
Hydroxyapatite coating on porous zirconia
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Abstract

Since hydroxyapatite has excellent biocompatibility and bone bonding ability, porous hydroxyapatite ceramics have been intensively studied. However, porous hydroxyapatite bodies are mechanically weak and brittle, which makes shaping and implantation difficult. One way to solve this problem is to introduce a strong porous network onto which hydroxyapatite coating is applied. In this study, porous zirconia and alumina-added zirconia ceramics were prepared by ceramic slurry infiltration of expanded polystyrene bead compacts, followed by firing at 1500°C. Then a slurry of hydroxyapatite-borosilicate glass mixed powder was used to coat the porous ceramics, followed by firing at 1200°C. The porous structures without the coating had high porosities of 51% to 69%, a high pore interconnectivity, and sufficiently large pore window sizes (300μm–500μm). The porous ceramics had compressive strengths of 5.3~36.8MPa and Young’s moduli of 0.30~2.25GPa, favorably comparable to the mechanical properties of cancellous bones. In addition, porous hydroxyapatite surface was formed on the top of the composite coating, whereas a borosilicate glass layer was found on the interface. Thus, porous zirconia-based ceramics were modified with a bioactive composite coating for biomedical applications.

Keywords: Zirconia; porosity; hydroxyapatite; coating; compressive strength

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1. Introduction

Porous bioactive ceramics such as hydroxyapatite are attractive for bone regeneration or reconstruction due to their bone bonding ability and good bone ingrowth property. However, porous hydroxyapatite ceramics are extremely weak. On the other hand, porous zirconia is relatively strong and tough compared to other porous bioceramics, but has the problem of biological inertness to bone tissues. Therefore, it is desirable to combine the mechanical properties of zirconia with the bioactivity of hydroxyapatite. Various studies have indicated the difficulty in obtaining dense zirconia-hydroxyapatite composites due to poor sinterability [1] and the decomposition of hydroxyapatite [2]. Thus it is not feasible to produce useful porous zirconia-hydroxyapatite composite scaffolds. This could be the reason why many studies have turned to hydroxyapatite coatings (or bioactive ceramic coating) on porous zirconia (or bioinert ceramics).

When coating porous zirconia with hydroxyapatite, two problems should be considered. Firstly, hydroxyapatite coatings on sintered porous zirconia would result in poor interfacial bonding due to the sintering shrinkage occurring during heating and the thermal shrinkage due to cooling. To
improve the interfacial strength of the hydroxyapatite coated porous zirconia, a lesson can be learnt from dense zirconia implants which have been coated with bioactive glass [3] or glass-ceramic composite [4]. Secondly, plasma spraying technique cannot be used to coat porous ceramics, although it is a good method for dense ceramic substrates. However, coatings on porous ceramics can be applied by methods of sol-gel coating, ceramic slurry dipping, or biomimetic approach using a simulated body fluid. For example, Kim et al. [5] studied porous zirconia that were coated with multilayer composite coatings consisting of fluorapatite (FA), hydroxyapatite, and tricalcium phosphate (TCP) with the FA as an intermediate layer.

However, there are few studies on hydroxyapatite coating on porous zirconia with the coating being modified by a glass phase. Our previous work [6] suggested that boerisilicate glass (bioinert and with a low melting point) – hydroxyapatite composite could be used as the coating material for porous zirconia. The purpose of this study was to prepare porous zirconia using an in-house developed method, correlate the mechanical properties with the porous structures, and make the porous zirconia ceramics more useful by coating with a bioactive hydroxyapatite–borosilicate glass coating. In the study, borosilicate glass was added into the HA matrix to enhance the interfacial bonding between the coating and the substrate. Borosilicate glass was used because it was readily available, biocompatible, and bioinert.

