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In Vitro Study of Iron Doped Hydroxyapatite

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Abstract. The effect of iron substitution on the bioactivity of hydroxyapatite (HAp) under the physiological conditions was investigated. Five samples of iron doped hydroxyapatite (FeHAp) with different iron concentrations (0, 0.05, 0.1, 0.2, and 0.3 mol %) were synthesized by wet chemical method. The formation of bone like apatite layer on the surface of the samples was detected using X-ray diffraction (XRD) and Fourier transforms infrared (FTIR). The changes of the pH of SBF (*simulated body fluid medium*) were measured at pre-determined time intervals using a pH meter. The dissolution of calcium, phosphorus and iron ions in SBF medium was determined by single beam scanning spectrophotometer. XRD and FTIR results exhibit the formation of carbonate apatite layer on the surface of the immersed samples, which increased with the increase of iron content. The solubility and dissolution rat than pure sample, which indicate that iron increase the bioactivity of HAp in vitro.

Keywords: Hydroxyapatite, iron doped hydroxyapatite, bioactivity, bone like apatite, solubility.

1. Introduction

Hydroxyapatite (HAp) which has the chemical formula $[Ca_{10} (PO_4)_6(OH) _2]$ is well known as the mineral component of bones and teeth. There is considerable interest in this mineral in the dental and medical community due to its good osteoconductivity and osteoinductivity [1]. Bioactivity of HAp is its ability to bond directly to bone and forming a new bone after implanted in the living body. The mechanism of bone like apatite layer formation on the surface of the sample contains two main processes:

- Dissolution process
- Precipitation process

According to these processes the soaked sample will dissolve and with the increase in the immersion time the dissolution rate decrease and the precipitation rate increase until equilibrium reach then the Ca, P ions concentrations saturate. The needed time for formation of apatite layer on the surface of the sample can be varied from material to another. The faster of the formation of apatite layer, the greater the bioactivity of the material [2].

Many substitutions can introduce to HAp structure to improve its biocompatibility and bioactivity. For example the presence of strontium (Sr) enhances osteoblast activity and differentiation, while it inhibits osteoclast production and proliferation [3]. Also Silicon (Si) enhances the formation of a poorly crystalline surface apatite layer of HAp after immersing in simulated body fluids (SBF) [4].

The substitution of iron into the structure of HAp is known to decrease the formation of HAp [5-6], however little known about the behaviour of iron doped HAp under the physiological conditions. This work aims to study the effect of iron on the bioactivity and solubility of HAp under the physiological conditions.

2. Experimental Procedure

2.1 Synthesis of HAp and FeHAp

Five samples of hydroxyapatite were synthesized by wet chemical methods. The chemical reagent and starting materials concentrations for the precipitation are listed in Table (1).

Sample	Code	Ca(No ₃) ₂ .4	4H ₂ O	(NH ₄) ₂ HPC	D ₄	Fe(No ₃) ₃	.9H ₂ O
		gm	Mol	gm	mol	gm	mol
1	HAp	36.8	0.16	12.66	0.096		
2	FeHAp ₁	36.8	0.16	12.66	0.096	2.02	0.005
3	FeHAp ₂	36.8	0.16	12.66	0.096	4.04	0.01
4	FeHAp ₃	36.8	0.16	12.66	0.096	8.08	0.02
5	FeHAp ₄	36.8	0.16	12.66	0.096	12.12	0.03

Table (1) The chemical reagent of the prepared samples.

2.2 Preparation of simulated body fluid (SBF)

The preparation of SBF was adopted according to the protocol developed by Kokubo et al [7]. In order to prepare 1000 ml of SBF, first of all 700 ml of ion-exchanged and distilled water with a stirring bar into 1000 ml glass beaker is set on a hot plate covering with a watch glass maintaining the temperature at $36.5\pm2^{\circ}$ C under stirring. The first 8 order reagents given in Table (2) are dissolved in water one by one. The 9th and 10th order reagents are dissolved to adjust pH to 7.4. After the pH adjustment, the solution is cooled down to 20°C and enough ion-exchange and distilled water is added to complete the solution to 1000ml. The test specimens were immersed individually in test tubes in 50 mL of (SBF), in a thermodynamic incubator (shaking-water bath) at 37 oC. A 3-mL sample of SBF was taken from each test tube after 6, 12, 18, 24 and 28 days from the beginning of the immersion. The test tubes were discarded after the samples had been taken. The 9-mL sample was stored frozen until analyzed for Ca, P and Fe ions concentrations.

