

Basic Study

Utility of tricalcium phosphate and osteogenic matrix cell sheet constructs for bone defect reconstruction

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Supported by Grant-in-Aid for Young Scientists (KAKENHI).

Ethics approval: The animal protocol was designed to minimize pain or discomfort to the animals. The animals were acclimatized to laboratory conditions (23 °C, 12h/12h light/dark, *ad libitum* access to food and water).

Institutional animal care and use committee: All procedures involving animals were reviewed and approved by the Institutional Animal Care and Use Committee of the Nara Medical University.

Conflict-of-interest: The authors state that they have no conflicts of interest to disclose.

Data sharing: No additional data are available.

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Received: September 30, 2014

Peer-review started: October 1, 2014

First decision: October 28, 2014

Revised: January 27, 2015

Accepted: March 18, 2015

Article in press: March 20, 2015

Published online: June 26, 2015

Abstract

AIM: To determine the effects of transplanting osteogenic matrix cell sheets and beta-tricalcium phosphate (TCP) constructs on bone formation in bone defects.

METHODS: Osteogenic matrix cell sheets were prepared from bone marrow stromal cells (BMSCs), and a porous TCP ceramic was used as a scaffold. Three experimental groups were prepared, comprised of TCP scaffolds (1) seeded with BMSCs; (2) wrapped with osteogenic matrix cell sheets; or (3) both. Constructs were implanted into a femoral defect model in rats and bone growth was evaluated by radiography, histology, biochemistry, and mechanical testing after 8 wk.

RESULTS: In bone defects, constructs implanted with cell sheets showed callus formation with segmental

or continuous bone formation at 8 wk, in contrast to TCP seeded with BMSCs, which resulted in bone non-union. Wrapping TCP constructs with osteogenic matrix cell sheets increased their osteogenic potential and resulting bone formation, compared with conventional bone tissue engineering TCP scaffolds seeded with BMSCs. The compressive stiffness (mean \pm SD) values were 225.0 ± 95.7 , 30.0 ± 11.5 , and 26.3 ± 10.6 MPa for BMSC/TCP/Sheet constructs with continuous bone formation, BMSC/TCP/Sheet constructs with segmental bone formation, and BMSC/TCP constructs, respectively. The compressive stiffness of BMSC/TCP/Sheet constructs with continuous bone formation was significantly higher than those with segmental bone formation and BMSC/TCP constructs.

CONCLUSION: This technique is an improvement over current methods, such as TCP substitution, and is useful for hard tissue reconstruction and inducing earlier bone union in defects.

Key words: Bone marrow stromal cells; Osteogenesis; Bone regeneration; Tissue engineering; Calcium phosphate

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Core tip: The treatment of bone defects is a common clinical problem for orthopedic surgeons. Artificial bone combined with cultured bone marrow stromal cells (BMSCs) has been examined and applied for the treatment of clinical cases such as small bone defects. In the present study, we developed constructs of BMSCs and beta-tricalcium phosphate (TCP) combined with osteogenic matrix cell sheets (BMSC/TCP/Sheet constructs), which showed a vigorous osteogenic potential compared with conventionally engineered bone tissue structures consisting of BMSC/TCP constructs. The BMSC/TCP/Sheet constructs will be useful for reconstruction of hard tissues, such as bone defects, as new tissue-engineered bone.

Ueha T, Akahane M, Shimizu T, Uchihara Y, Morita Y, Nitta N, Kido A, Inagaki Y, Kawate K, Tanaka Y. Utility of tricalcium phosphate and osteogenic matrix cell sheet constructs for bone defect reconstruction. *World J Stem Cells* 2015; 7(5): 873-882 Available from: URL: <http://www.wjgnet.com/1948-0210/full/v7/i5/873.htm> DOI: <http://dx.doi.org/10.4252/wjsc.v7.i5.873>

INTRODUCTION

The treatment of bone defects is a common clinical problem for orthopedic surgeons. Autologous bone grafts including vascularized bone grafts are performed for treatment of osteonecrosis and bone defects in clinical practice^[1,2]. However, it is necessary to sacrifice intact bone such as the pelvis or fibula. Moreover, there is limited bone available for autologous transplantation.