2. Experimental procedure

Starting materials
The starting materials used to prepare the porous bioceramics were 3mol% yttria stabilized zirconia powder (Aldrich Chemical Company, Inc.) with an average particle size of 0.265μm and a specific surface area of 4.01 m²/g. Alumina powder (AKP-30) (Sumitomo, Tokyo, Japan) with an average particle size of 0.357μm and a specific surface area of 6.28m²/g was also used. Ammonium salt of polymethacrylic acid (NH₄PMAA) solution containing 75wt% water and 25% active substance, with molecular weight of 12000 and pH 8 (Darvan C, R. T. Vanderbilt) was used as a dispersant for ceramic slurries. HCl (0.1 M) and NaOH (0.1M) were used to adjust the pH of the slurries. Expanded polystyrene (EPS) beads (Styropor® from BASF Singapore Pte. Ltd) were used as a polymer preform. The average particle sizes of the EPS beads were 2.95mm, 2.58mm, 2.18mm, 1.85mm, and 1.40mm. The starting materials used for coating were hydroxyapatite (HA) powder (Riedel-de Haën®) with an average particle size of 0.11μm and a Pyrex® borosilicate glass powder with a particle size of 0.50μm. The borosilicate glass had the following standard composition: 81 wt% of SiO₂, 4 wt% of Na₂O, 0.5 wt% of K₂O, 13 wt% of B₂O₃, and 2 wt% of Al₂O₃. The softening point of the glass was 821°C.

Preparation of uniform porous structures
An in-house designed apparatus for vacuum casting was used to prepare porous structures. The apparatus is schematically shown in Fig. 1. The plastic plate with holes as the bottom was used to support the gypsum plate as a filter. The holes in the plastic plate were to remove the water in the ceramic slurry. The EPS beads were packed in a plastic container before casting the slurry. Another plastic plate with holes was used to cover the EPS beads. The plate was then pressed by tightening the two sets of screws and nuts. The pore sizes of the final porous ceramic structures were controlled by using different sizes of EPS beads: 2.95mm, 2.58mm, 2.18mm, 1.85mm, and 1.40mm and the porosities of the porous structures were controlled by adjusting the screws and nuts. In the casting process, yttria stabilized zirconia (TZP) or 20 wt% Al₂O₃ added TZP (ZA20) slurry with a solids loading of 50 wt% and a PMAAA content of 0.5% and 0.7%, respectively, was poured into the plastic container so that the interstices between the beads were filled with ceramic particles. The green body was then dried in air for 2 days and sintered at 1500°C for 2 hours. A heating rate of 1°C/min. Was used for heating till 500°C.
Preparation of HA/glass coating on porous TZP substrate

HA - 50 wt% glass mixed powder slurry with a solids loading of 25 wt% was prepared by planetary ball milling at 100 rpm for 2 hours. A dip-coating process was used for the HA-glass coating. Specifically, the porous TZP samples were immersed into the HA-glass slurry for 1.5 minutes. The samples were then centrifuged to remove the extra slurry. After drying in air, the coated samples were sintered at 1200°C in air for 1 hour.

Characterization

The total porosity and the open porosity of the porous structures were measured by a geometric method and via wax infiltration. The morphology of the porous structures was observed by a stereozoom microscope (Leica MZ6). Scanning electron microscopy (SEM; JEOL, JSM-6310 LV) was used to examine the as-sintered surfaces and the as-polished coating layers on porous TZP substrates. The mechanical properties of the porous structures were studied by an Instron Tester (Model 5567) using the crosshead speed of 0.5 mm/min. The compressive strengths of porous TZP and ZA20 ceramics sintered at 1500°C with various pore sizes and porosities were measured using rectangular blocks with dimensions of 10 x 10 x 20mm³.

3. Results and discussion

3.1 Morphology of uniform porous TZP

Variation in pore size

The infiltration of a packed EPS bead bed with the TZP slurry resulted in porous TZP after sintering and the porosities of the porous TZP structures with different pore sizes were about 69%, depending on the applied load on the EPS beads. Porous structures with different pore sizes were obtained by using EPS beads of different sizes. Fig. 2 shows the porous TZP structures with two different pore sizes. The spherical macropores in the porous TZP ceramics were basically interconnected through pore windows of much smaller sizes, which were typically from 300 μm to 500 μm, estimated from the SEM images. Thus the obtained porous TZP ceramics had a high porosity and large enough pore window sizes. Since the porous structures were designed for bone ingrowth, the pore window (or neck) sizes were important. The oversized pore windows could allow subsequent coating of the bioactive hydroxyapatite layer, which would make the pore window sizes smaller. However, the bioactive layer coated porous zirconia should have a pore window size larger than 100 μm, as various studies have indicated optimal pore sizes of 100 μm to 200 μm for bone ingrowth. The proper pore size of a porous bioceramic material should also be determined by considering the mechanical properties such as compressive strength.