Table	(2).	Order,	amounts,	weighing	containers,	and formu	ila weights o	of reagents	for prepari	ing of	SBI	ď,
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Order	Reagent	Amount (g/L)	Container	Formula weight
1	NaCl	8.035	Weighing paper	58.4430
2	NaHCO ₃	0.355	Weighing paper	84.0068
3	KCl	0.225	Weighing bottle	74.5515
4	K2HPO4.3H2O	0.231	Weighing bottle	228.222
5	MgCl ₂ .6H ₂ O	0.311	Weighing bottle	203.3034
6	1.0m - HCl	39 ml	Graduated cylinder	
7	CaCl ₂	0.292	Weighing bottle	110.9848
8	Na ₂ SO ₄	0.072	Weighing bottle	142.0428
9	Tris	6.118	Weighing paper	121.1356
10	1.0m - HCl	0-5 ml	Syringe	

To examine the formation of apatite layer in the surface of the prepared samples, the samples were taken out from the SBF solution and gently washed with pure water. The samples should be dried in a desiccator without heating. The dried discs were crashed by a mortar and tested by XRD and FTIR techniques.

2.3 X-ray diffraction (XRD)

X-ray diffraction (XRD) pattern data were obtained at room temperature using powder samples in an **X'Pert PRO-PAN** anlytical diffractometer (with Cu-K α radiation, λ = 1.54056Ű) at 40 kV and 30mÅ. Intensity data were collected by the step-counting method (step 0.020 and a time per step of 0.4 s) between 4° and 80° (20)

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2.4 Fourier transform infrared (FTIR)

Fourier transform infrared (FTIR) spectroscopy was employed to determine the chemical functionality of CHAp. The potassium bromide (KBr) disk technique was used for analysis using 2 mg of CHAp powder compacted with 200 mg of KBr under hydraulic pressure.All the spectra were measured by using (**Model 6100**, **Jasco-Japan**) infrared spectrometer with a resolution of 4.00 cm⁻¹ at room temperature, which covers the wave number range of 4000 – 400 cm⁻¹ to evaluate the function group of the samples.

2.5 pH measurement

The change of the pH of the prepared samples was examined in SBF under a physiological condition (pH 7.4 at 37 ± 1.5 °C) and the pH was measured with (**pH meter model 3510 Jenway - Japan**) at pre-determined time intervals.

2.6 Concentrations of released ions

The quantitative analysis of released calcium, phosphorus and iron ion concentration after immersion in SBF are determined with single beam scanning spectrophotometer (Jenway 4600 UV-Vis Spectrophotometer -England) over range of 320-1100 nm using biological kits. All ions are measured using (Spectrum-diagnostics - Egypt).

3. Results and Discussions

3.1 XRD of the immersed samples

XRD patterns of pure HAp and FeHAp samples after soaking in SBF for 24 days are shown in Figure (1). The pure HAp sample doesn't reflect any phases except HAp as confirmed by the ICSD card no. 76-0694, while all iron doped HAp samples showed a hematite as a secondary phase beside HAp as confirmed by the ICSD card no. 89-0597. The presence of hematite may be attributed to the dissolution process of samples after soaking in SBF. This process introduces free ions to the solution such as Fe, Ca, P and O, which support the formation of hematite. A similar behavior was observed with iron doped HAp after sintering in air [8].



Figure (1). XRD of samples after immersion in SBF for 24 days.

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Figure (2) shows the percentage amount of hematite and apatite phases as a function of iron content. It was observed that hematite phase decreased, while apatite phase increased with the increase of iron content. This behavior suggested that iron enhance the formation of HAp, while it reduced the formation of hematite. To interpret this behavior, the condition of HAp nucleation must be considered. Peixin Zhu et al [9] reported that the nucleation of HAp induced by the electrostatic attraction force between the sample surface and the cations dissolved in the solution. According to this Fe³⁺ ions which have three positive charges may be more effective in inducing an electric field than Ca²⁺ ions, which have two positive charges only. Thus after dissolution, most iron ions accumulated on the sample surface due to the electrostatic force between the negatively charged surface of samples and the positively charged iron ions and hence the more iron ions the more is the nucleation sites for HAp. These result was found in good agreement with the results that have been already published [10- 11]. This effect combined with the decrease in the probability of hematite formation.



Fig. (2). Phase percentage of hematite and HAp of the prepared samples after soaking in SBF as a function of iron content.

Figure (3) shows the calculated lattice parameter of the studied samples after soaking in SBF for 28 days. The values of both *a* axis and unit cell volume of pure HAp are larger than that of all iron doped samples, while the value of *c* axis of HAp is smaller than that of all iron doped samples. Also figure (3) shows that the values of both *a* axis and unit cell volume of iron doped samples tend to decrease with the gradual increase of iron content, while the value of *c* axis tend to increase with the increase of iron content. This behavior may be attributed to the formation of carbonated HAp layer on the surface of the soaked samples, as its believed that the substitution of carbonate in the phosphate group sites (β - type) can cause an expansion in the unit cell length along the *c* axis and a contraction along the *a* axis.

