Biocompatibility and osteoconductivity of the pyrost bone substitutes

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Summary. The purpose of this study was to re-evaluate the bone regeneration power and the in vitro biocompatibility of the Pyrost bone substitute. Twenty-four adult New Zealand White rabbits were used. Bone defects were filled over both iliac crest and mid-diaphyseal portion of the ulna bone were created. Appropriate sized-block of Pyrost bone substitute were implanted. Four of the animals were killed at each postoperative month to evaluate its bone regeneration power by histologic study. The Pyrost bones were co-cultured with osteoblasts to evaluate its biocompatibility. The results showed that Pyrost bone substitute was quite stable and incorporated well with active bone regeneration. The Pyrost heal better at the iliac crest than at the ulnar defect. The Pyrost was compatible to the osteoblasts. Osteoblasts had successfully seeded and mitotically expanded on the porous surface of the Pyrost bone graft. The result showed that Pyrost bone obviously exerts an intense stimulus on osteo-regeneration in the presence of osteoblasts. We consider Pyrost to be an alternate to the conventional preserved allografts that is occasionally necessary.

Key words: Pyrost bone substitute, Biocompatibility, Osteoconductivity

Introduction

Congenital or acquired bone defect is a major problem in the orthopedic surgery. In order to accelerate the healing of bone defect, bone grafting is required in about 15% of the reconstructive operations on the locomotor system (Urist et al., 1994). Autografts are preferably used because of their superior efficacy and capability of avoiding transmission of infection. However, it does have drawbacks including donor site morbidity and limited availability, particularly in children (Glowacki et al., 1981; Dutting et al., 1988; Grob, 1988). The effectiveness of allogeneous bone graft is limited by the problems of high cost of bone banking, potential of graft-related disease transfer, the high rates of nonunion and infection, and allograft fractures (Kotani et al., 1992).

A variety of bone grafts substitute materials are commercially available: both mineralized and demineralized forms (Doherty et al., 1994). Pure mineral bone, Pyrost, had been reported as a promising bone-replacement material (Katthgen and Mittelmeier, 1983). Osteoblasts have recently been grown in vitro on the bone-replacement materials (Guillemin et al., 1987; Matsuda and Daves, 1987; Gregoire et al., 1990; Sautier et al., 1990). In the study of Katthgen and Mittelmeier (1983), the experimental model used was mainly the condylar portion of distal femur. It is not a common location for harvesting bone graft. Filling with plenty of osteogenic cells, the iliac crest is one of the most commonly used donor site for harvesting autogenous bone graft. The significant difference between autologous bone and these bone-replacement materials is that autologous bone contained a variety of living cells, including osteoblasts. When being used to replace bony defects, the biocompatibility of bone substitutes is mandatory for achieving bone regeneration.

The purpose of this study were to re-examine the bone regeneration power of Pyrost bone substitute at the site of cancellous and cortical bone defects; also to determine the in vitro biocompatibility of the Pyrost bone substitute. Our hypothesis is that the Pyrost is a suitable carrier system for osteoblasts proliferation and also a potential bone substitute for the iliac bone defect created after harvesting bone graft.

Materials and methods

Animal experiment

Twenty-four adult New Zealand White rabbits, weighing 2.0 to 2.5 kg., were used in this experiment. The animals were fed Purina Laboratory Chow ad
They were then transferred to 5% containing solutions containing NaCl, KCl, NaH₂PO₄, KCI, 3mM Na₂CO₃, and 137 mM NaCl, and sequentially rehydrated down to distilled water. The specimens were fixed with 3% phosphate-buffered formaline for subsequent histological examination. The specimens were stained with Hematoxyline-Eosin and then observed with light microscopy. The control samples were the same as the experimental samples.

For light microscopy examination, sections were prepared using a diamond saw with water cooling (Buehler lsometTM, Lake Bluff, Illinois, USA), and were then coated with Entellan and examined by fluorescence microscopy (Nikon-Microphot-Fx, Japan).

