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CELL-DEGRADATION OF CALCICUM PHOSPHATE CERAMICS

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Abstract

Calcium phosphate ceramics are used in bone surgery under different forms: dense or porous ceramics as bone substitute, thin ceramic coatings on metallic implants as an osseointegration enhancer. Their degradation depends on their physico-chemical properties and particularly on their chemical composition. Natural calcium phosphates of bone are degraded by mononuclear or multinuclear cells and the extracellular matrix induces the differentiation of the degrading-cells. Hydroxyapatite, which is one of the most used calcium phosphates, is known as a low degradation material. However, the histological analysis of implanted HA-materials both in animals and in humans showed that a cellular degradation took place on the surface of the material.

Key Words: Calcium phosphate ceramics, hydroxyapatite, hydroxyapatite-coatings, macroporous ceramics, cell degradation, osteoclast, macrophage, remodelling, dissolution.

Introduction

Hydroxyapatite (HA) ceramics are used daily in human and animal surgery under different forms [31, 36]. This material is classified as a surface-active biomaterial [33]. It means that an interfacial bond is formed between the material and the tissue in which it is implanted. Calcium phosphate materials, currently used in human surgery, are assumed by surgeons to be non- or slowly degradable materials. Depending on its use, a degradation may be necessary to improve osseointegration [20].

HA-materials such as HA-coatings and macroporous-HA can be considered to have a chemical composition which is close to bone mineral [42]. All other characteristics are radically different. The macroporous ceramic can be composed of a single calcium phosphate phase. HA-coatings can contain other calcium phosphate phases in their amorphous phase [61]. The size of the crystallites of bone is much smaller than the size of ceramic grains, and the structure and the texture of bone and ceramics are totally different [51]. Despite these differences, based on its close chemical composition to that of the bone mineral, HA is assumed to be a bio-degradable material.

The purpose of this article is to discuss some observations made on HA ceramic degradation in a biological environment.

Calcium phosphate ceramics

Ceramic material can be defined as all solid materials which are neither metal nor organic. Ceramics are heterogeneous material constituted by grains which are assembled using a thermal process (sintering stage). They can be used in the form of bulk or porous material obtained by slip casting or machining. Moreover, thin coatings of calcium phosphate ceramic on metal alloys, polymers and composites can be obtained by plasma-spraying of HA-powder. The characteristics of the ceramic will have a great influence on the resorption of the material. Moreover, different calcium phosphates can be used to manufacture bioceramics. The HA and the β-tricalcium phosphate (β-TCP) are the most commonly used. The choice between these two compounds depends
Figure 1. X-ray diffraction patterns of: HA-powder before plasma-spraying (A), and of the obtained HA-coating (B). The peaks are broader after the plasma-spraying indicating the presence of an amorphous phase after the spray.
on the solubility required for the material. HA is considered to be quite insoluble and β-TCP soluble when tested in vitro.

Different phases can appear in the HA-ceramics according to the sintering protocol adopted and the purity of the powder used to manufacture the ceramic. β-tricalcium phosphate (β-TCP) can appear during the manufacturing process of HA-ceramics. An amorphous phase, which has a Ca/P ratio identical to the Ca/P ratio of HA but which cannot be identified as HA by X-ray diffraction, can also be found in HA-ceramic coatings. These phases do not have the same solubility as pure crystalline HA. The crystallinity of the ceramic is measured by X-ray diffraction (XRD). The broadening of the XRD pattern peaks after plasma-spraying is representative of the amount of amorphous phase in the ceramic. Amorphous phase which can contain other calcium phosphate phases, such as β-TCP or calcium oxide, is more soluble than the crystalline phase [61] (Fig. 1).