Artificial bone such as hydroxyapatite (HA) and beta-tricalcium phosphate (TCP) can be applied as a scaffold for the treatment of small bone defects. Although these biomaterials provide a physical matrix for deposition of new bone and may guide the extension of bone, they do not induce ectopic bone formation^[3]. To achieve osteogenesis in artificial bone, constructs of artificial bone combined with cultured bone marrow stromal cells (BMSCs), which are designated as engineered bone tissue structures^[4-6], have been examined and applied for the treatment of clinical cases such as avascular necrosis of the femoral head and benign tumors^[7-9]. However, some cases require a long period of time to replace autologous bone, which subsequently delays bone union.

Previously, we developed a mechanical retrieval method to fabricate cell sheets of cultured BMSCs with osteogenic potential, designated osteogenic matrix cell sheets, that could be transplanted without a scaffold, and resulted in bone formation at subcutaneous sites^[9-12]. The utility of transplantation of engineered bone tissue structures for hard tissue reconstruction will increase if such treatment can achieve sufficient bone formation for early bone union after transplantation. In our previous study, we reported that a construct of osteogenic matrix cell sheets and artificial bone results in extensive bone formation compared with a construct containing a BMSC suspension. A construct with osteogenic matrix cell sheets allows bone formation inside of the pores as well as outside of the artificial bone. Outside bone formation can unite two artificial bones by bridging bone formation^[11]. These results suggested that transplanting a combination of osteogenic matrix cell sheets and engineered bone tissue structures may advance bone formation for reconstruction of bone defects. Marcacci *et al*^[9] reported the clinical outcomes of long bone repair using macroporous bioceramics with BMSCs. However, they used a BMSC suspension to prepare BMSC/bioceramic composites. Therefore, in the present study, we prepared TCP constructs with or without osteogenic matrix cell sheets and assessed their osteogenic potential after transplantation subcutaneously and at bone defect sites. We compared the bone formation patterns between TCP constructs with or without osteogenic matrix cell sheets.

MATERIALS AND METHODS

Preparation of bone marrow cells

The experimental procedures were approved by the Animal Experimental Review Board of Nara Medical University. Male Fisher 344 (F344) rats were purchased from Japan SLC (Shizuoka, Japan) for use as donors and recipients. The method for bone marrow cell preparation has been reported previously^[12-14]. Briefly, bone marrow cells were obtained from the femoral shafts of 7-wk-old F344 rats by flushing with 10 mL of

culture medium. The cells were collected in two T-75 flasks (Costar, Cambridge, MA, United States) containing 15 mL of minimal essential medium (MEM; Nacalai Tesque, Kyoto, Japan) supplemented with 15% fetal bovine serum (FBS; Gibco Life Technologies, Carlsbad, CA, United States) and antibiotics (100 U/mL penicillin and 100 µg/mL streptomycin; Nacalai Tesque), and incubated at 37 °C in a humidified atmosphere with 5% CO₂. At confluency (approximately day 14), the primary cultured cells were harvested using a 0.25% trypsin/EDTA solution (Nacalai Tesque) to obtain in a cell suspension.