Sequential digestion of newborn Wistar-rat calvaria was performed by using a modification of the methods described by Wong and Cohn (Boonekamp et al., 1984). Briefly, after pretreatment of the dissected calvaria with 4mM Na₀ EDTA in a pre-warmed (37 °C) solution containing 137mM NaCl, 2.7mM KCl, 3mM NaH₂PO₄, pH 7.2 (solution A), for 10 minutes three times, the fragments were sequentially digested with collagenase (180 U/ml, Sigma Co.) in solution A with EDTA. The sequential digestion was performed for 5 minutes followed by two times treatments for 10 minutes. The cells released after each treatment were immediately harvested by centrifugation and re-suspended in culture medium. The osteoblastic phenotype of the cells was assessed by the tests of alkaline phosphatase assay (Doty and Schofield, 1976; Begley et al., 1993).

Three Pyrost bones (10x10x0.5 mm) were placed in sterile dishes. Confluent rat osteoblast cultures were passaged by trypsin-EDTA and were seeded into each well on top of the implant materials at a seeding density of 3x10⁵ cells/cm². A large area of each well was left uncovered to allow assessment of cell plating efficiency by microscopy. The culture media used was Dulbecco’s modified Eagle’s medium supplemented with 10% fetal calf serum (Gibco, UK), penicillin (100 units/ml) and streptomycin (100 mg/ml). The dishes were incubated at 37 °C in an atmosphere supplemented with 5% CO₂ and 95% air and fed with complete changes of medium twice a week. The days of plating was considered as the first day of culture. The test samples were removed from wells at days 3, 5, 7, 9, fixed in 3% formaldehyde in 0.1M PBS buffer (pH 7.4). Each test composed of triplicate samples and separate control at each of the time points. The samples were stained with Hematoxyline-Eosin and then observed and photographed with inverted microscopy.

For electron microscopy examination, the Pyrost bone substitute was fixed with 3% phosphate-buffered formaline solution, dehydrated through a graded ethanol series and critical dried from CO₂ (Ladd Research Industries Inc., Burlington, VT, USA). Specimens were sputter-coated with gold (Model IB-2 Ion Coater, Eiko Engineering, K.K., Japan) and examined in a ISABT (model SX-30 E, International Scientific Instruments, Inc., CA, USA) scanning electron microscope at an accelerating voltage of 15 kV.

Results

In vivo gross morphological and histological evaluation

Pyrost bone grafts were stably fixed into rabbits’ pelvis. After implantation of Pyrost for six months, the location of Pyrost still can be identified, but had been closely incorporated into the host bone. It was further confirmed by the roentgenographic assessment that did not show any radiolucent space surrounding the Pyrost bone substitute. The sham-operated site of pelvic wing showed a bony defect filled with fibrous tissue. The defect was quite obvious in the roentgenogram.

At the ulnar defects grafting with the Pyrost bone substitute, visible callus formation was observed at the end of the first month. The amount of callus formation was much increased at the end of the 6th month. At the control limb without Pyrost bone substitute, callus...
Fig. 1. Light micrograph of bone regeneration on Pyrost at the iliac defect. The darkly colored Pyrost implant (P) is surrounded everywhere, by regenerated bone (R) and osteoid tissue (OT). Von Kossa stain. Bar: 200 μm.

Fig. 2. Fluorescence micrograph of bone regeneration on Pyrost. Vigorous bone regeneration on the surface of the Pyrost implant (P). There is no connective tissue layer between implant and bone, adjacent to it is inconspicuous regenerated marrow tissue. Von Kossa stain. Bar: 200 μm.

Fig. 3. Vigorous bone regeneration on the surface of the Pyrost implant (P). There is minimal connective tissue layer between implant and bone. Note that even after 3 months' implantation, the darkly colored Pyrost is still almost totally preserved. The entire bone-defect area is filled with regenerated bone. Pyrost and regenerated bone are closely linked together. Von Kossa stain. Bar: 80 μm.

Fig. 4. At high power magnification, Pyrost (P) surrounded by regenerated bone (R). Intensive bone regeneration with massive osteoblasts (OB) can be seen. Von Kossa stain. Bar: 20 μm.