The porosity of the material is often divided into microporosity and macroporosity. The microporosity is due to the small spaces existing between the grains of the ceramic which were left by the shrinkage occurring during the material diffusion process at the grain boundaries (Fig. 2). They have a diameter of less than a few micrometers and are not interconnected. The macroporosity results from the shaping of the ceramic done in order to obtain a bone tissue ingrowth inside the ceramic. The macropores should be interconnected and have a diameter of from 100 μm up to several millimeters. The porosity of the material determines the size of the surface which is in contact with the body fluid. The surface in contact with the tissue depends also on the interconnection of the material pores. It is generally admitted that a pore diameter of more than 100 μm is necessary for bone ingrowth [30] depending on the volume of the implanted ceramic. If the interconnection of the pores is poor, the size of the exposed surface will be smaller than if the pore connection is high.

The size and the shape of the grains obtained after the sintering stage or plasma-spraying can vary and have an influence on the amount of the surface of the crystalline phase exposed to the body fluids.

Degradation of bone calcium phosphate

Bone mineral of the adult is composed of an A and AB carbonated non-stoichiometric apatite [42]. Nucleation of carbonated apatite crystals within the bone extracellular matrix can be due to the removal of biological inhibitors, to interaction between Ca²⁺ and calcium-binding proteins, or to apatite precipitation out of locally supersaturated calcium phosphate solution made by body fluids inside some microdomains [4, 27, 45, 58]. Bone mineral crystals are first formed either inside "the holes" existing between the fibrils in the collagen bundle or inside extracellular matrix vesicles synthesized by osteoblasts. The bone mineral nucleation taking place inside these organic compartments is an heterogeneous secondary nucleation controlled by the molecules contained inside these compartments. Subsequently, the mineralization spreads out of the nucleation site throughout the extracellular matrix, this, making the bone an organo-mineral composite.

A high degradation rate of the organo-mineral composite constituting the bone tissue by bone cells is already known in several pathological and physiological processes. Osteoclasts [12, 44] are known to resorb bone tissue during remodelling phenomena which occur during the life of the bone. Bone tissue is resorbed by osteoclasts or giant cells, during Paget's disease. Some forms of osteoporosis, idiopathic osteolysis, and hyperparathyroidism are also characterized by the presence of many osteoclasts resorbing bone tissue. From the histological studies carried out in these pathologies, two distinct kinds of cells were shown to degrade bone tissue and calcium phosphate mineral: osteoclasts and mononuclear cells [13]. Histological analysis of bone allograft shows, in the same resorption sites, mononuclear cells, giant cells, and osteoclasts.

Osteoclasts: Osteoclasts are multinucleated (2 to 50 nuclei) giant cells (20 to 100 μm). Once in contact with the bone, the newly formed osteoclast begins to erode a cavity of characteristic shape and dimension. They have a central ruffled border with many cytoplasmic folds, surrounded by a clear zone where the cell is closely attached to the bone surface, forming a temporary seal around the site of bone destruction [17]. Cell attachment could be mediated by oligosaccharides [48]. The chemical processes mediating bone resorption are
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numerous and complex. Dissolution of the mineral is the main task of the osteoclast which produces biochemical changes that includes release of acid, particularly lactic acid and citric acid [43], coupled with increased synthesis and release of lysosomal enzymes and other proteases [32]. Carbonic anhydrase, which hydrolyses carbonate into proton and carbon dioxide, plays a role in the acidification of the medium at the interface of bone and osteoclasts. The ruffled border membrane contains a protein identical to H^+ -pump found in the lysosomal membrane [7]. The decrease of the pH induced by the proton-pump in the extracellular compartment facing the ruffled border seems to be responsible for the bone mineral degradation. Lysosomal enzymes secreted in this compartment could help to degrade the organic matrix. Furthermore, collagen fibrils could be degraded by other cells containing large amounts of collagenase such as osteoblasts [32] although osteoclasts could resorb the organic and inorganic components of bone as reported by Blair et al. [9]. However, the digestion of the collagen coating, existing on bone surface, by osteoblasts and lining cells could be necessary for osteoclast digestion of bone [15]. Complement (C₄) adsorption on the exposed mineralized bone surfaces could mediate the recruitment of mononuclear preosteoclasts to this site [46]. Osteoclasts have been known for a long time to be involved in relation with osteoblasts in the normal remodelling of bone tissue. Osteoclast precursors may respond to specific humoral signals emanating from other bone cells such as active osteoblasts, lining cells, or immune cells synthesizing interleukin 1α and β, tumoral necrosis factor α and β, and transforming growth factors α and β [5, 55].

