . 4 H₂O] and Sucrose – Synth (C₁₂H₂₂O₁₁).

The hydroxyapatite Ca/P: 1.67 and β-tricalcium phosphate Ca/P: 1.5.

2.2. Methods

Hydroxyapatite was prepared from two aqueous solutions of sucrose, one with phosphoric acid and, the other with calcium nitrate tetrahydrate. The Ca/P ratio was maintained equal to 1.67. Sucrose was used as a gel forming and dispersant agent. Then, the gel was fired at 700ºC.

Beta-tricalcium phosphate was prepared of same way that hydroxyapatite, but maintaining the Ca/P ratio in 1.5. The gel was fired at 1050ºC.

![Fluxogram of sol-gel process with sucrose. Both bioceramics were calcined, but hydroxyapatite at 700°C and β-TCP at 1050°C.](image)

2.3. Characterizations

The morphology and microstructure were analyzed by scanning electron microscopy (SEM). The equipment used was a JOEL model JXA-840A of DEMA-FEM-UNICAMP, SEM-FEG JEOL JSM 6330F belonging to LNLS-LME.

The X-ray diffraction was performed to verify the crystallinity and phases present in the material. The equipment belong to the DEMA-FEM-UNICAMP, model DMAX 2200-Rigaku, Cu-Kα radiation with Ni filter. Analyses were performed at 40kV and 30mA in a 2θ range from 20º to 50º, with step size of 0.02º and 2 second of integration time.

The crystallite size were calculated using the Scherrer Equation “Nag et al. (2007)” (Eq. 1) after fitting a gaussian curve on the principal peak to determine their width at the half height intensity “Briks et al. (1946)”.

\[
t = \frac{0.9\lambda}{\beta \cos \theta}
\]

Where, “t” is the crystallite mean size, “λ” is the X-ray wavelength, “β” given by Eq. (2) is the peak width at half height intensity, and “θ” is the Bragg angle.

\[
\beta = \beta_{exp} - \beta_{inst}
\]

Where, “β_{exp}” is the peak width at half height intensity of the sample and “β_{inst}” is the instrumental correction obtained by the width at half height intensity of the monocrystal pattern.
The X-ray fluorescence spectroscopy was used to determine the impurities quantitative and the equipment used was Rigaku RIX 3100 spectrometer. The sample were conformed into 25mm disks and analyzed by a semi-quantitative routine.

Infrared spectra analysis was performed with a Fourier transform infrared spectroscopy (FTIR) and the equipment used was Thermo Scientific Nicolet IR100 FT-IR with spectral resolution of 128 to 4 cm\(^{-1}\). The spectral used was 4000 to 400 cm\(^{-1}\) and the samples were ground with dried potassium bromide (KBr) powder, and tablet pressed into a disc for this analyses.

3. RESULTS AND DISCUSSION

3.1 X-ray diffraction

The standard graphics of XRD were obtained in database Joint Committee on Powder Diffraction Standards (JCPDS) and the HA has the number 09-0169 and β-TCP has the number 09-0432.

![Figure 2. X-ray diffraction of sample. (a) Graphic of HA with a peak contamination of B-TCP. (b) Graphic of β-TCP.](image)

After heat treatment the HA shown the peaks in Fig. 2 (a) indicating a contamination with beta-TCP. The sample B-TCP shown in Fig. 2 (b) the peaks of beta-TCP and Hydroxyapatite, this problem occurred because the sol-gel process with sucrose to synthetize beta-tcp is a new application. This process for synthesis of calcium phosphates as hydroxyapatite and beta-tcp is new and it was developed based on the article “Souza et al. (2007)”. Sucrose is introduced into the process to decrease the size of the particles.

<table>
<thead>
<tr>
<th>Sample</th>
<th>Crystallite (nm)</th>
</tr>
</thead>
<tbody>
<tr>
<td>HA</td>
<td>~58</td>
</tr>
<tr>
<td>B-TCP</td>
<td>~95</td>
</tr>
</tbody>
</table>

The results of X-ray diffraction were used to perform analysis of crystallite size by Scherrer equation. The results were consistent due to temperature difference of sample calcination. The Hydroxyapatite crystallite had an average size of about 58nm and beta-TCP calcined at 1050°C had an average size of 95nm.

