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Fabrication and characterisation of a calcium phosphate/zirconia composite

M. K. Hilal¹, M. Y. Shareef¹ K. A. Maher^{*2} and R. Van Noort¹

¹Department of Restorative Dentistry, University of Sheffield, UK ²Department of Chemistry, Faculty of Science, University of Zakho, Iraq

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The aim of this study is to fabricate high strength calcium phosphate/ yttrium artially stabilised zirconia (Y-PSZ) composites using a simple powder pressing technique. Various compositions of hydroxyapatite® and zirconia powder©, ranging from 10 to 90 wt% Y-PSZ, were prepared using a wet mixing process. Ten compacts (13.5mm×3mm) were made for each composition, pressed and fired in air at sintering temperatures from 1100 to 1450°C for up to 12 hours. The bulk density, true porosity, linear shrinkage and biaxial flexural strength (BFS) were measured using standard methods. Microstructural changes were examined using XRD, DTA and SEM. The true porosity reduced and linear shrinkage increased with increasing sintering temperature and amounts of zirconia, whereas the firing time had little effect. The highest mean value achieved for the BFS was 269±20 Mpa for a composition of 70% Y-PSZ, fired at 1450°C for 6 hours, however with the hydroxyapatite (Hap) decomposed to α -TCP. XRD and DTA data show an interaction between the zirconia and the Hap at temperature lower than changes normally encountered for pure Hap. We conclude that sintering temperature and composition affect densification behaviour of calcium phosphate/Y-PSZ composites. The BFS increases with a reduction of porosity and increases with both sintering temperature and additions of zirconia. It was noted that there is considerable scope for improvement in the BFS values by reducing the porosity of these composites. Further work is needed to elucidate the microstructural changes.

Keywords: Fabricate, Zirconia, Hydroxyapatite, Composition, α-TCP. XRD, DTA, Temperature, Porosity.

INTRODUCTION

Hydroxyapatite (Hap) with the chemical composition $Ca_{10}(PO_4)_6(OH)_2$ has been extensively study as a bone substitute. It shows an excellent biocompatibility when implanted in either soft tissue (Ogiso *et al.*, 1982; Jansen *et al.*, 1985; van Blitterswijk et al., 1990), or hard tissue (van Blitterswijk et al., 1985; Jarcho et al., 1977; Denissen et al., 1980; de Groot, 981) and can form strong and intimate bond with bone (van Blitterswijk *et al.*, 1990; Jarcho *et al.*, 1977; Denissen *et al.*, 1980). Driskell *et al.* (Driskell *et al.*, 1973) was the first to report that a chemical bond exists between bone and Hap.

Hap, when sintered at high temperatures of up to 1250°C, was initially believed to be non-resorbable (Driskell et al., 1973). However, it is currently generally accepted that degradation of Hap can occur to a certain extent (van Blitterswijk et al., 1985; Jarcho et al., 1977; Denissen et al., 1980; de Groot, 981; Driskell et al., 1973; van Blitterswijk et al., 1989a). LeGeros, *et al.* (LeGeros et al., 1992) reported that the bioactivityof Hap may be related to its dissolution rate.

Hap is mechanically weak and unsuitable for use in stress bearing areas. Two approaches have been explored to overcome this problem (Ducheyne et al., 1980; Takagi et al., 1992).

One of the most promising applications of calcium phosphate technology revolves around the use of Hap

*Corresponding Author E-mail: Maher-333@hotmail.de

coatings conventional metallic prostheses. on Among the advantages of this approach is the potential for stronger, more permanent fixation of the metal implant directly to surrounding bone via Hap bone-bonding and the elimination of leaching of metallic trace elements into adjacent tissues. In the case of major orthopaedic appliances such as total hips and knees, a strongly adherent Hap coating on the metallic devices could obviate the need for methyl methacrylate bone cement, which has proved to be the limiting factor in long-term survival of such devices. However, all commonly used coating techniques depend on mechanical adhesion (rather than a chemical bond) to the underlying metal which makes this metal-Hap interface the weakest point of the system (Ellies et al., 1992; Jarcho, 1992) Ducheyne et al. (Ducheyne et al., 1980) used Hap as surface layer on titanium implants, but this approach has several problems that include separation of the coating layer from the underlying bioinert matrix and the coating process can reduce the strength of the bioinert materials (Takagi et al., 1992). In vivo studies (Hayashi et al., 1989; Gottlander and Albrektsson, 1991) have assessed bony adaptation to Hap-coated and non-coated pure titanium control implants. One study demonstrated an increased amount of direct bone to implant contact with coated specimen at six weeks (Gottlander and Albrektsson, 1991). Sharp edges or deep threads observed on the base metal also reduce the adherence properties of these ceramics. Another concern is the dissolution of improperly applied coatings with time. and consequences that both bond strength and coating integrity will be affected (Ellies et al., 1992; Hayashi et al., 1989; Gottlander and Albrektsson, 1991; Kay, 1992).