Cell sheet preparation

To prepare an osteogenic matrix cell sheet, a cell suspension was seeded at a density of 1×10^4 cells/cm² in a 100-mm cell culture dish (100 × 20 mm; BD Falcon, NJ, United States) and cultured with medium containing 10 nmol/L dexamethasone (Dex; Sigma, MO, United States) and 82 µg/mL ascorbic acid phosphate (AscP; L-ascorbic acid phosphate magnesium salt n-hydrate; Wako Pure Chemical Industries, Kyoto, Japan) until confluency (approximately 2 wk)^[10,13]. The cells were then rinsed twice with phosphate-buffered saline (PBS; Gibco), and the sheet was lifted by a mechanical retrieval method using a scraper. Because the osteogenic matrix cell sheet was sufficiently robust to handle with a scraper and forceps, handling procedures for the cell sheet were performed easily, such as combination of the cell sheet with TCP. Although β-glycerophosphate (β-GP) was added to create osteogenic matrix cell sheets in our initial study^[10], recent investigations have revealed that osteogenic matrix cell sheets with high osteogenic potential can be prepared without β-GP^[15,16]. Therefore, the osteogenic matrix cell sheets in the present study were created without β-GP.

Construction of TCP containing BMSCs

Sterilized porous TCP ceramics (Osferion; disks: 5 mm diameter and 2 mm thick; cylinders: 5 mm outer diameter, 1 mm inner diameter, 5 mm in length, and 60% porosity) were provided by Olympus Terumo Biomaterials (Tokyo, Japan). After immersion in MEM, the air bubbles in the TCP were aspirated and each ceramic was soaked in a cell suspension (1×10^6 cells/mL) for 2 h in a CO₂ incubator at 37 °C^[13]. The constructs were then cultured with osteoinductive supplements including Dex, AscP, and 10 mmol/L β-GP (Sigma) for 2 wk in a CO₂ incubator at 37 °C to obtain BMSC/TCP constructs.

Experimental groups

We prepared three experimental groups according to the type of implant received. The C group received BMSC/TCP constructs, the S group received TCP combined with an osteogenic matrix cell sheet (Sheet/TCP constructs), and the SC group received BMSC/

TCP constructs combined with an osteogenic matrix cell sheet (BMSC/TCP/Sheet constructs). Constructs with an osteogenic matrix cell sheet were prepared by simply wrapping the sheet around the constructs just prior to transplantation. The TCP ceramics used for subcutaneous transplantation and bone defect model experiments were the disk and cylinder types, respectively.

Subcutaneous implantation of constructs

Constructs with a TCP disk were implanted subcutaneously onto the back of syngeneic 7-wk-old male rats using a previously described procedure^[10,17]. Each group included six disks for one recipient rat as the subcutaneous implantation model. After 4 wk, the implanted constructs were harvested and evaluated radiographically, histologically, and biochemically. For radiographic and histological evaluations, the harvested disks were fixed in buffered formalin (Wako Pure Chemical Industries). After obtaining X-ray images, each disk was decalcified with a KCl solution (K-CX solution; Falma, Tokyo, Japan), embedded in paraffin, cut at the middle of the disk parallel to the base, and then stained with hematoxylin and eosin (HE) for histological evaluation.

Biochemical analysis

The alkaline phosphatase (ALP) activities and osteocalcin contents of harvested constructs were measured as described previously^[17,18]. Briefly, each disk was crushed and homogenized using a microhomogenizer in 1 mL of 0.2% Nonidet P-40 (NP-40), and then centrifuged at 12000 rpm for 10 min at 4 °C. A 10-µL aliquot of the supernatant was combined with 56 mmol/L 2-amino-2-methylpropanediol buffer containing 10 mmol/L p-nitrophenylphosphate and 1 mmol/L MgCl₂. After incubation for 30 min at 37 °C, 2 mL of 0.2 N NaOH was added to stop the reaction and the ALP activity was calculated by measuring the absorbance of p-nitrophenol released at 410 nm using a spectrophotometer.

After extraction with 0.2% NP-40 in 4 mL of 20% formic acid for 2 wk at 4 °C, osteocalcin was extracted from the sediment. A 0.5-mL aliquot of the formic acid extract was applied to a prepacked Sephadex G-25 column (NAP-5 column; Amersham Pharmacia Biotech AB, Uppsala, Sweden) and eluted with 10% formic acid. The protein fractions were pooled and dried. After dissolution in 0.5 mL of ELISA sample buffer, the osteocalcin content was assayed using a Rat Osteocalcin ELISA Kit (DS Pharma Biomedical, Osaka, Japan)^[18].

