Pumice and perlite co-substituted hydroxyapatite: Fabrication and characterization

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ABSTRACT

The purpose of this study was to combine hydroxyapatite (HA) with pumice and volcanic silicate perlite. Doped hydroxyapatite is inexpensive and easy to produce. A precipitation procedure was applied for synthesizing pure and doped hydroxyapatite. Samples were sintered at 1100°C for 1 h. These doped structures are characterized by X-ray Diffraction (XRD), Scanning Electron Microscopy (SEM-EDS) and X-ray Fluorescence. Higher SiO$_2$ and Al$_2$O$_3$ content was shown by XRF analysis. Nanoscale HA is obtained with density of 3.056 g/cm$^3$. The XRD results revealed the existence of the HA, β-TCP and Ca$_5$(PO$_4$)$_2$SiO$_4$ phases. SEM images confirmed the sintering temperature and number of dopants had significant effect on grain sizes of the samples.

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1. Introduction

In recent decades, porous ceramics materials are widely used in biomedical applications. Orosity provides reduced weight and widens surface area which allows better mechanical bonding with the host tissues. Hydroxyapatite (HA), Ca$_{10}$(PO$_4$)$_6$(OH)$_2$, which is in the group of calcium phosphate ceramics, considered biocompatible materials due to their ability to form strong bonds with osteogenic and hard tissues. HA is also used as a coating element in orthopedic and dental implants because of the structural similarities of the HA to apatite minerals in bone and tooth enamel [1-3]. Owing to the low fracture toughness and mechanical properties, HA is frequently used as a filling material in not load-bearing areas. Compressive strength and microhardness values of HA varies between 12 MPa-64 MPa and 85 HV-170 HV, respectively [4]. Recently, studies were carried out to strengthen the low mechanical properties of porous HA to improve its biological properties. One of these studies is doping HA with different elements [5]. In order to improve internal structure, mechanical, antibacterial and biocompatibility properties HA doped with different cations such as Ti, Y, Ag, Mg$^{2+}$, Sr$^{2+}$ and Al$^{3+}$ instead of Ca ions in the microstructure [6,7]. Mostly, synthetic HA powders are produced by hydrothermal, precipitation, hydrolysis of calcium phosphates, or sol-gel methods. Researches revealed that the mechanical properties of HA are primarily depending on the sintering temperature and particle size. Due to the longer mixing time in the precipitation method compared to other methods, materials with smaller particle sizes ($\leq$ 50 nm) produced, and therefore secondary phases not formed. The presence of secondary phases significantly reduces the mechanical strength of the material [8]. One of the methods that can be used to improve the mechanical properties of HA is doping with glass-ceramics obtained from silica or phosphate-based bioactive glasses. Studies revealed that composite materials strength and biocompatibility properties increased [9]. When the titanium implant is coated with silicon dioxide doped HA, the corrosion, surface roughness, and osseointegration properties increase compared to the pure HA [10].

In this study, HA has been doped separately with natural pumice and perlite. Pumice is a volcanic rock type that is porous, spongy, light, and highly resistant to atmospheric conditions. As the gases in its body leave the structure as it cools very quickly during its formation, numerous pores formed. Because of these pores, its permeability is low [4]. Pumice, which is resistant to physical and chemical factors, has a porosity ranging from 45% to 70%, and its solubility in water is very low. The density of natural pumice varies between 1.2-1.4 gr/cm$^3$, and its compression strength varies between 1.72 ± 0.12 MPa. The pH value of pumice is around seven and does not show any toxic properties [11]. In vitro tests performed on MG63 cells have shown that pumice increases the proliferation and adhesion of the cells [12]. In another study, it has been demonstrated that the biocompatibility and mechanical properties of HA, which is
doped with 5% natural pumice by weight, compared to pure HA, have increased [13].

Perlite is a silica-based volcanic glass, and it is an amorphous, inert material with low bulk density. Perlite consists of 71-75% silicon dioxide, 12.5 to 18.0% alumina, 1 to 4% sodium and calcium oxide, and traces of metal oxide. The density of perlite, which does not show toxic properties, varies between 0.6-2.30 gm / l, and particle size around 0.2-4 mm [14]. Although, there are many studies focusing on different aspects of perlite, a few studies found on calcium phosphates. From a biomedical point of view, porous composite (ecopore) with perlite surface modified by etching, aminosilane, and combined with fibronectin showed non-toxic and promote human primary osteoblasts growth [15].