Fig. 5. Histological examination of the Pyrost bone grafted at the ulnar defect. At six months after implantation, there is little callus formation (C) of the adjacent cortical bone into the Pyrost (P) observed with intervening fibrous tissue (F). Von Kossa stain. Bar: 200 μm.
formation at both ends of the defect toward the center was observed. However, complete bridging of the defect did not occur. Neither the ulnar defects with Pyrost block nor that without Pyrost block was completely united with autogenous bone formation.

Pyrost bone substitute, incorporated well with the pelvis, showed active bone regeneration. Histological examination revealed that the Pyrost was infiltrated with regenerated bone trabeculae (Fig. 1). The entire bone-defect area is filled with regenerated bone. There is vigorous bone regeneration on the surface of the Pyrost with little connective tissue layer (Fig. 2). After deplastication, vigorous bone regeneration on the surface of the Pyrost implant with massive osteoid tissue can be seen. Pyrost and regenerated bone are closely linked together. Also there is inconspicuous regenerated marrow tissue adjacent to this regenerated bone (Fig. 3). At higher magnification, early phases of bone regeneration with many osteoblasts at the surface of the Pyrost implant were visible (Fig. 4).

In the ulnar defects grafted with Pyrost block, solid incorporation of the Pyrost block into the adjacent cortical bone was not observed. Even after six months' implantation, there is little callus formation of the adjacent cortical bone into the Pyrost observed with intervening fibrous tissue (Fig. 5). This suggested that the bone regeneration and ingrowth was incomplete at the end of the six month.

In vitro biocompatibility of the Pyrost bone graft

At the 3rd day of culture, the Pyrost block exhibited a sparse population of osteoblasts (Fig. 6). The number of cells had increased by the 5th day, the cells began to contact with each other. On the more porous surfaces, the osteoblasts were observed to suspended across the pores by the Pyrost block (Fig. 7). By the 9th day, a nearly complete layer of osteoblasts coated over porous surface (Fig. 8).

The cells, growing on the Pyrost block, typically exhibited numerous pseudopodial and filopodial extensions anchoring the cells to the surface (Fig. 9). Later, layer of flattened polygonal cells developed and covered almost the whole surface of the Pyrost bone substitute (Fig. 10). The osteoblasts successfully seeded and mitotically expanded on the porous surface of Pyrost bone substitute block. The Pyrost bone graft exhibited good biocompatibility in the osteoblast culture.

Discussion

The basic functions of a bone graft or bone substitute are (1) the osteogenic performance, the induction or conduction of bone regeneration, (2) the bony fixation and stabilization and (3) the filling of the bone defect itself (Morscher, 1982). In many cases there are problems with the filling of fairly large bone cavities, so soughing of the alternatives to conventional bone grafts are indicated. In the past decade, many synthetic biomaterials have been reported for bone substitutes, including bio-inert ceramic and bio-active ceramics (Kotani et al., 1992), synthetic calcium phosphates (Kitsugi et al., 1988), plastic and metal implants (Kathagen, 1986). Because in case of requiring bone graft or bone substitute, the ideal approach is not long-term or permanent bone replacement, but the stimulation of bone regeneration followed by filling the defect with the body's own bone. This principle still cannot be adequately realized with above-mentioned synthetic biomaterials. In this study, we focused on the osteoconduction induced by Pyrost bone at the region of iliac crest for its possible role in bone substitute after harvesting autogenous iliac bone graft.