Variation in porosity

By adjusting the applied load on the bed of the EPS beads, the deformation of the EPS beads or in effect the contacting area between the EPS beads could be controlled and thus uniform porous structures with different porosities and different pore window sizes could be prepared. Fig. 3 shows the porous TZP structures with a macropore size of 1.29mm but two different porosities. From the pictures, one can see that with the increase of the porosity, the struts of the porous structure became thinner. Furthermore, the porous structure with a lower porosity had less uniformly distributed pores. However, with the increase of the porosity, the macropores became better distributed, the cell edges became more slender and more uniform, and the whole porous structure became more foam-like. The pore window size also increased with the increase of porosity, ranging from less than 100 μm to about 500 μm. Thus, too low a porosity was not proper as it resulted in too small a pore window size or pore isolation. On the other hand, if the porosity was too high, the interstices
between the contacting EPS beads became too small and thus it became more difficult for ceramic slurry infiltration. Thus the porosity should be also proper and again the final compressive strength of the porous structure should be considered.

3.2 Compressive strength versus porous structure

**Effect of pore size on compressive strength**

Compressive strength and compressive modulus of the porous ceramics were measured for the project, but only the compressive strength results are presented. Fig. 4 shows the effect of pore size on the compressive strength of the porous TZP and porous ZA20 structures sintered at 1500°C and with a porosity of 51%. From the figure, one can see that the compressive strength for the TZP and the ZA20 porous structures decreased with the increase of the pore size. For the porous TZP ceramics, the compressive strength decreased from 23.6MPa to 15.9MPa, when the average pore size increased from 0.98mm to 2.09mm. For the porous ZA20 ceramics, the compressive strength decreased from 36.8MPa to 21.8MPa, when the average pore size increased from 1.03mm to 2.22mm. In spite of the large pore sizes, the present porous ceramics had rather high compressive strengths due to the intrinsically high strength of the materials and also due to the dense ceramic struts as a result of the processing conditions.

Although it is not easy to understand why larger pores led to lower strength, this trend was repeatedly observed by the authors. In addition, Liu [7] reported a similar compressive strength-pore size relationship in porous hydroxyapatite ceramics. However, Bose et al. [8] showed that the strength of the porous alumina and the porous tricalcium phosphate structures (44% porosity) did not flow a trend of monotonous increase or decrease. In other words, when the pore size was too small (305 μm), the compressive strength was decreased. This decreased strength could be due to the incomplete infiltration of the ceramic slurry, resulting in a porous structure with a large number of defects. The decrease of compressive strength due to increasing pore size was also found in porous polymers and porous metals. For example, Ma et al. [9] noticed the increase of compressive modulus of porous poly (L-lactic acid) (PLLA) foam (94.5% porosity) with the decreases of pore size (from ~450 μm to ~300 μm). On the other hand, for porous magnesium foam (45% porosity), when pore size varied from 100 μm to 400 μm, the compressive strength changed from 16 MPa to 10 MPa [10].

**Effect of porosity on compressive strength**

The compressive strengths of the uniform porous TZP and ZA20 ceramics with similar pore sizes (~1.3 mm) but different porosities (from 51% to 69%) are shown in Fig. 5. From the figure, one can see that the compressive strength decreased with the increase of porosity. Specifically, for the porous TZP ceramics, the compressive strength varied from 23.6MPa to 3.3MPa, whereas for the porous ZA20 ceramics, the compressive strength varied from 36.8MPa to 5.3MPa. The mechanical properties of these porous TZP and ZA20 ceramics were higher than those of cancellous bones (compressive strength 2 - 12MPa).

