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Effect of CO_2 Ion Doping on Characteristics of Nanostructure Hydroxyapatite

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Abstract

In this research work, carbonated hydroxyapatite samples with different carbonate contents were prepared using ion implantation method. Pure nano-structure hydroxyapatite samples were implanted by 80 keV CO_2^+ ions during different times of 4, 8, 16, and 24 min using an ion implantation devise. The resulting materials were tested by X-ray diffraction, Fourier transform Infrared spectroscopy, and transmission electron microscopy, electron spin resonance systems before and after the implantation. The results confirmed the doping of carbonate group into the hydroxyapatite structure, and show that the presence of carbonate in the structure of hydroxyapatite decreases the degree of crystallinity. Moreover, it was observed that the increasing of ion implantation time caused to decrease the average grain size of the hydroxyapatite sample.

Keywords: Hydroxyapatite; Ion implantation; Carbonate; FTIR; XRD; TEM; ESR

Introduction

Hydroxyapatite (HAP) is a biocompatible ceramic having the physical and chemical properties similar to the bones and teeth [1,2]. This bio-ceramic have numerous applications including ceramic coating, manufacturing the implants substituted for the bones and teeth, cosmetic foundation [3], radiation dosimetry [4,5] etc. The calcium hydroxyapatite produced by living tissue is called biological HAP. In fact the synthetic type of HAP with the formula of $Ca_{10}(PO_4)_6$ (OH), differs slightly from the natural one. Researches show that the biological HAP contains the carbonate groups [6-8] and a small amount of other elements. During the formation of HAP as the hard part of the bones and teeth, some organic groups such as carbonate (CO_3^{2}) have been replaced over the time in the position of hydroxyl (OH^{-}) or phosphate (PO_4^{3-}) groups in the HAP crystals. Therefore, they have differences in terms of physical, chemical, mechanical properties as well as their crystallinity, and solubility in comparison to the artificial HAP.

Ion implantation is one of the ways to change the physical structure of thin films. Replacement of ion in the network structure will change the chemical composition of the substrate material, thus other properties of material could be changed subsequently. In this method, high-energy ions produced by an ion implantation device with perfect vacuum penetrate into the substrate material. Planted ions may lead to three processes within the material that are: scission of primary bond and replacement of network atoms, entering the channel (space between the structure), and locating the position of network defects [9,10].

In the present work the CO_2^+ ions entered into the HAP crystal structure in different implantation times and the changes in structure were studied and compared.

Materials and Methods

Sample preparation

Polyvinyl alcohol (PVA) powder (2 w%) was dissolved in water at room temperature, to prepare an adhesive solution. The resulted glue was mixed with HAP powder prepared from Merck Company. The produced paste was kept for two days in air evacuated nylon to agglomerate and become suitable to use. The obtained substance was milled for half an hour to obtain a smooth and homogeneous powder. Then the pill shape samples with 13 mm in diameter were prepared using a hydraulic press, punch and matrix made of non-corrosive alloy under 700 MPa.

Ion implantation procedure

The prepared pills without any hand contact and using the cleaned tools were transferred to the special sample place in the ion implantation system. Before each implanting operation, the chamber was scoured out with alcohol and acetone. When the vacuum pressure reached to 2×10^{-5} Torr, the valve connected to the CO₂ gas capsule was opened till the vacuum reached the desired values (Table 1). Then the appropriate voltage and current were applied to produce an 80 keV CO₂⁺ ion beam. The samples were exposed to the ion beam in different time periods of 4, 8, 16 and 24 min. All the system parameters during the ion implantation are depicted in Table 1. An ion implantation system model MBM100, made in China was used in this research work.

Characterization

Fourier transmission infrared spectroscopy (FTIR) was carried out in the wave number range of 400-4000 cm⁻¹ using a Perkin Elmer, series 100 spectrometer.

System parameters	Sample 1	Sample 2	Sample 3	Sample 4
implantation time (s)	240	480	960	1440
Ion beam current (mA)	16	18	18	20.5
Beam voltage (kV)	80	80	78	83
Vacuum Pressure (2 × 10 ⁻⁵ Pa)	3.7	3.2	3.2	3.7
Beam current on sample (µA)	0.5	0.5	0.5	0.5

Table 1: Ion implantation system parameters.

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X-ray diffraction (XRD) analysis was performed by a Philips Analytical X-Ray B.V., using the Cu-K_a radiation (1.5456 Å wavelength), in 2 θ range of 20°-60°. The grain size of the prepared products was calculated using the Scherer's equation indicated as below:

$$t(nm) = (0.89 \times \lambda(nm)) / (B(rad) \times cos\theta)$$

where t is sample grain size, λ is the X-ray wavelength, B is full width at half maximum of a peak in the X-ray pattern, and θ is the Bragg's angle.

Transmission electron microscopy (TEM) studies were also carried out using a Zeiss EM10C microscope at 80 kV.

ESR measurement

Electron spin resonance (ESR) spectroscopy is an extremely sensitive method for studying materials containing unpaired electrons. The samples were put into the quartz thin-wall ESR tubes (4 mm diameter) and measured with a Bruker EMS-104 spectrometer operating in x-band. The ESR signal intensities were measured as peak to peak height for the most intense ESR lines (first derivative of the absorption spectra) per sample mass. The used ESR spectrometer parameters for this study were 0.285 mT modulation amplitude, 100 kHz modulation frequency, 3.0 mT scan width, 1024 point field resolution, 164 msec time constant, 21 sec sweep time, and 50 dB receiver gains.

