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Studying the effect of hydroxyapatite particles in osteoconductivity of Ti-HA bioceramic

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Abstract

Ball milling method and powder metallurgy technology were employed to synthetize metal matrix composites (MMC) for bone grafts' applications. The raw powder of the MMC was prepared by mechanical alloying of pure titanium (Ti) powder with hydroxyapatite (HA) particles. The biocompatibilities of the sintered Ti-HA composites were examined after immersing the samples in simulated body solution (SBF) for different periods of time. SEM image and XRD results analysis were utilized to study the effect of HA on osteoconductivity of the Ti-HA composite. To this purpose, several composites were synthetized from different Ti-HA raw powder combination based on the HA particle size, milling time, and the mass fraction of HA content (% w/w) in the MMC. In vitro analysis of Ti-HA composite shows that composite with 30% w/w HA has higher bioactivity in comparison with composite containing pure Ti with 10% w/w HA.

Keywords Powder technology \cdot Synthetic bone grafts \cdot Metal matrix composites (MMC) \cdot Hydroxyapatite (HA) \cdot Simulated body solution \cdot Bioceramic

Introduction

Titanium and its alloys are the most common metal in medical devices and implants due to their impressive corrosion resistance,

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osteoconductivity, bio-adhesion (bone ingrowths), modulus of elasticity, etc. Nevertheless, pure titanium (Ti) shows lack of desirable bioactive properties, particularly, bone-growth. Recent researches showed that it is possible to improve titanium's orthopedic implant fixation and surface bioactivity by coating it with biomaterial ceramics, such as hydroxyapatite (HA) or other calcium phosphate (CP) forms [1, 2]. HA with Ca_{10} (PO₄)₆(OH)₂ chemical formula has been widely used as bioactive material for a bone substitute because of its chemical similarities with the mineral components of bones. However, the current coatings techniques are prone to wearing or delamination [3].

The metal matrix composites (MMCs) are comprising rigid ceramic reinforcements embedded in a ductile metal or alloy metal matrix [4, 5]. The properties of MMCs are widely recognized to be controlled by the size and volume fraction of the reinforcements as well as by the nature of the matrix/ reinforcement interfaces. During the last decades, considerable research efforts have been directed to deploy MMCs production methods to develop reinforced Ti matrix composite using HA particles to improve the bioactivity and mechanical properties of the composite for medical applications [6–8].

In the present study, several Ti-Ha composites were sintered by applying ball milling method and powder metallurgy technology to serve as synthetic bone grafts. A series of experiments were conducted to study the effect of HA particle characterizations, introduced by initial HA particle size, specified HA percentages in the raw powder mixture, and the mechanical alloying milling time on bioactivity of sintered Ti-Ha composite. The in vitro studies of Ti-HA composites were performed using simulated body fluid (SBF) solution and surface morphology. Phase constituent of the composites are determined through SEM and XRD results. SEM provides the phase constituent of the composites before immersion in SBF while XRD gives the phase constituent of the composites after immersion in SBF.

Sample preparation

HA powders with 150-nm particle size were prepared by rapid microwave synthesis according to Bovand et al.'s methodology [9]. The micron-size HA with 15- μ m particle size was synthesized by co-precipitation method described in Sousa et al.'s research work [10].

Figure 1 shows SEM morphologies of the HA powders for the nano-size (Fig. 1a) and micron-size (Fig. 1b) HA powders with spherical- and rod-like morphology. The Ti powder (< 45μ m) from Merck KGaA was mixed with 10 and 30% w/wHA powders with 150 nm as well as 15- μ m size to have four Ti-HA mixture types. Half of each Ti-HA mixture group was milled for 20 h, and the other half was milled for 50 h to have eight Ti-HA powder combinations, labeled as C1 to C8 in Table 1.

Ti-HA powder combinations with 15-µm HA particles are labeled as Ti-mHA, while Ti-HA powder combinations with 150-nm HA particles are labeled as Ti-nHA. The mechanical milling was occurred in a planetary ball mill equipped with hardened steel jar and stainless steel balls. The milling was performed under argon atmosphere to avoid unexpected reactions during ball milling. The ball mill parameters were 300 rpm for rotation speed and 1:10 for the mass ratio of powder to the balls. Subsequently, the milled powders were pressed mechanically at a pressure of 600 MP in uniaxial steel dies to produce pellets with diameter of 20 mm and 10 mm. The pellets were sintered at 1150 °C with approximate heating rate of 7 °C/min for 2 h in a laboratory electric furnace under argon media. All sintered pellets were mechanically polished with SiC paper in water, and then, ultrasonically rinsed with distilled water.