The second approach is to produce Hap/ Ceramic composite with better mechanical properties with Hap (Ducheyne et al., 1980; Takagi et al., 1992; Ellies et al., 1992; Jarcho, 1992; Hayashi et al., 1989; Gottlander and Albrektsson, 1991; Kay, 1992; Wu and Jeh, 1988) such as ZrO_2 -Hap, Ti-Hap, Al_2O_3 -Hap composites. The even distribution of the apatite phase as islets in a strong matrix may contribute to mineralization and direct bone apposition onto this type of ceramic composite (Wu and Jeh, 1988; Li et al., 1993; Takagi et al., 1992).

The objective of this study concerned with structural ceramics is the generation of materials having high reliability. To achieve this objective, there are two fundamentally different approaches: flaw toughening. The flaw control and control approach accepts the brittleness of the material and attempts to control the large extremes of processing flaws. The toughening approach attempts create microstructures that impart sufficient to resistance the strength fracture so that becomes insensitive to the size of flaws. The former has been the subject of considerable research that identifies

the most detrimental processing flaws, as well as the processing step responsible for those flaws (Large, 1989; Evans, 1982; Evans., 1989). The latter has emerged more recently, and has the obvious advantage that appreciable processing and postprocessing damage can be tolerated without compromising the structural reliability (Evans., 1989; Cook et al., 1985; Marshall, 1986).

Composite materials which combine the properties of a ceramic matrix and ceramic reinforcement or aggregate phase, otherwise known as ceramic matrix composites, are the subject of intense research interest. With careful processing improved strength and toughness can be obtained when compared to unreinforced ceramics. Composite such as zirconia toughened Al₂O₃ (ZTA), TiC-Al₂O₃ and SiC-Al₂O₃ are currently used as wear resistant bearings and cutting tools. A variety of ceramic composites are been explored for medical applications e.g. ZrO₂-HAp, Ti-HAp, Al₂O₃-HAp composites.

Molecularly designed composite materials that mimic the structure and properties of bone offer the greatest potential for solving the problems of interfacial stability and stress shielding of load-bearing prostheses. This is because the modulus of elasticity and fracture toughness of a composite is a function of many variables which can be controlled during processing. Various combination of a low elastic modulus bioactive materials, and a higher elastic modulus bioinert materials, can yield composites with properties that are similar to those of bone. This type of composite should eliminate stress shielding of bone because there is no mismatch in elastic modulus across the implant-bone interface. Use of a dispersed phase that has a bioactive property can ensure that rapid bone bonding occurs.

In a study perfomed by Jianguo et al. (Jianguo and Bahman, 1992), a range of high-strength hydroxyapatite composites (HAp/Oxide of Alumina, Titania or Zirconia) as well as pure hydroxyapatite, alumina, titania and zirconia were densified by hot isostatic pressing. The mechanical evaluation showed that the strength of the composite materials was 3-8 times higher than that of pure hydroxyaptite and close to that of the corresponding pure oxide. These ceramics were implanted in the femora of 16 rabbits for three months. Eight rabbits were sacrificed for histological evaluation and the others used for push-out test in order to study the bond strength between bone and ceramic implants. Histological evaluation under the light microscope showed direct contact between bone and ceramics for all ceramic materials studied except for alumina. The measured bond strengths in this experiment increased the following order: in alumina. titania, zirconia/hydroxyapatite, alumina/hydroxyapatite, titania/hydroxyapatite, hydroxyapatite. However, the authors did not give the sintering temperature to produce the fired ceramics, neither did they characterise their materials to show if there was decomposition of HAp or

interaction between HAp and other oxide and so it is not clear if the processed materials were genuinely composites of the two starting materials or their reaction product.