Osteoclasts originate from hemopoietic tissue. They derive from mononuclear phagocytes as precursors. They arise from a derivative of the hemopoietic stem cell that can become committed specifically to osteoclast generation [14]. The separation between osteoclasts and macrophages or giant cells lineage could come early [11]. Alternatively, it remains possible that osteoclasts could develop from mature mononuclear phagocyte by induction of osteoclastic differentiation in the special microenvironment of bone surface. Osteoclast phenotype has been well characterized and shown to express many of the antigens of the monocyte-macrophage lineage [5]. Mononuclear cells identified in histological sections of bone, applied to the bone surface, show a similar immunophenotype to that of the multinucleated osteoclasts [6].

Marrow-derived giant cells express osteoclast-specific cell surface antigens when cultured with calvariae. In the mesenchymal cell containing chorioallantoic membrane, intact bone matrix induced the formation of giant cells that express osteoclast-specific antigens [54]. De-mineralized matrix and matrix components such as type I collagen, osteocalcin, osteopontin, osteonectin, bone-derived growth factors, cartilage proteoglycan, hyaluronic acid, laminin, and fibronectin do not induce the formation of giant cells expressing osteoclast markers (antigen 121 F) on their membrane [54]. In the same experiment, a culture medium, conditioned by chick hatched calvariae, induced the formation of giant cells having osteoclast markers from the culture of a monocytic or marrow mononuclear cell population. It showed that the terminal stages of osteoclast differentiation were influenced by soluble factors found in the bone environment. Glowacki et al. [28] showed that osteocalcin-deficient bone particles were not associated with osteoclasts when implanted subcutaneously in rats. However, in the same experiment, it was shown that synthetic microcrystalline HA recruited multinucleated cells lacking the osteoclastic features [28].

Thus, osteoclast recruitment at the bone contact depends on specific or non-specific humoral signals and factors emanating from bone extra cellular matrix (ECM). However, ECM calcium phosphate materials can have an influence on osteoclast recruitment mediated by non-specific humoral signals such as C₃ adhesion on the naked calcium phosphate surface.

**Other cells:** It is not clear whether macrophages contribute directly to bone remodelling in vivo. Evidence has been presented that cells of the mononuclear phagocytic system are capable of bone resorption in various culture system. Cells of the mononuclear phagocytic system were shown to resorb implanted isogenic bone matrix on animal. These cells are also present at the point of contact of bone tissue allograft in humans. Mononuclear cells could be responsible for the digestion of the bone collagen [32] and a part of the erosion cavity dug by the osteoclast [21]. It is unclear if these cells are precursor cells having failed to fuse or resorbing monocytes [52] helping osteoclasts by the release of cytokines.

Furthermore, macrophages are interrelated with osteoblasts and synthesize cytokines which are able to regulate their activity [34, 56].

**Degradation of ectopic natural HA-crystals**

Ectopic calcium phosphate crystals found in soft tissues induce a cellular response involving phagocytic cells. Calcium oxalate and dicalcium pyrophosphate deposits in soft tissues are surrounded by histiocytes and giant cells [10]. HA-crystals, identical to the HA-crystals found in the human joint disease, having a diameter of between 0.1 and 5 µm, induce prostaglandin E₂ release by macrophages [2]. Apatite-containing particles are frequently found in osteoarthritic joints. Hydroxyapatite has been reported to induce acute inflammatory reaction in human skin [18] and synovitis after injection into dog's knee joints [60]. Articular chondrocytes were shown to secrete collagenase, protease, and prostaglandins E₂ and F₂ after phagocytosis of HA-particles. Cheung et al. [16] showed that the phagocytosis of HA-crystal found in osteoarthritic joint induced the multiplication of synovial cells. These results were attributed to the increase of cytoplasmic Ca²⁺ suggested by a degradation of the crystals once phagocytosed.
Degradation of hydroxyapatite-ceramics