The purpose of this study is to synthesize hydroxyapatite (HA) doped with different contents of pumice and perlite to develop new nanocomposite bioceramics and to determine the optimum pumice and perlite contents which yield the best properties in terms of microstructures and mechanical properties. Pure and doped HA synthesized by a precipitation method. The synthesized materials were sintered at 1100°C for 1h. Presence of phases and bonds were characterized by x-ray diffraction (XRD). Grain sizes of the samples were obtained by scanning electron microscopy (SEM). Chemical components of the samples obtained by X-ray fluorescence (XRF).

2. Materials and Method

During the production of samples, HA was doped separately with pumice and perlite particles by using the precipitation method at 1%, 2.5%, and 5% by weight. Due to the longer mixing time in the precipitation method compared to other methods, materials with smaller particle sizes (≤ 50 nm) are produced and therefore secondary phases are not formed. The precursor used in experimental studies were supplied from Sigma-Aldrich. In order to synthesize pure HA, precursor chemicals Ca(NO$_3$)$_2$4H$_2$O and (NH$_4$)$_2$HPO$_4$ were separately dissolved in deionized water. NH$_4$OH was added dropwise to the solution containing (NH$_4$)$_2$HPO$_4$ in order to pH value 11. NH$_4$OH was added in (NH$_4$)$_2$HPO$_4$ and two solutions mixed together to form one solution. The solution, which was stirred for one day at room temperature (24-25°C), was left for one day for precipitation. The wet cake obtained as a result of filtering was dried at 200 °C for 12 hours and then sintered at 1100 °C for 1 hour. After drying and sintering process, pure and doped powders were ground in agate mortar. Pumice and perlite powders were added separately to the ammonium phosphate solution in the amounts indicated in Table 1. The steps used in the production of pure HA are repeated in the production of doped HA. The XRF result, main element oxide percentages (%) of the pumice and perlite powders used in this study are given in Table 2 in terms of weight.

| Sample   | Pumice%mol | Perlite%mol |
|----------|------------|-------------|
| HA1.0Pumice | 1.0        | -           |
| HA2.5Pumice | 2.5        | -           |
| HA5.0Pumice | 5.0        | -           |
| HA1.0Perlite| -          | 1.0         |
| HA2.5Perlite| -          | 2.5         |
| HA5.0Perlite| -          | 5.0         |

Table 1. The ratio of pumice and perlite added to the doped HA.

| Oxides (Percentage by weight %)  |
| SiO$_2$ | Al$_2$O$_3$ | Fe$_2$O$_3$ | CaO | CaCO$_3$ | K$_2$O | Na$_2$O |
|--------|-------------|-------------|-----|---------|-------|--------|
| Pumice | 74.12       | 13.12       | 1.66| 6.04    | 3.82  | 1.25   |
| Perlite| 76.42       | 13.94       | 0.90| 3.10    | -     | 4.71   |

Table 2. The main oxide percentages elements of pumice and perlite powders (XRF).

XRD analysis was carried out to determine the phases in the produced powders. For this purpose, phases were determined by using a scanning speed of 2 °/min at 10-80° 2θ angles in a Panalytical/Empyrean device. Zeiss brand Sigma 300 device was used in SEM analysis. The chemical components of the phases in the microstructure of the powders were determined by the SEM-EDS.

3. Result and Discussion

XRD patterns of the sintered samples (1100°C) were given in Figures 1-2. XRD peaks of the figure 1 consisting of the ICDD main standard peak group with the HA, Ca$_5$(PO$_4$)$_2$SiO$_4$ and β-TCP (ICDD:09-0432, 40-0393, 55-0898). After doping with pumice (1, 2.5 and 5 wt. % Pu-HA) composite composed of mainly HA, with β-TCP as a secondary phase. The diffraction lines of 1, 2.5 and 5 wt. %Pu-HA samples diminished and broadened in comparison with pure HA. That means, 1, 2.5 and 5 wt. %Pu-HA have smaller grain size than pure HA.
Figure 1. XRD pattern of pure-HA and pumice doped HA (1 wt. %, 2.5wt. % and 5wt. %).

Figure 2. XRD pattern of pure-HA and perlite doped HA (1 wt. %, 2.5wt. % and 5wt. %).