Implantation of constructs into the bone defect model

Bone defects of the femur were created in F344 rats (300-350 g body weight) under anesthesia. Briefly, a lateral incision was made on the hind limb and the vastus muscle was divided longitudinally to expose the femur. The femoral shaft was removed from

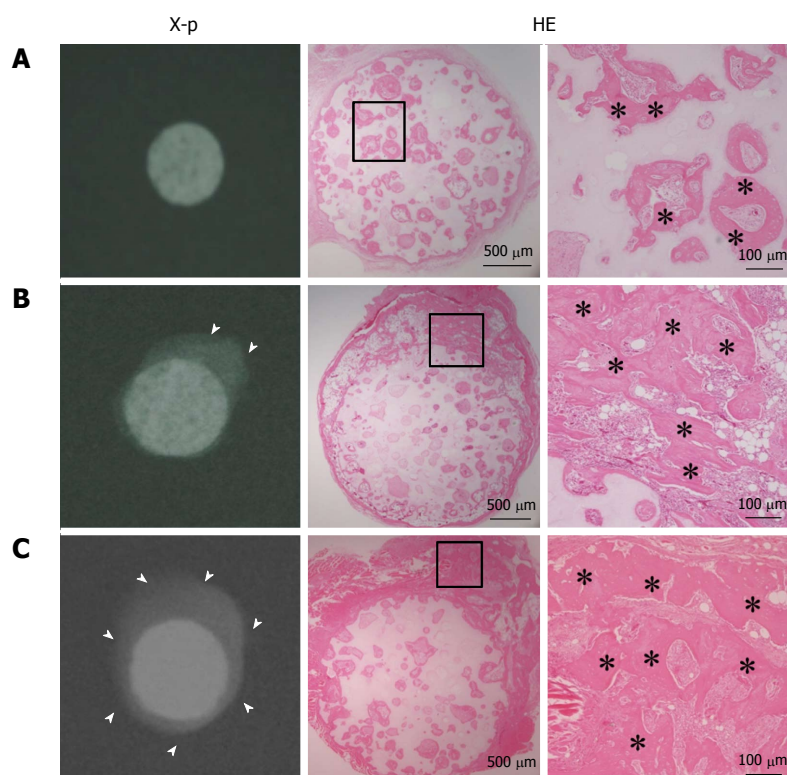


Figure 1 X-ray images and hematoxylin and eosin-stained sections at 4 wk post-implantation. A: BMSC/TCP constructs (C group); B: Sheet/TCP constructs (S group); C: BMSC/TCP/Sheet constructs (SC group). The S and SC groups showed abundant calcification around the TCP disk. Arrowheads indicate the area of calcification. Bone formation was observed histologically in the pores of the TCP disk in all groups. The S and SC groups showed bone formation on the surface of the disk. Higher magnification images of the boxed areas are shown at the right side of each panel. Asterisks indicate bone tissue. HE: Hematoxylin and eosin.

the distal lesser trochanter to a length of 10 mm to create a bone defect that was filled with a TCP-cylinder construct. The cylinder was fixed with an 18 G needle inserted into the intramedullary femoral shaft to achieve ridged fixation. A BMSC/TCP cylinder (BMSC/TCP construct) was transplanted into the bone defect of the left femur, while a BMSC/TCP cylinder with an osteogenic matrix cell sheet (BMSC/TCP/Sheet construct) was transplanted into the defect of the right femur. To evaluate bridging bone formation, X-ray images were taken under anesthesia at 2, 4, and 8 wk postoperatively.