Chemical, biochemical and biological evidence of Hydroxyapatite-solubility: Hydroxyapatite is one of the less soluble calcium-phosphates used as bone substitute material. The solubility phase diagram of the most frequently used calcium phosphates as biomaterials shows that although HA is the most stable phase under many conditions, it becomes less stable than dicalcium phosphate dihydrate (DCPD) if the solution is sufficiently acidic. Such ranges of acid pH are commonly found in some cell compartments and particularly in cells responsible for the bone degradation. It is considered that both the endocytic and exocytic vacuolar apparatus in cells are maintained at low pH [3]. Endocytic vesicles, lysosomes, portions of the trans-Golgi apparatus, certain acid-producing secretory vesicles, and the "resorption chamber" of the osteoclast are low pH cell compartments. Using fluorescein-labelled dextran, pH values of 4.7-4.8 were measured in macrophage lysosomes [57]. Similar pH values were measured by Baron et al. [7] in the extracellular resorption region of osteoclasts. A recent report showed that cultured fibroblasts phagocytosed HA-particles inside or in close proximity to their acidic intracellular compartment identified by neutral red uptake [24]. Alliot-Licht et al. [1], Kwong et al. [41], and Orly et al. [53] have demonstrated that calcium phosphate grains having a size less than 50 μm can be phagocytosed by several kinds of cells such as fibroblasts, osteoblasts, established osteoblast-like cell lines, synovial cells, and macrophages then degraded in intracellular vesicles. Subsequent to the internalization of HA-particles, it has been observed that the growth rate and DNA synthesis of osteoblast were significantly inhibited [1]. The HA-particle size was found by Evans [22] to have an effect on the cytotoxicity of HA in vitro. Finely ground powders of HA reduced the growth rate of fibroblast line. The effect was dose-related and only occurred with small particles (3-7 μm). A physical effect could interfere with the cell growth. In particular, the elevated surface at the point of contact of the cell culture medium, when the ceramic powder was finely ground, could induce an ionic modification of the culture medium responsible for the cytotoxicity of the ceramic as demonstrated by Hyakuna et al. [35]. However, these results were not consistent with the work of Mitchell et al. [50] who reported a mitogenic effect of a mixture of calcium phosphate crystals (HA, calcium phosphate, and calcium pyrophosphate dihydrate) on synovial cells. Surface characteristics and solubility of the materials tested could explain this difference. The increase in free cytosolic Ca²⁺ concentration with a simultaneous release of inositol(1,4,5) triphosphate seems to be necessary for a mitogenic effect [49]. This is consistent with the higher mitogenicity of free Ca²⁺ compared with HA crystals as reported by Praeger and Gilchrest [59]. Based on earlier reports [36-38] published on the implantation of calcium phosphate ceramic, HA-ceramics were supposed to be a non- or slowly-degradable material in a biological environment. HA was considered as non-resorbable, and β-TCP was considered as slowly degraded. However, the biological effects of the HA on isolated cells related to the release of Ca²⁺ from the ceramic, shows that HA is not absolutely stable and may be degraded in living tissue. It is particularly interesting to know the factors affecting the degradation of HA-coatings which are supposed, in some cases, to last as long as possible, once implanted. Klein et al. [39] have demonstrated that the solubility of coatings was much higher than sintered particles. Degradation of calcium phosphate coatings in a buffered solution in vitro was shown for tetracalcium phosphate, tricalcium phosphate and HA. Maxian et al. [47] recently reported that a greater in vitro Ca²⁺ dissolution, and a greater surface, physical, and chemical changes were observed for poorly crystallized coatings compared to amorphous coatings, suggesting that amorphous coatings could be more advantageous if coating longevity is desired. This is not consistent with what has been previously published [61]. When we investigated the in vitro behaviour of animal cells grown at the point of contact of HA-coating having different crystallinities, we found that more Ca²⁺ dissolved from coatings having a low crystallinity (40-70%) compared to the coating having a high crystallinity (100%) [25]. HA-particles were found inside cells grown on both poorly and highly crystalline coatings. The highly crystalline coatings induced a cytostatic effect. It is interesting to note that a coating degradation was found in vitro and during short periods of time for all the crystallinities tested.