XRD patterns of the HA doped with perlite and sintered at 1100°C (figure 2) are attributed to the phases of β-TCP and/or HA because peaks of β-TCP and HA are overlapping each other. In the case of 2.5wt. %Pe-HA and 5wt. %Pe-HA an increase of the β-TCP observed. Perlite addition increased the crystallinity. Due to perlite doping into peak apatite lattice, peak shift to higher 2θ values with an increase in perlite (5wt. %Pe-HA) content. As seen from the XRD patterns peaks are attributed to β-TCP and/or HA and as a third phase Ca₅(PO₄)₂SiO₄ was also observed. In the sample doped with 2.5wt. %Pe-HA ions, most of the HA peaks transformed into β-TCP phase. Because HA, pumice and perlites different thermal expansion coefficients of the crystalline phases (α) may also cause high stresses at the grain boundaries and increase the thermal decomposition of the HA phases [16]. The XRD results reveals that the decomposition rate of HA depends on the mass fraction of pumice among the composites [17].

Figure 3 and figure 4 shows the surface morphology of the pure-HA, (1wt. %, 5wt. %) Pu-HA and (1wt. %, 5wt. %) Pe-HA composites, respectively. The 1wt. %Pu-HA, 2.5wt. %Pu-HA and 5wt. %Pu-HA composites have porous surface with sponge-like views (Fig. 3b, c and d). When the pumice ratio increased the structure looks differs from homogeneous surface to scarce needle-shaped crystals. The sizes of the pores seem to be increased with respect to the pure-HA.

Figure 3. Scanning electron micrograph of the pure-HA (a), 1wt. %Pu-HA (b), 2.5wt. %Pu-HA (c) and 5wt. %Pu-HA(d).
The changes that occurred in the grain size and morphology of HA due to the ion dopings were observed in SEM images and average grain size measurements. SEM images revealed that temperature and ion amounts of dopants had significant effect on grain sizes of the samples. There is a relationship between amount of dopant and sintering temperature with grain sizes of apatites. SEM results shows that perlite doped HA, especially, 1 wt. %Pe-HA and 2.5wt. %Pe-HA composites have porous surface with sponge-like views (Fig. 4b, c and d). In fig.4b it obvious that there are some cracks on the grains. The reason for cracks could be related to the transformation of HA into β-TCP [18].

The SEM-EDS spectra along with pure-HA, Pu-HA and Pe-HA results were given in fig. 5 and fig. 6. Figure 5'a (pure-HA) comprised of mainly Ca (43.08 at. %), O (37.75 at. %) and P (19.16 at. %). In fig.5(b,c,d) pumice doped HA composed of Ca (25.36-32.18 at.%), O (44.61-51.58 at.%), P
Table 3. Density distributions of pumice and perlite doped HA.

| Sample            | Density (g/cm³) |
|-------------------|-----------------|
| HA                | 3.056           |
| HA1.0Pumice       | 2.778           |
| HA2.5Pumice       | 2.727           |
| HA5.0Pumice       | 2.565           |
| HA1.0Perlite      | 2.578           |
| HA2.5Perlite      | 2.282           |
| HA5.0Perlite      | 2.031           |

Several studies have been conducted to determine the density of HA. According to literature, the density of HA varies between 3.05-3.1 g/cm³. In those studies, the density of nanoscale HA is about 3.056 g/cm³; this result is compatible with our results (3.056 g/cm³). The lowest density among the doped samples was HA5.0Pumice and HA5.0Perlite which in combination with second phases in SEM results [19].

4. Conclusion

In this study, HA doped with pumice and perlite were synthesized by a precipitation method. Doping amount of pumice and perlite were kept at 1.0–5.0 wt.%. All samples were sintered at 1100°C for 1h. Microstructural and mechanical properties of the sintered samples were investigated via XRD, SEM-EDS and XRF methods. XRD patterns peaks are attributed to β-TCP and/or HA and as a third phase Ca₅(PO₄)₂SiO₄ was observed. The diffraction lines of 1, 2.5 and 5 wt. %Pu-HA samples diminished and broadened in comparison with pure HA. That means, 1, 2.5 and 5 wt. %Pu-HA have smaller grain size than pure HA. In this study, Pu-HA and Pe-HA non-toxic composites revealed a remarkable potential for biomedical applications as coating element and bone graft.

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