In our study, Pyrost bone graft incorporated well with the bone actively regenerated at the defect of pelvic osteotomy site. The implanted material was resolved slowly, and was still present almost in its entirety after 6 months. Histological examination revealed vigorous bone regeneration on the surface of the Pyrost implant (Fig. 1). The regenerated bone contains little connective tissue layer between implant and bone (Fig. 2) with massive osteoid tissue (Fig. 3) and many osteoblasts at the surface of the Pyrost implant (Fig. 4). The resulting network of regenerated bone surrounding the implant and filling the interstices was as dense as, and sometimes even denser than the surrounding intact bone structures. Since the bone defect chosen in the rabbit cannot be filled spontaneously with regenerated bone, the bone regeneration observed in the experimental groups were mainly under the influence of the Pyrost substitutes (Karaca et al., 1994; Matsuda et al., 1995). In those of ulnar defects, there is little callus formation of the adjacent cortical bone into the Pyrost even after six months' implantation. This suggested that ingrowth was incomplete even after six months. The Pyrost heal better at the iliac crest than at the ulnar defect. In the clinical medicine, the usual site for harvesting autogenous bone graft was the iliac crest. The loss of bone in the donor site and the related cosmetic problem may bothering in some degree. The previous harvested site was also impossible for further harvesting graft. The present results provide the evidence that Pyrost bone substitute is good osteoconductive at the iliac crest. We suggest that in appropriate indications, implantation of Pyrost substitute can be a very useful and effective bone substitute in the donor site after harvesting bone graft. This makes this site possible to reuse in later situation.

An important prerequisite for a bone graft substitute material is that the material must be able to support osteoblast attachment and even able to promote proliferation of the attached cells. Doherty suggested that cell attachment and proliferation is faster and dimineralized bone as opposed to mineralized form (Doherty et al., 1994). In the present study, we elucidated that the biocompatibility of the porous surface of mineralized Pyrost bone substitute to osteoblasts was rather good (Figs. 5-8). The osteoblasts had successfully seeded onto and mitotically expanded on the Pyrost bone
Fig. 6. In vitro biocompatibility of the Pyrost bone graft with osteoblast. After 3 days in culture, the Pyrost bone samples exhibited only a sparse population of osteoblasts (OB). H&E stain. Bar: 20 μm.

Fig. 7. In vitro biocompatibility of the Pyrost bone graft with osteoblast. At the 5th day, the cells exhibiting contact with each other. On the more porous surfaces, osteoblasts (OB) were observed to suspended across the pores by their processes. H&E stain. Bar: 20 μm.

Fig. 8. Stereomicroscopic examination for biocompatibility of the Pyrost bone graft with osteoblast. At the 9th day, an almost complete layer of osteoblasts coated the porous surface of the Pyrost bone. H&E stain. Bar: 200 μm.

Fig. 9. Scanning electron microscopic examination of the in vitro biocompatibility of the Pyrost bone substitutes with the osteoblasts. At the end of 1 day's culture, the cells that grew on the materials exhibited many pseudopodial and filopodial extensions anchoring the cells to the surface of the Pyrost. Bar: 5 μm.

Fig. 10. Scanning electron microscopic examination of the in vitro biocompatibility of the Pyrost bone substitutes with the osteoblasts. At the end of 7 day's culture, layer of flattened polygonal cells was present.
substitute (Figs. 9, 10). The attachment of osteoblasts on the Pyrost materials exhibited numerous cellular processes (Figs. 7, 9) suggests that the cells tend to attach regardless the topography of the materials. It suggested that porosity may be important to allow fluid circulation, bone ingrowth and mechanical stability at the implantation site (Brunette, 1988). Such observations are in accord to those of Begley et al. (1993), and suggested that migratory osteoblasts adhere to any firm substrate presented in vitro (Jones and Boyd, 1979). The Pyrost demonstrated a high biocompatibility to osteoblasts, i.e., it does not only allow the attachment of the osteoblasts but also the proliferation of the osteoblasts (Matsuda et al., 1995). The Pyrost bone substitute is possible to behave as a delivery system for bone growth. However, the mechanism whereby Pyrost implant affects healing remained to be solved.

Conclusion

In this study, Pyrost was used as a substitute after harvesting autogenous iliac bone graft. The result showed that Pyrost bone obviously exerts an intense stimulus on osteo-regeneration and in-vitro study also showed good biocompatibility to the osteoblasts. We consider Pyrost to be an alternate to the conventional preserved allografts that is occasionally necessary.

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References


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