<table>
<thead>
<tr>
<th>Materials</th>
<th>Sr</th>
<th>Fe</th>
<th>Na</th>
<th>S</th>
<th>Cl</th>
<th>Mg</th>
<th>Si</th>
<th>Al</th>
<th>Ni</th>
<th>K</th>
<th>Mn</th>
<th>Zr</th>
</tr>
</thead>
<tbody>
<tr>
<td>HA (wt.%)</td>
<td>0.028</td>
<td>0.004</td>
<td>0.018</td>
<td>0.007</td>
<td>0.028</td>
<td>b.l.d.</td>
<td>b.l.d.</td>
<td>b.l.d.</td>
<td>b.l.d.</td>
<td>b.l.d.</td>
<td>b.l.d.</td>
<td>b.l.d.</td>
</tr>
<tr>
<td>B-TCP (wt.%)</td>
<td>0.018</td>
<td>b.l.d.</td>
<td>0.011</td>
<td>0.006</td>
<td>b.l.d.</td>
<td>b.l.d.</td>
<td>b.l.d.</td>
<td>0.005</td>
<td>0.003</td>
<td>b.l.d.</td>
<td>b.l.d.</td>
<td>0.006</td>
</tr>
<tr>
<td>HA-VETEC (wt.%)</td>
<td>0.012</td>
<td>0.015</td>
<td>0.074</td>
<td>0.021</td>
<td>0.032</td>
<td>0.325</td>
<td>0.020</td>
<td>0.009</td>
<td>0.002</td>
<td>0.012</td>
<td>0.009</td>
<td>b.l.d.</td>
</tr>
</tbody>
</table>

b.l.d.: below the limit of detection
The X-ray fluorescence showed a very important result for the use of these materials in the biomedical area because it has not significant amounts of contaminants and quantities of each element is small compared to commercial material as HA-VETEC shown in the Tab. 2.

![Image 1](image1.png)

**Figure 3.** Image of SEM. (a) Shown the HA nanoparticle aggregates. (b) Shown the β-TCP nanoparticle aggregates.

The scanning electron microscopy proved that the samples have the approximated results obtained with Scherrer equation. The Fig. 3 (a) shown the HA nanoparticle clusters with nanometer size. Fig. 3 (b) shown Beta-TCP with larger particle size when compared with HA, but the results are similar the results of Scherrer equation when compared the particle size with scale.

![Image 2](image2.png)

**Figure 4.** FTIR. First line: Hydroxyapatite. Second line: Beta-tricalcium phosphate and hydroxyapatite.

The FTIR spectrums exhibited the characteristics absorption peaks of $\text{PO}_4^{3-}$. The band OH (600 cm$^{-1}$) reduced in spectrum of the mix beta-tricalcium phosphate and the hydroxyapatite, the vibration bonds decrease in spectra because was forming hydrogen phosphate. In addition, enlarge the bond attributed to the stretching vibration of $\text{HPO}_4^{2-}$ at 1221 cm$^{-1}$. The P-O stretching and bending is not clear at range of 863-732 cm$^{-1}$ and 504–475 cm$^{-1}$ in the HA and Beta-TCP.
4. CONCLUSION

In the results of X-ray diffraction, the hydroxyapatite only have a Beta-TCP peak contamination, this is due to Ca/P has not been perfect during the synthesis. The Fig. 2 (b) shown peaks of beta-TCP and HA, this problem occurred due to sol-gel process with sucrose to synthesis of beta-TCP it is a new method and it needs modifications in your process parameters.

The results of Scherrer equation shown the crystallite size below 100nm, therefore these are nanomaterials. The HA with ~58nm and beta-TCP with ~95nm.

The X-ray fluorescence to semi-quantitative analysis showed a low amount chemical elements presents in the samples and it showed also low wt.% of each element when to compare HA and B-TCP with commercial HA.

The scanning electron microscopy showed the morphology and an idea about the particle size. The hydroxyapatite showed the agglomerate structure with nanoparticles bellow 100nm. The beta-TCP with HA contamination showed nanoparticles and large dense blocks due to calcination temperature.

By FTIR, the evidence the key factors influence at morphologies nonconventional in the case were non-stoichiometric phosphocalcic in HA and Beta-TCP.

3. ACKNOWLEDGEMENTS

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4. REFERENCES


5. RESPONSIBILITY NOTICE

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