Experimental procedure

The chemical compositions of the composite surface were analyzed by EDXA, and its morphology was examined by LEO-1455VP SEM. The SBF was prepared to have ionic concentration near to human blood plasma based on the procedure developed by kokubo to investigate biocompatibility of Ti-HA composites [11]. The SBF was prepared by sequentially dissolving of NaCl, NaHCO₃, KCl, K₂HPO₄.3H₂O, MgCl₂.6H₂O, CaCl₂, Na₂SO₄ and tris buffer in 1 l of deionized water. pH of solution was adjusted to 7.4 by addition of 1 M HCl solution at 36.5 °C. Table 2 gives the ion concentration of SBF solution and its comparison with human blood plasma.

The wettability of the composites was determined by measuring of contact angle through sessile drop method. The wettability measurement was performed using data physicsoca 15 plus equipment (Filderstadt, Germany) with CCD camera at room temperature in air with 0.5 μ l droplet volume. The composite surface was polished by 1000 SiC paper in one direction and rinsed by ultrasonic in distilled water bath to remove any contamination. The contact angle was measured at three different points on the surface of composite and pure Ti sample.

Figure 2 exhibits the SEM micrograph of C2 and C3 samples. It is observable in Fig. 2 that 30% w/w HA content titanium composite has higher porosity in compare to 10% w/w HA content composite. Therefore, higher percentage of HA content lowers the density of the composite. TEM bright field microstructure images of C2 sample in Fig. 2c is the evidence of crashing Ti and HA particles to form nanocomposite materials. The pellets of Ti-HA composite were soaked in SBF solution with 40 ml/g ratio of solution volume (ml) to weight of powder sample (g) to investigation the microstructure. Therefore, three different series of pellets were immersed in plastic jars containing SBF with 0.1 cm²/ml ratio of surface

Fig. 1 SEM image of starting powders. a Nano-size HA. b Micron-size HA



Iable 1 I1-HA mixture combinations' characterizations									
Ti-HA mixture ID		C1	C2	C3	C4	C5	C6	C7	C8
Ti-HA mixture characterizations	HA particle size w/w (%) Milled time (h)	15 μm 30 50	150 nm 30 50	15 μm 10 20	150 nm 10 20	15 μm 30 20	150 nm 30 20	15 μm 10 50	150 nm 10 50

area (cm²) to solution volume (ml). Then, the jars were put into an air oven thermostat at 36.5 °C to soak the sintered Ti-HA powder samples. The sample were removed every 7 days to investigate the effect of soaking time on formation of apatite layer. The solution's pH was measured in order to investigate phase constituent and pH of SBF solution. Likewise, the powder sample of Ti-HA composite were soaked in SBF solution for 2 months for phase analysis with X-ray diffraction. The SBF solution refreshed every 24 h to accelerate the apatite layer formation, and also, to make easier identification of the apatite layer. The samples were retrieved after 7, 14, 21, 28, and 56 days in contact with SBF for characterization of constituent phase formed during immersion in SBF.

Results and discussion

Figure 3 shows the effect of immersion time on the pH value of SBF solution. The pH of solution was 7.4 before immersion which is similar to human body. At the early stage of the process, it can be seen that the pH value increases slightly as the immersion time increases. Initially, OH⁻ accumulation on the surface of the composite is responsible for increasing the pH value of the composite. At the later stage, the pH value reaches to its maximum value of 7.86, 7.71, and 7.60 for the samples containing 30% HA (w/w), 10% HA (w/w), and 0% HA (w/w), respectively. Longer immersion time causes the pH of the composition decreasing gradually. After 16 days' immersion time, the pH for pure Ti sample approaches the pH of human blood plasma, i.e., approximately 7.4. The pH of the sample continues to decrease slightly but the reduction is so small that may not harm the body. The pH value of Ti-HA 30% (w/w) composite was found to be higher than Ti-HA 10% (w/w) and pure Ti samples. Dissolving of Ca²⁺ ion by SBF solution results in increasing of pH in early stage of the immersion. Exchange of Ca²⁺ and H⁺ pileups the OH⁻ on the surface of composite, which is in favor of apatite nucleation. Accumulation of OH⁻ and PO₄³⁻ ions attract H⁺ and Ca²⁺,

and induce negative charge on composites' surfaces. This process proceeds through spontaneous reactions contain precipitation, nucleation, and growth of apatite layers on the surface of the composite [12–15].