Evaluation of bone formation in the bone defect model

At 8 wk post-transplantation, the rats were euthanized and both hind limbs were harvested. After removal of the intramedullary pins, micro-CT images were taken using a Microfocus X-ray CT system (SMX-160CTS; Shimadzu, Kyoto, Japan). Five of the harvested femurs were then dissected from the surrounding muscle, fixed in 10% formalin, decalcified with K-CX solution, and embedded in paraffin. For histology, the femurs were cut longitudinally and stained with HE. Eight of the harvested femurs were applied to compression testing using a universal testing machine (5566; Instron, Canton, MA, United States) equipped with a computer for data acquisition. The harvested femur was compressed longitudinally and the primary

yield point for fracture of the TCP was calculated. The crosshead speed was 10 mm/min. Compressive stiffness was calculated by dividing the compressive stress by the compressive strain in the linear portion to the primary yield point of the stress-strain curve.

Statistical analysis

The values for ALP activity, osteocalcin content, and compressive stiffness were calculated as the mean and SD. Statistical significance was determined by one-way ANOVA with post-hoc multiple comparisons using Tukey’s test. A value of $P < 0.05$ was considered statistically significant.

RESULTS

Subcutaneous implantation

Figure 1 shows the radiographic images and histology (HE staining) at 4 wk after subcutaneous implantation. Abundant calcification around the disks was observed in the radiographic images of the S and SC groups, whereas no calcification was observed in the C group. Histology showed bone formation in all harvested disks. Although the C group showed bone formation only in the pores of the ceramics, the S and SC groups showed bone formation in the pores and around the disks. Less bone formation appeared to be in the central area of the disks in the S group compared with

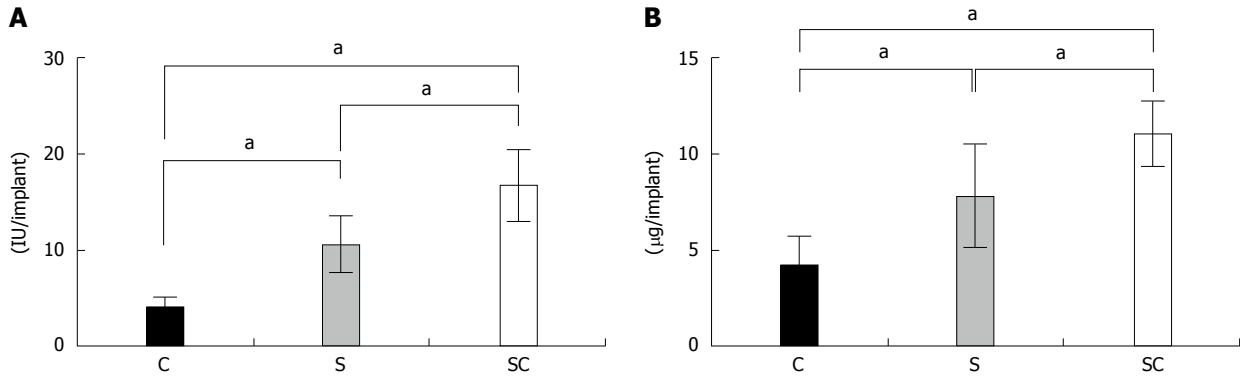


Figure 2 ALP activity (A) and osteocalcin content (B) of harvested constructs. The C, S, and SC groups indicate BMSC/TCP (black column), Sheet/TCP (gray column), and BMSC/TCP/Sheet (white column) constructs, respectively. Values are means \pm SD ($n = 6$). ^a $P < 0.05$. BMSCs: Bone marrow stromal cells; TCP: Tricalcium phosphate.