Result and Discussions

Figure 1 shows the XRD patterns of the samples were implanted at different times in comparison to the non-implanted one. Evaluation and comparison of the diffraction patterns before and after the implanting indicate the existence of the peaks in all samples which are fully in accordance with the standard cards of hydroxyapatite (card no. 00-0350-0180). Among these peaks, the Miller indices of the most intense 9 peaks are indicated in the figure. The XRD patterns show that the peak positions before and after implantation have not changed and no new peak appeared afterward. Considering the intensity of the peak in position of 25.9° with Miller indices of (002), it is obvious that an increase in implantation time (carbon increase), decreases the peak





height and increases the peak width as shown in Figure 2. This result is in agreement with those reported in some references [11-15].

Therefore, it can be concluded that shortening the peak height and increasing the peak due to intensification of the implantation time, will be followed by crystallinity reduction. This variation was calculated and given in Table 2 and shown in Figure 3.

Figure 4 shows the FTIR analysis results for the samples before and after the implantation. All of the observed bands are related to the carbon apatite which can be arranged in three types of phosphate, carbonate and hydroxyl groups. A broad band in position of 1043 cm⁻¹ and two narrow bands in positions of 567 cm⁻¹ and 603 cm⁻¹, and another one in position 962 cm⁻¹ are related to the phosphate compounds. Two bands in positions 603 cm⁻¹ and 1051 cm⁻¹ are also reported for phosphate compounds. The bands were observed in positions of 883 cm⁻¹ and 1417 cm⁻¹ till 1576 cm⁻¹ were associated with carbon compounds that





Implantaion time	Peak position	Peak width	Peak height
0 min	25.981°	0.19 6°	149
4 min	26.022°	0.198°	94
8 min	25.985°	0.275°	76
16 min	25.947°	0.305°	63
24 min	25.952°	0.332°	47

Table 2: Data related to (002) peak.





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became more intense by increasing the implantation time. This means that the carbon entered in to the structure of hydroxyapatite samples via this method. Another broad band around the position of 3436 cm⁻¹ was related to the hydroxyl ions (water) which means that the samples containing little water were probably related to the production stage of the pills [15].

Chemical structure of the hydroxyapatite and CO_2^+ ion is shown in Figure 5 [13]. According to this figure, the CO_2^+ ions can enter into the channels or bonds with the oxygen in one of the two indicated positions, and eventually form carbon apatite.

Therefore, it can be concluded that the ion implantation technique is a physical method to insert the carbonate group into the structure of hydroxyapatite, which it leads to the production of carbon apatite. Also, in this method the implantation time, sample moisture content, and the ion energy can be considered as the process variables to improve the result.

The grain sizes of the HAP samples were calculated via equation 1 using the data extracted from the XRD patterns for the main XRD peaks. The averages over the same implantation time are reported in Table 3.

These calculations were done via fitting the Gaussian function to each single peak and calculating their overlap, with respect to the





Implantation time	Average particle size		
0 min	22.24 ± 3.17 nm		
4 min	21.19 ± 3.51 nm		
8 min	19.22 ± 2.61 nm		
16 min	18.66 ± 2.51 nm		
24 min	17.98 ± 2.82 nm		

Table 3: Average grain sizes of the HAP samples implanted in different times.

position and peak widths. The indicated standard deviation means that there is a distribution over the particle sizes in each set of the HAP

Page 3 of 4

samples. These results demonstrate that the average grain size decreases while the ion implantation time increases. In the author's view the decreasing of sample grain sizes is associated with the sputtering of grains due to the hitting of accelerated carbon ions. In fact, an increase in implantation time causes more spattering and a smaller particle size. TEM micrographs of the HAP samples are demonstrated in Figure 6. The particle sizes in the samples are obvious in TEM-associated image which confirms the obtained results indicated in Table 3.

Figure 7 shows the ESR responses of the samples after different ion implantation times. According to this figure, the ESR signal increases strongly after 4 min ion implantation as a result of carbon ion incorporation into HAP crystal, and remained approximately constant at the higher implantation time.

In fact the carbonated impurities are incorporated into, or attached to the surface of HAP crystal during formation, and are converted to radicals through absorption of ionizing radiation. Thus, the nonimplanted sample, do not show an intensive ESR response where this response value could be due to the native or mechanical induced signals. At the higher implantation time the amount of induced radicals remains constant due to the fact that the carbonate ions mostly contribute in sputtering of the particles and loss their energy afterwards. It means that the saturation occurs after a few minutes of implantation.

Conclusions

The FTIR analysis indicates that as carbon entered into the



Figure 6: Magazine logo TEM provided microstructure of HAP samples a) before implantation and b) implanted with CO_2^+ ions in 24 min.



structure of HAP the carbon apatite was formed. Increasing the implantation time decreases the (002) peak height and increases the peak width in the XRD patterns of the HAP samples which means crystallinity reduction.

The obtained results from the XRD and TEM analyses demonstrate that the average grain size decreases while the ion implantation time increases. Therefore, the ion implantation technique is a physical method to insert the carbonate group into the structure of HAP and it leads to the production of carbon apatite.

The EPR response of the HAP samples implanted by carbonate ions show the higher intensities in comparison to non-implanted sample. But the saturation occurs after a few minutes of implantation.

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