Figure 4 demonstrates the representative XRD of Ti-HA composite before and after immersion in the SBF for various periods of time. The apatite phase concentration was not noticeable in XRD results even after 2 weeks since it was lower than the X-ray instrument detectability. Some weak peaks in graph B of Fig. 4 can be noticed for the apatite phase of pure Ti after immersion in SBF for 4 weeks, as compare to XRD result for pure Ti sample. The apatite peak related to Ti-HA composite becomes more intense after 16 weeks, whereas, some weak peaks related to apatite were observed for the pure Ti sample. The rate of increase in intensity of peak related to apatite phase in composite was more in 30% (w/w) HA composite compare to 10% (w/w) HA composite, regardless of milling time and initial HA particles size. This is the result of the greater apatite formation of the sample with higher HA content due to the higher concentration of Ca^{2+} and TiO_2 in comparison to the pure Ti sample or the 10 % w/w HA content composite.

Figure 5 shows the microstructure of Ti-HA composite before immersion in SBF. Figure 5 also gives evidence of non-homogenous distribution of α -Ti + Ca₃ (PO₄)₂, known as TCP, CaTiO₃, and Ti₅P₃ in the Ti-HA composite, which was discussed in the authors' previous article [16].

The reaction product of HA and Ti (consist of Ca₃(PO4)₂, CaTiO₃, and Ti₅P₃) formed during sintering process in Ti-30% w/w HA is accumulated in the outer layer of titanium particle (Fig. 5a). This can be attributed to higher degree of non-homogeneity in higher HA content composite. Images of the EDXA spectra of the composite in Fig. 7 reveals that high concentration of Ca, P, and O exists in pores and the intergranular spaces, while Ti is the dominant element in the middle of the grain.

Figure 6 shows SEM image of pure Ti sample and Ti-HA composite which immersed in SBF solution for different time

Table 2 Ion concentrations of SBF and human blood plasma (mmol/l)

Sample	Na ⁺	K ⁺	Ca ²⁺	Mg ²⁺	HCO ₃ ⁻	Cl	HPO4 ²	SO4 ²⁺
Blood plasma	142.0	5.0	2.5	1.5	27.0	103.0	1.0	0.5
SBF	142.0	5.0	2.5	1.5	4.2	147.8	1.0	0.5





span. From Fig. 6, it can be noticed that after 2 weeks the sphere-like apatite distributed irregularly with different sphere size and precipitate on the surface of composite. Figure 6d, f suggests that more apatite was precipitated for longer immersion time, such that after 4 weeks the surface of composite completely covered with layer of apatite (Fig. 6c and e). While no considerable apatite layer was found for pure Ti sample even after 2-week immersion, some sphere-like apatite were observed after 4 weeks' immersion. Unlike Ti sample, the apatite layer of Ti-HA composite can be distinguished easily after 2 weeks (Fig. 6d and f). However, precipitated layer of apatite in Ti-HA 30 wt% was more than Ti-HA 10 wt.%, regardless to the initial HA size and milling time.

The samples were examined for the pollution with related to the milling jar and balls. No contamination was found in the XRD results of the samples; however, EDXA analysis shows small trace of Cr and Fe in some regions of the sample with 50 h milling.

Figure 7 compares the mapping of elements on the surface of pure Ti, C2, and C8 samples. In case of pure Ti, small

amount of Ca was detected after 2 weeks which cannot be visible in SEM image because of low concentration. Nevertheless, more Ca were detected by EDXA in case of C2 and C8 samples. In Fig. 7b and c, irregular distribution of Ca can be found for Ti-HA composite, and hence, some regions within the microstructure can be distinguished as calcium-poor and calcium-rich areas. The selective precipitation of apatite seems to be as result of forming different phases in Ti-HA composite during sintering process [16].

The Ti rich region has lower bioactivity, and therefore, selective precipitation of apatite layer was done in areas with higher HA phases. Since higher concentration of ceramic phases are on the pores and inter-granular areas, the apatite layers tend to precipitated in these locations rather than on the middle of the grain with higher Ti phases. Table 3 lists the measured contact angle for different Ti-HA composite. Ti-HA composite with 30% *w/w* HA shows higher wettability than Ti-HA 10% *w/w* and pure Ti sample. Thereof, it is reasonable to conclude that the wettability of the composite will improve with increase in HA content.