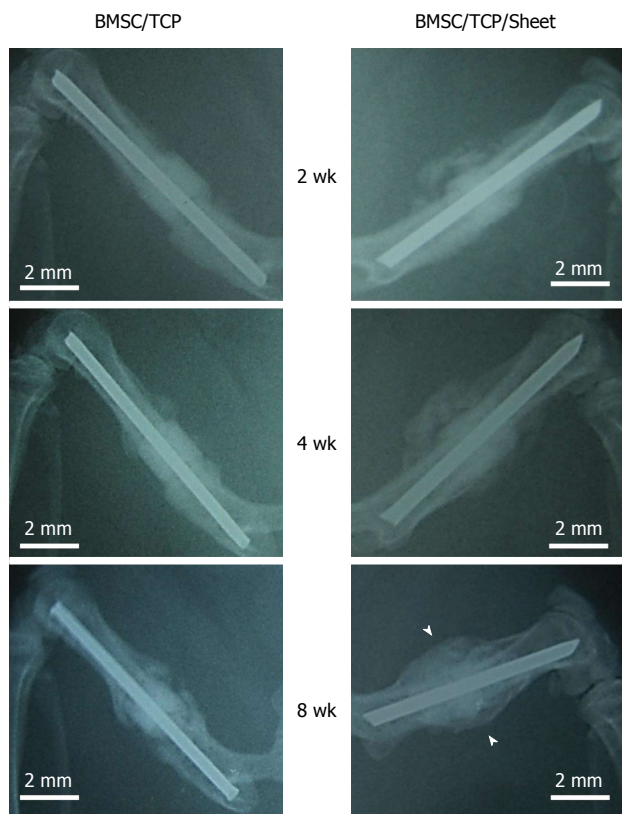


Figure 3 X-ray images of constructs implanted into femurs. The femoral shaft was removed and replaced with BMSC/TCP constructs with or without cell sheets. Callus formation appeared around the BMSC/TCP constructs with cell sheets (BMSC/TCP/Sheet) at 2 wk post-implantation, was extended at 4 wk, and continued to the host bone at 8 wk. In contrast, the BMSC/TCP constructs showed non-union at 8 wk. Arrowheads indicate continuous bone formation. Bar = 2 mm. BMSCs: Bone marrow stromal cells; TCP: Tricalcium phosphate.

the SC group.

The ALP activities (Figure 2A) and osteocalcin contents (Figure 2B) in the SC group were significantly higher than those in the other groups at 4 wk post-implantation. In the S group, these values were significantly higher than those in the C group.

Bone defect model

Radiographic images were taken at 2, 4, and 8 wk

post-implantation (Figure 3). At 2 wk, obvious callus formation was observed around the implanted constructs with osteogenic matrix cell sheets. At 8 wk post-implantation, callus formation had developed and bridging callus formation between host bones resulted in bone union. In contrast, no bridging callus formation was observed around constructs without osteogenic matrix cell sheets at 8 wk post-implantation.

Figure 4 shows the micro-CT images and histology of each construct implanted into femurs at 8 wk. Implanted BMSC/TCP/Sheet constructs showed bone formation around the TCP cylinder ($n = 8$). However, micro-CT revealed two patterns of bone formation, continuous bone formation to host bones (Figure 4A, $n = 4$) and segmental bone formation over the cylinder (Figure 4B, $n = 4$). The ratio of segmental and continuous bone formation among the samples was 1:1. We defined BMSC/TCP/Sheet constructs showing bridging bone formation between host bones across the TCP by micro-CT as continuous bone formation. Bone formation observed around the constructs, but not continuing to the host bones, was defined as segmental bone formation. Bone formation was not observed around implanted BMSC/TCP constructs ($n = 8$) in which soft tissue interposition between the cylinder and the host bone resulted in non-union (Figure 4C). Micro-CT also revealed that BMSC/TCP/Sheet constructs with either continuous or segmental bone formation were intact, whereas broken TCP cylinders were found in BMSC/TCP constructs (Figure 4C).