Microscopic evidence of material degradation: HA is used under different forms: dense, porous, pow­dered or thin coatings. Gomi et al. [29] showed that osteoclast-like cells resorb sintered synthetic hydroxyapatite in vitro and that the nuclearity of the resorptive cells is influenced by the surface topography of the substrate. Koerten et al. [40] demonstrated that spray dried HA-spheres were degraded when injected into the mouse peritoneal cavity. Müller-Mai et al. [51] showed the microscopic degradation of HA-coating. Recently Baslé et al. [8] demonstrated that macroporous calcium phosphate elicited, at the ceramic contact, the recruitment of a double multinucleated cell population, giant multinucleated cells and osteoclasts, having a resorption activity. This result is not consistent with that obtained by Glowacki et al. [28]; it could be due to the difference in features chosen for osteoclast characterization.

We have been able to perform histological analyses of HA materials implanted in human and animal bone in the form of coating or macroporous ceramics in order to evaluate the material degradation in vivo.

Materials and Methods

Fifteen hip prostheses coated with plasma spray­ed HA coatings (characteristics given in Table 1) and implanted for periods from 5 days up to three years (Table 2) were retrieved post-mortem from patients deceased for reasons which were not directly related to the
Table 1. Characteristic of the HA-coatings analyzed.

<table>
<thead>
<tr>
<th>Plasma-sprayed powder</th>
<th>more than 97%</th>
</tr>
</thead>
<tbody>
<tr>
<td>Crystallographic structure</td>
<td>Ca₁₀(PO₄)₆(OH)₂, other compounds admitted: tricalcium phosphate (traces), calcium oxide (traces), amorphous phase</td>
</tr>
<tr>
<td>Crystallinity</td>
<td>60% ± 10%</td>
</tr>
<tr>
<td>Coating thickness</td>
<td>150µm ± 50µm</td>
</tr>
<tr>
<td>Trace elements</td>
<td>As &lt; 3 ppm; Cd &lt; 5 ppm; Hg &lt; 5 ppm; Pb &lt; 3 ppm</td>
</tr>
<tr>
<td>Mechanical strength (pull-out test)</td>
<td>35 MPa</td>
</tr>
</tbody>
</table>

Table 2. Extreme values of the thickness (in µm) of coating implanted for various period (thickness before implantation: 100-150 µm).

<table>
<thead>
<tr>
<th>Implantation</th>
<th>Metaphysis</th>
<th>Middle</th>
<th>Diaphysis</th>
</tr>
</thead>
<tbody>
<tr>
<td>12 months</td>
<td>99-220</td>
<td>50-164</td>
<td>40-184</td>
</tr>
<tr>
<td>24 months</td>
<td>0-183</td>
<td>0-180</td>
<td>0-188</td>
</tr>
<tr>
<td>43 months</td>
<td>58-164</td>
<td>104-204</td>
<td>119-163</td>
</tr>
<tr>
<td>1.5 months</td>
<td>104-143</td>
<td>123-182</td>
<td>164-119</td>
</tr>
<tr>
<td>7 months</td>
<td>163-137</td>
<td>213-155</td>
<td></td>
</tr>
</tbody>
</table>

Table 3. Characteristics of the macroporous HA-ceramics.