Kasuga, et al. (Kasuga et al., 1993) have reported on glass-ceramic (apatite/wollastonite) toughened а with zirconia which were prepared for widespread prosthetic applications. The strength of the composite increased with increasing zirconia content. This bioceramic exhibited extremely high bending strength (400-1000 MPa) for 30-80 vol.% zirconia. The bioactivity of these zirconia toughened glass-ceramic composites was evaluated by their bond strength to living bones, and it was found that the bond strength of composite containing 30 vol.% zirconia was as high as that of the glassceramic.

Recently investigators have demonstrated that the mechanical reliability of HAp can be improved by reinforcing HAp with Al_2O_3 (Hayashi et al., 1992; Huaxia and Marquis, 1992; Dimitrova-Lukacs and Gillemot, 1993; Dimitrova-Lukacs, 1994; Dimitrova-Lukacs, 1994; Rieu and Goeuriot, 1993). The main problems in producing dense composites in this system are connected with the big difference in sintering temperature of the components, the enhanced tendency of HAp to decompose during sintering and the consequent interaction between HAp and reinforced phase. An attempt at producing an HAp/Al₂O₃ composite (Terry et al., 1974) sintered at 1400°C in air, resulted in the formation of calcium aluminates. due to decomposition of hydroxyapatite into TCP and CaO with H₂O vapour. CaO then reacted with alumina to form calcium aluminates.

Several researcher have reported (Suda et al., 1990; Garvie and Nicholson, 1972; Garvie et al., 1975; Lange et al., 1986) on hydroxyapatite-zirconia composites with different volume ratio of zirconia particles. Most of these works reflect the difficulties of retaining the HAp and ZrO_2 phases in the composites due to difference in densification temperature (e.g. HAp start to densify at 1250°C, whereas ZrO_2 start to densify at 1400°C), and decomposition of HAp at 1400°C into TCP. As a consequence the desirable initial phases cannot be preserved in the composites unless expensive high technology processing at temperatures as low as 1250°C

Suda al. 1990) et (Suda et al., have produced composite ceramic contained either 1.6 or 50% of zirconia by volume addition to HAp, using cold isostatic pressure (CIP), then fired at 1300-1450°C in air atmosphere for one hour. Although the bending strength of their composite is 220-270 MPa, and the fracture toughness is 3.6-4.5 MPa.m^{1/2}, they did not characterise their materials to show if there is decomposition of HAp or interaction between HAp and zirconia. Since even high purity HAp decomposes into

tri-calcium phosphate (TCP) at 1250° C (Shareef et al., 1990), and the fact that they fired their materials at sintering temperatures higher than 1250° C, it has to be concluded that the materials obtained could not possibly be HAp/ZrO₂ composites.

Takagi *et al.* (Takagi et al., 1992) produced HAp/ZrO₂ composites using hot isostatic pressing (HIP) at 800-1150°C at 100 MPa for 2 hours. No phase change was found in zirconia nor in the HAp phase. The strength and the toughness achieved were respectively 190 MPa and 2.3 MPa.m^{1/2}. These value were approximately 20% and 100% higher than the corresponding value for hydroxyapatite ceramics without zirconia. Li *et al.* (Denissen et al., 1980) also used HIPing (1225°C at 160 MPa for 1 hour) to produce HAp/ZrO₂ composite that has excellent fatigue resistance in addition to high strength, but they did not characterised the fired composite.

The objective of this study is to fabricate and develop a bioactive, high strength and toughness calcium phosphate/zirconia composite, which can be used in load bearing applications.

MATERIALS AND METHODS

Hydroxyapatite (HAp) and yttrium partialy stabilised zirconia (Y-PSZ) powders were used in this study, supplied by Plasma Biotal Ltd, Tidswell, and Unitec Ceramics, Stafford, UK respectively.

The standard methods of XRD (CuK α radiation), SEM, BSI, EDX, PSA (laser diffraction method) and DTA have been used to characterise the starting powders, and to identify the crystalline phases of the fired samples. The bulk density, true porosity, linear shrinkage and BFS were determined for the fired samples.