Dependence of compressive strength on porosity has been established by a few models. For highly porous ceramic structures with interconnected pores, Gibson and Ashby [11] considered the macrofracture (crushing) of a porous structure as a result of the bending microfracture of the struts and derived the following formula:

\[
\sigma = \sigma_0 (1 - P)^{3/2}
\]

(1)

where \(\sigma\) is the compressive strength of a porous structure at porosity \(P\) and \(\sigma_0\) is the bending strength of the strut material, which may contain micropores. It is obvious from equation (1) that increasing porosity leads to decreasing compressive strength.
3.3 Effect of alumina addition on zirconia

In the current study, not only pure porous TZP but also porous TZP - alumina composite were prepared. Studies on dense TZP - alumina composites indicated that alumina could improve the mechanical strength and fracture toughness of pure TZP. As shown in Fig. 4 and Fig. 5, the compressive strengths of the porous ZA20 ceramics were higher than those of the porous TZP structures. The effect of alumina was mainly because of the thermal expansion coefficient mismatch between the TZP and alumina phases, which caused tensile residual thermal stresses in the composite. The residual stresses tended to reduce the critical stress level required for the tetragonal to monoclinic zirconia phase transformation, which is a well-known mechanism of strengthening and toughening for ceramics.

3.4 Microstructure of the HA/glass composite coating

Fig. 6 shows a coated porous TZP ceramic with a pore size of 1.54mm and a porosity of 64%. The HA/glass coating layer was about 50 μm thick and attached to the pore wall surface of the porous TZP structure. Although the coating layer was relatively thick, the pore windows (up to ~ 500 μm) were not blocked. Fig. 7 shows the top surface of the HA/glass coating on the porous structure. The top surface was porous and rich in hydroxyapatite phase. However, the interior part of the coating layer was dense due to the existence of borosilicate glass (softening point 821°C), which melted at the sintering temperature (1200°C) and wetted the TZP pore wall surfaces. Thus, the borosilicate glass rich layer served as an intermediate layer between the bioinert zirconia and the bioactive hydroxyapatite coating. Indentation results indicated good interfacial bonding between the glass coating and the zirconia substrate, as indentation cracking occurred only within the glass layer rather than along the interface.

4. Conclusions

Porous TZP and ZA20 structures had high porosities of 51% to 69%. Most macropores were interconnected with pore sizes of 1-2 mm and with pore window sizes of 300 – 500 μm. The compressive strength and Young’s modulus of the porous TZP and ZA20 ceramics all decreased with the increase of pore size and porosity. Depending on pore size and porosity, the porous TZP ceramics showed compressive strengths of 3.3~23.6MPa and Young’s moduli of 0.24~2.20GPa. On the other hand, the porous ZA20 ceramics had compressive strengths of 5.3~36.8MPa and Young’s moduli of 0.30~2.25GPa. These data were similar to or better than the mechanical properties of cancellous bones (2~12MPa for compressive strength and 0.2~5GPa for tensile modulus).

The porous TZP structures were further coated with an HA/borosilicate glass composite layer. The average thickness of the coating layer was 50μm. Indentation tests demonstrated good interfacial bonding between the coating layer and the TZP substrate. Porous HA surface was formed on the top of the coating and dense borosilicate glass was found in the interior of the coating layer due to the low softening point of borosilicate glass (821°C) and the high sintering temperature (1200°C). Thus, porous TZP ceramics with various pore sizes and porosities could be modified by the bioactive composite coating.

References


Figure captions

Fig. 1 In-house designed apparatus for preparing the porous ceramics.

Fig. 2 Stereo-optical micrographs showing the porous TZP structures sintered at 1500°C that had a porosity of 69% but different average pore sizes: (a) 1.54mm and (b) 0.98mm.

Fig. 3 Stereo-optical micrographs showing the porous TZP ceramics sintered at 1500°C that had a pore size of 1.29mm but different porosities: (a) 62% and (b) 69%.

Fig. 4 Compressive strength versus average pore size for porous TZP and ZA20 structures sintered at 1500°C and with a porosity of 51%.

Fig. 5 Compressive strength versus porosity for porous TZP and ZA20 ceramics sintered at 1500°C.

Fig. 6 SEM micrograph showing the HA/glass coating on a porous TZP ceramic.

Fig. 7 SEM micrograph showing the as-sintered surface of the HA/glass coating layer on the porous TZP structure.
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