Fig. 4 XRD spectra of sintered HA-Ti samples. a Pure Ti sample. b Pure Ti sample after 4 weeks immersion in SBF. c Sample C2. d C2 after 4 weeks immersion. e C2 after 8 weeks immersion. f Sample C7. g C7 after 4 weeks immersion. h C7 after 8 weeks immersion



Fig. 5 SEM image of **a** C8: TinHA 10% (*w*/*w*) (milled for 50 h.), **b** C4 Ti-nHA 10% (*w*/*w*) (milled for 20 h.), and **c** C6: TinHA 30% (*w*/*w*) (milled for 20 h) **Fig. 6** SEM image of **a** pure Ti sample after 1 week immersion, **b** pure Ti sample after 2 week immersion, **C** C3: Ti-mHA 10% (*w*/*w*) (milled for 20 h.) after 4 week immersion, **d** C3 after 2 week immersion, **e** C2: Ti-nHA 30% (*w*/*w*) (milled for 50 h) after 4 immersion, and **f** C2 after 2 week immersion



The chemical and topographical properties of the surface are important factor for biomedical materials (Fig. 8). Higher wettability improves the implant performance in the body [17].

The contribution of milling time and initial HA powder size is through determining the final surface roughness of the ceramic in the composite whose influence enhances the Ti-HA composite wettability. The wettability can be measured by contact angle of substrate with the given liquid according to the balance at solid, liquid, and vapor phases given by young equation:

$$\gamma_{sv} = \gamma_{sl} + \gamma_{lv} \text{Cos}\theta_{\gamma} \tag{1}$$



Fig. 7 SEM image. Mapping of Ca and EDXA related to a pure Ti sample, b C2: Ti-nHA 30% (*w/w*) (milled for 50 h.), and c C8: Ti-nHA 10% (*w/w*) (milled for 50 h)

where γ_{sv} , γ_{sl} , γ_{lv} , and θ_{γ} are interfacial tension of solidvapor, solid-liquid, liquid-vapor, and the contact angle, respectively.

It is worth to note that surface roughness will enhance the wettability caused by the chemistry of the surface

Table 3 Measured contact angle of Ti-HA composite and pure Ti	Sample	Mean value of contact angle	Standard deviation	
samples	c 1	23.93	1.74	
	c2	24.56	2.44	
	c3	33.73	2.90	
	c4	30.64	2.96	
	c5	26.75	1.54	
	c6	23.12	2.90	
	c7	45.01	3.43	
	c8	39.23	3.51	
	Ti	61.56	1.91	

defined by Wenzel [18], and it can be computed according to the equation in below:

$$\cos\theta_m = r \, \cos\theta_\gamma \tag{2}$$

where θ_m is measured contact angle, θ_γ is the contact angle on ideal surface (young contact angle), and *r* is the surface roughness ratio which can be calculated from a 3D roughness parameter and varied from below one for rough surface to one for smooth surface.

Ceramics are hydrophilic, and consequently, enhance the surface wettability [17]. Increasing HA content improves wetting properties due to the higher surface roughness induced by the structure of HA, and also, by higher amount of ceramic phase in Ti matrix. In conclusion, two factors will increase the wettability of Ti-HA composite: (a) ceramic phases which formed during sintering and (b) increase of the surface roughness induce by HA. The wettability was better in case of Ti-HA composite with 30% w/w HA than the other samples, regardless to its initial HA powder size and milling time. This is because of higher roughness of sample and higher amount of ceramic phases.

Fig. 8 Images of sessile droplets of a C6, b C8, c C4, and d pure Ti sample



Conclusion

A synthetic bone grafts was introduced based on titanium matrix composite reinforced with hydroxyapatite (HA) particle prepared by mechanical alloying and powder metallurgy method.

The in vivo bioactivity of composites was scrutinized by immersing the samples in simulated body solution (SBF). Results show that increasing HA may improve bioactivity of composite since the sample with 30% w/w HA content shows better bioactivity compatibility in compare to the 10% w/w HA and 0% w/w HA content samples.

The sample with 30% w/w HA shows better wetting properties than other samples, and therefore, it can be concluded that increasing HA content could improve wettability of composite. The contribution of milling time and initial HA powder size on in vivo bioactivity of Ti-HA composite are through determining the final surface roughness of the ceramic in the composite whose influence enhance the Ti-HA composite wettability. Results showed that the pH for pure Ti sample approach the pH of human blood plasma, which indicates possible osteoconductivity of the suggested synthetized Ti-HA composite.

Further studies on the effect of various bioceramic associated with different sintering techniques, such as spark plasma sintering (SPS), may offer better optimization for MMC applicable for synthetic bone prosthesis.

Compliance with ethical standards

Conflict of interest The authors declare that they have no conflict of interest.

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