<table>
<thead>
<tr>
<th>Characteristic</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Chemical composition</td>
<td>100% HA</td>
</tr>
<tr>
<td>Porosity</td>
<td>70%</td>
</tr>
<tr>
<td>Pore diameter</td>
<td>500-1500 µm</td>
</tr>
<tr>
<td>Grain size</td>
<td>3 µm</td>
</tr>
<tr>
<td>All pores interconnected</td>
<td></td>
</tr>
</tbody>
</table>

hip surgery. The femurs containing the HA-coated prostheses were cut into three parts which were fixed into a 4% formaldehyde solution into PBS (phosphate buffered saline). They were dehydrated in alcohol solutions and embedded into PMMA (polymethyl methacrylate). Two-thick sections were made using a low speed cooled diamond saw. They were re-embedded at the surface of a polystyrene block and polished until a 50-100 µm thickness. Sections observed using a light microscope were stained with a fuscin-toluidine solution after etching the block in a 1% formic acid solution and a 20% methanol solution. Sections observed using scanning electron microscopy (SEM) were stained with a silver methenamine solution [23]. Briefly, sections were stained using a 1% silver methenamine solution (brought to pH 9.6 with NH₂OH) for 4 minutes in a microwave oven at 20% power (2450 MHz). Sections were then examined using the backscattered mode of a SEM operated at 25 kV. The degradation of the coating was measured on sections of 5 HA-coated prostheses using an image analysis system. The maximum and minimum coating thickness was determined on sections made at three levels on each prosthesis.

On thick sections of two HA-coated prostheses implanted three weeks and one year, the prosthesis was detached from the bone and treated with a 2% NaOCl solution for 24 hours to remove the organic matrix adsorbed on the coating. They were dehydrated and gold-palladium coated, then they were observed in the SEM at 25 kV.

Twenty macroporous HA ceramics composed of pure HA (characteristics given in Table 3) were implanted into 9 mm holes drilled into the cortical bone of the medium diaphysis of sheep femurs for periods from one month to ten months. Five other ceramics were implanted for two months into defects made into the medium diaphysis of dog’s ulnae. Once retrieved, the implants were processed as described above for histological analysis.

Results and Discussion

A decrease in the coating thickness was noted after a few months of implantation (Table 2). This degradation can appear after a few weeks of implantation and was not dependent solely on time. The degradation could be completed after a two years implantation period or was non-detectable after three years. SEM observations of specimens, whose organic matter had been removed by NaOCl, showed that the surface of the one year implanted coating was very rough (Fig. 3) compared to the surface of the three weeks implanted one (Fig. 4). Higher magnifications made it possible to see the crystals of the coating suggesting a dissolution of the amorphous phase between the grains. Grains also showed some degradation marks on some of their facets (Fig. 5).

Multinucleated cells and mononuclear cells were found at the coating contact point in all the sections examined except the sections taken from the prosthesis implanted for five days. These cells were located at the coating contact in regions facing medullary cavities. Most of them, after a few months implantation period, had HA-grains included in their cytoplasm showing a phagocytosis of calcium phosphate grains coming from the coating (Fig. 6). Several types of coating degradation could be described morphologically and were observed both by SEM or light microscopy on prostheses coating implanted for a few months (5 months or more). Several of these types could be observed on the same prosthesis: a) coating having a non-altered thickness in contact with multi- or/and mono-nucleated cells containing HA-particles (Fig. 7); b) coating that was of an altered thickness in regions facing medullary cavities although coating in contact with bone tissue is not altered (Fig. 8); c) coating having an homogeneously degraded thickness; and d) coating degraded regions invaded by bone tissue (Fig. 9).
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Figure 3. Scanning electron micrograph of a one year implanted HA-coated prosthesis in human. The organic phase in this sample had been removed by using NaOCl. The coating surface (C) is rough suggesting that degradation occurred during the implantation period. Bone trabeculae (B) are apposed on the coating surface. Bar = 550 μm.