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Conversely, when carbonate substitutes for the hydroxyl group (A- type) there is an expansion along the a axis and a contraction along the c axis [12]. Thus as we mentioned before that the probability of apatite layer in the samples surface enhanced by increasing iron content in the prepared samples, which can contributed to the increase the observed increase and decrease in the values of lattice parameter of the soaked samples.



Fig. (3). Lattice parameter of the soaked samples in SBF for 24 days.

The crystal size of the grown apatite layer in the surface of the soaked samples was calculated using sherrer's equation and listed in Table (3). These values ranged from about 29.1 to 18.9 nm, where the largest value (29.1 nm) is due to pure HAp sample. As the iron content was gradually increased, the crystal size showed gradual decrease to reach its minimum value (18.89 nm) due to the sample FeHAp₂, then the crystal size starts to increase again to reach (26.62 nm) due to the sample FeHAP₄. These results suggested the formation of carbonate substituted HAp (CHAp) and the observed decrease in the values of the crystal size of the first two samples may be due to the substitution of carbonate in the phosphate sites (β - type) [9- 12]. The increase in the values of the last two samples can be attributed to the substitution mechanism as the increase of carbonate content can lead to the replacement of phosphorus ion by more than one carbonate ion and hence enlarge the crystal size.

Table (3) represents the crystallinity of the soaked samples as calculated from XRD data. There are two observations from these data; the first one is the value of the crystallinity of the pure HAp sample, which is smaller than that of iron doped

samples. This may be due to the presence of iron since it can enhance the crystallization and growth of HAp in the physiological conditions [10-11].

The second observation is the gradual decrease in the crystallinity of all iron doped samples with the increase of iron content, except in the case of FeHAp₃ sample which showed a remarkable decrease compared with the decrease of all iron doped samples. The observed gradual decrease may be due to the formation of carbonated substituted HAp which prevents the formation of more crystalline layers as well as to prevent the dissolution of ions from the soaked samples. This in turn tends to decrease the crystal growth also. These results were found to agree well with the published results for carbonate substituted HAp [8-12]. The smallest crystal size value due to the sample FeHAp₃ can be attributed to the highest carbonate content in this sample compared with all iron doped samples as confirmed by IR result (see next section).

Sample	Average crystal size (nm)	Xc (%)	Xc (%)	
НАр	29.10172	14.77		
FeHAp ₁	24.51612	34.86		
FeHAp ₂	18.89284	30.38		
FeHAp ₃	22.06824	22.4		
FeHAp ₄	26.6263	28.94		

Table (3) The calculated crystal size and crystallinity of soaked samples.

3.2 FTIR of the immersed samples

The IR spectra of the immersed samples for 24 days are shown in figure (4). The phosphate group bands appear in all samples at ~ 962, 471, 1035 and 603, 566 for v₁, v₂, v₃, v₄ respectively. The broad band observed at ~ 1635 and ~ 3430 in all spectra indicate absorbed H₂O in the lattice. Broad bands appear at wave numbers ~ 875, ~ 1423 and 1454 indicate that carbonate group partially replace the (PO₄)³⁻ group in HAp lattice, which was β -type carbonated HAp. The presence of hydrogen phosphate, such as HPO₄ and H₂PO₄ adsorption band were not visible in all spectra. These results confirm the XRD result.

The formation of apatite layer appears clearly in all samples as all possess the fundamental functional groups of HAp. We can observe that the intensities of all peaks of pure HAp sample are small than that of all iron doped samples, which may be attributed to the increase in the crystallinity of iron doped samples compared with that of pure HAp sample as confirmed by XRD results. Also the intensity of carbonate, phosphate and hydroxyl groups increased and the peaks is sharper as the iron content increased. These results may be due to the precipitation of CHAp layer in the surface of the soaked samples, which increased with the increase of iron content [10-11].

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3.3 Solubility measurement

Figure (5) shows the pH versus time curves for all samples including the control sample, which didn't have HAp. We can observe that the pH of samples decreased in the first 12 days, while it approximately constant during the remaining soaking period. That's may be attributed to the alkaline effect of dissolved HAp content [13]. At the end of the interaction the solution saturated and then pH of the solution will stabilized. These results are in a good agreement with the literatured results for HAP [13-14-15-16]. According to these results and with the aid of XRD result, it may be suggested that the solubility of the prepared samples increased with the increase in the iron content.