The bulk density values were calculated using the following equation:

Bulk Density =
$$\frac{Weight (g)}{Bulk Volume (cm^3)}$$

.....(1)

Hence, Bulk Volume = $r^2h\pi$, where r and h are the compact's radius and height respectively.

The % theoretical density values were calculated using the bulk density of the compacts and the theoretical density value for HAp and zirconia of 3.156 and 6.05 gcm⁻³ respectively ^[39,40,41].

Theoretical Density (%) =
$$\frac{Bulk \ Density \ (gcm^{-3})}{Theoretical \ Density \ (gcm^{-3})} \times 100$$
(2)

Also, the true porosity values were calculated using the following formula:

True Porosity (%) =
$$1 - \left(\frac{Bulk \ Density \ (gcm^{-3})}{Theoretical \ Density \ (gcm^{-3})}\right) \times 100$$

.....(3)



Figure 1. The effect of sintering temperature on the % theoretical density of Y-PSZ (U) and Y-PSZ (M) samples, fired for 6 hrs an in air atmosphere.



Figure 2. The effect of sintering temperature on the % linear shrinkage of Y-PSZ (U) and Y-PSZ (M) samples, fired for 6 hrs in an air atmosphere

The linear shrinkage values of the fired compacts were determined by measuring the initial and the fired diameters of each sample as follow:

$$Linear Shrinkage(\%) = \frac{Di - Df}{Di} \times 100$$

.....(4)

Where, D_i and D_f are the initial and the fired diameter of the compact.

The BFS for a range of firing schedules of each HAp and Y-PSZ powder was determined for 10 test-pieces. The surface of the test-pieces were made flat by grinding with 600 grit SiC paper to avoid an uneven load. Each test-piece was placed on an annular knife edge of 9 mm diameter and then loaded in an universal testing machine with a cross-head speed of 0.5 mm/min (Lloyd M5K). The compacts were loaded to fracture and the maximum load was recorded. The BFS was calculated using the following formula (Shareef et al., 1994):

$$\sigma_f = \frac{p}{h^2} \{ 0.606 \ln (a/h) + 1.13 \}....(5)$$

where σ_f is the BFS, p is the load to fracture, a is the radius of the knife-edge support and h is the sample thickness.

RESULTS

Characteristics Properties of the Starting Powders

The particels size analysis (PSA) using laser diffraction method showed that the mean particle size of the partialy stabilized zirconia Y-PSZ (U) was 0.5 μ m and of Y-PSZ (M) was 1.6 μ m. Both powders were agglomerated as confirmed by the SEM micrographs. The XRD data for both ZrO₂ powders shows that they are consist of a mixture of tetragonal (t), monoclinic (m) and cubic (c) phases, with the latter two phases representing a major proportion, whereas the characteristics properties of the starting powders of HAp have been reported previousely (Shareef et al., 2011).



Figure 3. The effect of sintering temperature on the BFS of Y-PSZ (U) and Y-PSZ (M) samples, fired for 6 hrs in an air atmosphere.



Figure 4. Theoretical density, Linear shrinkage and BFS plotted as a function of sintering temperature for samples prepared from composite of 45/55 vol.% HAp/Y-PSZ (U) fired for 6 hrs.



Figure 5. Theoretical density, linear shrinkage and BFS plotted as a function of sintering time for samples prepared from composite of 45/55 vol.% HAp/Y-PSZ (U) composite fired at 1450°C.

Sintering Behaviour of the Fired Samples

The sintering behaviour of the fired HAp samples for thetheoretical density, linear firing shrinkage and the BFS have been reported previousely by Shareef *et al.* (Shareef et al., 2011), whereas for the fired samples prepared from Y-PSZ (U) and (M) used in this study are presented in Figures. 1, 2, and 3. These were found ncrease with increasing the sintering temperature. The sintering behaviour of the fired HAp/Y-PSZ composite for the theoretical density, linear firing shrinkage and the BFS values were plotted against sintering temperature, time and addition of Y-PSZ as shown in Figures 4,5 and 6. These were found increase with increasing sintering temperature, time and addition of zirconia. As shown in Fig. 6, the highest mean



Figure 6. Theoretical density, linear shrinkage and BFS plotted as a function of zirconia addition, fired at 1450° C for 6 hrs.