Figure 4. Scanning electron micrograph of a three weeks implanted HA-coated prosthesis. The organic phase in this sample had been removed by using NaOCl. The surface is very smooth showing no degradation sign. Bar = 140 μm.

Figure 5. High magnification scanning electron micrograph of the surface of the prosthesis shown in Figure 3. The amorphous phase is resorbed and the grains constituting the crystalline phase of the coating are observed. Some of these grains show resorption marks on some of their facets (arrowheads). Bar = 10 μm.

Figure 6. Back-scattered electron micrograph of a silver methenamine stained section of a HA-coating resorption zone (HA) of a two years implanted prosthesis. Osteoclast and macrophages contains calcium-phosphate particles (arrowheads). Ti: titanium alloy. Bar = 35 μm.
A similar process of grain phagocytosis was observed on pure HA crystalline ceramics implanted in bone defects in dogs invaded by loose connective tissue. Mononuclear cells were found at the contact point of the ceramic surface. Some of these cells were filled with many calcium phosphate grains located inside cell vacuoles filling the whole cytoplasm (Fig. 10). They were located in close proximity to the ceramic, suggesting that the solubilization occurs rapidly inside the vacuoles. No morphological sign of cell damage was found. Marks of cell resorption were found in the ceramic zones where the cells were present. The same HA-ceramics implanted in an osseous site in sheep for periods ranging from two months up to ten months and invaded by bone tissue seemed both solubilized and cell resorbed (Fig. 11). Solubilization took place in the surface region in contact with body fluid (in the medullary cavity), and cell resorption took place both in the regions at the bone contact point and in the regions previously partially solubilized. Solubilized regions were identified, on histological sections observed using backscattered electron microscopy as a less dense zone located on the material surface surrounding a dense core of ceramic. The decrease of density was attributed to resorption occurring at the grain boundaries. The evidence of cell resorption of these ceramics was found in some regions where the structure of the ceramic was altered at the contact of some multinucleated cells without HA-grains found in their cytoplasm. This suggested that there was a dissolution process before the grain phagocytosis. Although some evidence of cell mediated dissolution exits, it is difficult to evaluate from this study, which part of the material degradation is due to a solution-mediated process and which part is due to a cell-mediated dissolution.

These results were consistent with those published by den Hollander et al. [19] who studied the degradation of 45Ca-labelled macroporous HA-ceramics. HA showed a small decrease in the specific activity of the implant suggesting a degradation of the material once implanted. However, the lack of radioactivity in the adjacent bone normally being an area of increased calcium metabolism, indicated the restricted availability of the liberated 45Ca for neogenesis of bone tissue. The presence of particles of high specific activity was noted in lymph nodes.

Thus, even HA-ceramics which do not contain an amorphous phase are cell degraded both in vivo and in vitro. HA-macroporous ceramics used in our observations had a very low microporosity (< 2%) with very small pores and could be considered as dense material at the cell level. The molecular mechanism of HA-degradation once implanted can involve surface processes controlling the dissolution rate of HA-crystal and cell excretion of ions, enzymes and proteins modifying the crystal environment. Although the molecular mechanisms ruling the mononuclear precursors of osteoclasts are not known, they are influenced by some of the HA-ceramic characteristic and could be indirectly involved in the HA degradation process.
Conclusions

Dissolution and cell degradation of synthetic HA-ceramic occur simultaneously and it is very difficult to distinguish between them in vivo. We could not establish a rate of degradation of these ceramics since their degradation is a very heterogeneous phenomenon. On HA-coatings, the rate of degradation could vary on the same prosthesis depending on the location where the section was made. Also, depending on the tissue (e.g., bone or bone marrow) in contact with the ceramic coating, the thickness of the coating facing these tissues is not the same. Moreover, the resorption varies from one patient to another.

References

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phages release a peptide stimulator of osteoblast growth.  


**Editor’s Note:** All of the reviewer’s concerns were appropriately addressed by text changes, hence there is no Discussion with Reviewers.