Fig. (4). IR spectra of the immersed samples.

3.4 Concentration of released ions

Figure (6) illustrates the concentration of calcium ions during the immersion time in SBF. The concentration of calcium ions in the original SBF is about 100 ppm [17]. The concentration of calcium ions in the prepared samples start with 103.27 ppm in the un-soaked SBF and increased with the increase of immersion time to reach 123.78, 121.86, 118.95, 115.03, 112.75 ppm for HAP, FeHAP₁, FeHAP₂, FeHAP₃, FeHAP₄ respectively. The increase in the Ca concentration may be attributed to the release of calcium from sample to the solution [17-18-19].

The concentration of calcium ions in FeHAp samples increase with the increase of immersion time, while it decreases with the increase of iron content. This may be to the increased CHAp in the surface of the samples as the iron content increased, which confirmed by XRD and IR results.

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The original SBF contain about 30 ppm of phosphorus ions [17]. The concentration of phosphorus in the prepared samples start with 37.04 ppm in the unsoaked SBF and decreased with the increase of immersion time to reach 33.86, 32.01, 30.42, 29.36, 28.66 for HAP, FeHAP₁, FeHAP₂, FeHAP₃, FeHAP₄ respectively see figure (7).

The decrease in the phosphorous concentration may be attributed to their uptake from the solution to the surface of the samples to form the apatite layer [17-18-19]. The decrease of both calcium and phosphorus ions in the solution of iron doped samples may be revels that the increase of apatite formation capability on the surface of iron doped samples. This may be due to the presence of iron in these samples as we mentioned before.

Iron is a component of FeHAp, but SBF does not have this ions. Therefore, the presence of iron ions in the reacted solution is due to the leaching from the sample. Figure (8) indicates the concentration of released iron ions during the immersion time for iron doped HAp. We can observe that the concentration of released iron decreased with the increase of immersion time. This may be attributed to the uses of iron ions as a nucleation sites for the formed apatite layer so the number of remnant ions in the solution decreased. Another interesting observation is the increase of the amount of released iron with the increase of iron content in samples. That decrease attributed to the formation of hematite, which decreased with the increase of iron contents observed in XRD results.



Fig. (5). Chang of pH versus time of the immersed samples.

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Fig. (6). Concentration of calcium ions in SBF after 24days of immersion.



Fig. (7). Concentration of phosphorus ions in SBF after 24 days of immersion.



Fig. (8) .Concentration of released iron ions In SBF after 24 days of immersion.

4. Conclusion

The effect of iron on the bioactivity and solubility of HAp under the physiological conditions was investigated. The overall results showed that iron induced an electric field, which can enhance the nucleation and crystallization of HAp. The formed apatite layer increased with the increase of iron content. The solubility and dissolution rate of iron doped samples increase with the increase of iron content, which indicate that iron improved the bioactivity and solubility of HAp.

5. References

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خصائص الهيدروكسي أباتيت المطعم بالحديد : دراسة معملية خارج جسم الانسان

) فدم للنشر بف 11/4/2113م ، وفبل للنشر بف 11/1/2113م (

ملخص المحث. المركب فيمد الدراسة هو من أكثر المركبات أهمىة بن جمال الطب والهلندسة الطبيمة ويمعرف باسم هيدروكسي أبانايت. وهذا المرركب بالرغم من اسنعماال له الطبيمة إال أنه مازال حيّاج التي الدراسات الماس نفيضة لنحس في خصائاصه احلاليمة ولى هل نطبيمة مع فنع جماالت أخري جديدة لالس نفادة منه بعد حنسن هذه اخلواص.

الدراسة احلالية استهدفت إحالل عنصر احلديد بن هذا الماركب لنحس في خصائص المر الذي استان دراسة مدي مالى عنه الماركب جيويًا باعد إضافة احلديد وجنت ظروف فسي ولوجية خسابفة ومن هنا فن د أفرنح أضافة نسب من احلديد))1، 1010، 101، 201، و 103 مول٪ (ومن نم فن د من حتمري مخسة عينات للدراسة بالنسب السابقة بطريق لكيما ناية معروفة. أمكن

منابعة الدغري بف خصائص المركب باس نخدام حيود أل شعة السينية واس نخدام األ شعة منت احلمراء)بنطبيقات فورير(. كذلك من دراسة مدي دأئر العينات اخلمس بالغمر أل زمنة خمنافة بف جهاز حماكي لسوائل جسم الل نسان. أظهرت النائاج عموما منس أما مل حوظاً بن المال نامة احليوية للمركبات بعد إضافة احل ديد هاا.