Table 1. BFS and % theoretical density data of composite prepared from composition of 30/70 vol.% HAp/Y-PSZ (U).

Parameters	HAp/Y-PSZ (U)	HAp/Y-PSZ (M)
BFS (MPa)	269 ± 20	176 ± 10
% Teoretical Density	95 ± 0.5	85 ± 0.8
% True Porosity	5	15



Figure 7. Fracture toughness of hot pressed HAp, 30/70 vol.% HAp/Y-PSZ (U) composite and Y-PSZ (U) samples.

value achieved for the BFS was 269 ± 20 MPa for a composition of 70% Y-PSZ (U), fired at 1450° C for 6 hours, because the densification process reached to the higest fired density and resulting lower porosity values as shown in Table 1. However, with the hydroxyapatite (HAp) decomposed to α -TCP. On the other hand the fracture toughness values for the hot pressed samples prepared from HAp, HAp/Y-PSZ composite and Y-PSZ are plotted against the materials used as shown in Figure. 7. Also, the main values of the BFS, fracture toughness and the fired theoretical density are shown in Table 2.

The XRD and DTA data for the fired composite are indicated that there is an interaction between the Y-PSZ and the HAp. However, the BSI examination and EDX analysis for polished section of composition of 30% HAp and 70% Y-PSZ, fired at 1450°C for 6hrs, shown that an aggregated crystalline regions represented the calcium phosphate phase surrounded by the Y-PSZ materials, which is conformed **Table 2.** The main values of BFS, fratures toughness and the fired theoretical density of hot pressed HAp, HAp/Y-PSZ (U) composite and Y-PSZ (U) ceramics.

Parameter	НАр	30/70 vol.% HAp/Y-PSZ	Y-PSZ
BFS (MPa)	142±17	269±19	1213±223
Fracture Toughness (MPa.m ^{1/2})	1.3±0.1	3.8±0.4	8.2±1.0
%Theoretical Density	94±4	95±0.5	99±0.8

by the EDXanalysis.

DISCUSSION

lt has been found that, by increasing the sintering temperature, time and addition of Y-PSZ, the theoretical density and linear shrinkage values were increased for the HAp/Y-PSZ composites. This cause an increase in the BFS values because the densification process increased with increasing these factors (Figures. 1, 2 and 3). The same observation has been found by Takagi et al. (Takagi et al., 1992), which is stated that the densification behaviour of HAp/ Y-PSZ composite increased with increasing firing temperature and the amount of zirconia content. The development of HAp toughened materials by dispersed zerconia particles has been found difficult as reported by Evans (Evans, 1982; Evans., 1989) and Wu et al. (Wu and Jeh, 1988). One major obstacle has been found the reaction between the matrix and dispersed particles during the sintering process. This was conformed by the XRD and DTA data for the fired composite. Calcium which is the major constituent of HAp, diffuse into Y-PSZ and change it to stable which cubic phase, for а transformation toughening mechanism is not expected to occur as reported by Takagi et al. (Takagi et al., 1992). To avoid the difficulty associated with the diffusion, temperature for densification has to be the minimised. However, this is difficult because the composites generally have inferior sintering characteristics, and require high temperature for densification.

Takagi *et al.* (Takagi et al., 1992) achieving considerable reduction of densification temperature in a composite ceramics containing Y-PSZ particles dispersed in HAp using combined application of a colloidal pressing technique and densification by hot isostatic pressing.

The calcium phosphate/Y-PSZ composite obtained in this study was contained approximately 5% porosity. Further work is needed by using a hot pressing technique and reducing the particles size to enhance the uniformity of the microstructures of the composite. This will eliminate the porosity on one hand and to reduce the sintering temperature on the other hand to level which keeps HAp thermally stable at certain sintering temperatures as reported earlier by Shareef *et al.* (Shareef et al., 2011).

CONCLUSIONS

- 1. Sintering temperature and addition of zirconia were affected the densification behaviour of HAp/ Y-PSZ composite, whereas the firing time had little effect.
- 2. The BFS values were increased with an increase both sintering temperature and additions of zirconia.
- 3. The XRD and DTA data suggested that there was interaction between the zirconia and the HAp or TCP.
- 4. There is considerable scope for improvement in the BFS and fracture toughness values by reducing the porosity and the particles size of these composites.

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