Figure 5 shows the results of biomechanical compression testing. The compressive stiffness (mean \pm SD) values were 225.0 ± 95.7 , 30.0 ± 11.5 , and 26.3 ± 10.6 MPa for BMSC/TCP/Sheet constructs with continuous bone formation, BMSC/TCP/Sheet constructs with segmental bone formation, and BMSC/TCP constructs, respectively. The compressive stiffness of BMSC/TCP/Sheet constructs with continuous bone formation was significantly higher than those with segmental bone formation. The stiffness of constructs with segmental bone formation was similar to that of constructs without osteogenic matrix cell sheets (BMSC/TCP constructs).

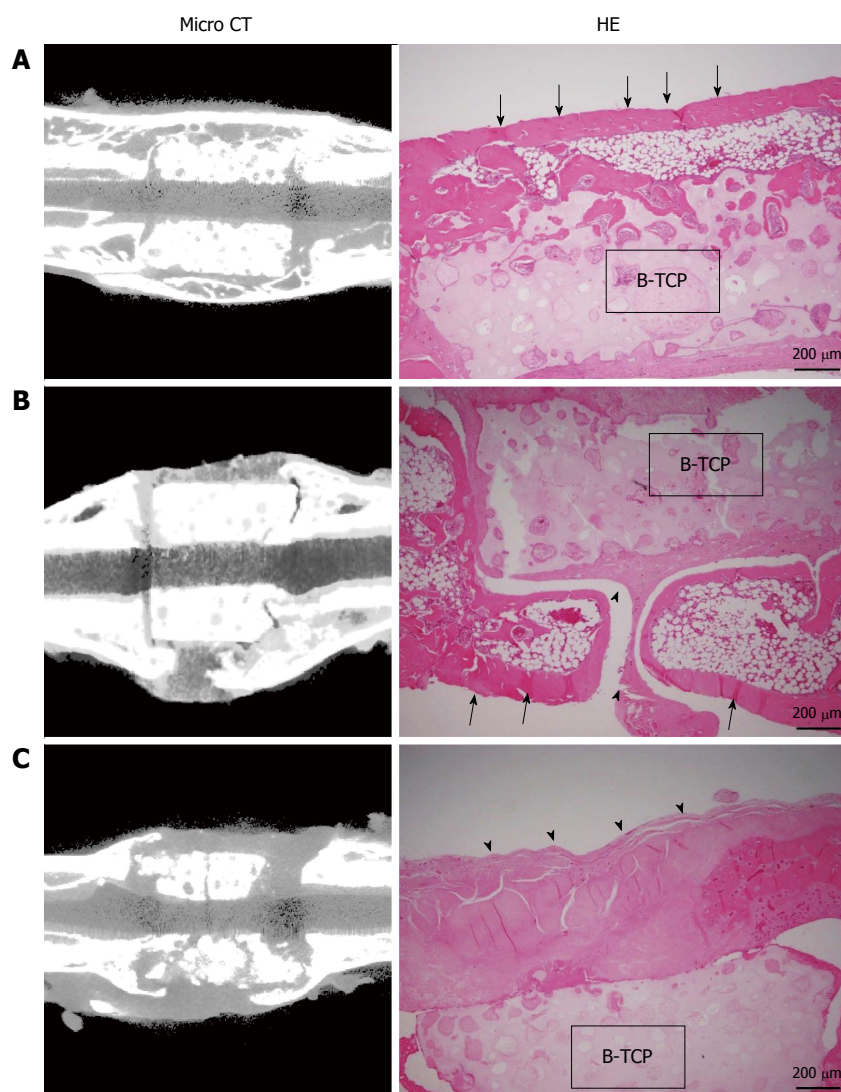


Figure 4 Micro-computed tomography images and hematoxylin and eosin-stained sections at 8 wk post-implantation. A: BMSC/TCP/Sheet constructs with continuous bone formation to host bone. Both micro-computed tomography (CT) and histology showed bridging bone formation over the TCP to host bone (arrows); B: BMSC/TCP/Sheet constructs with segmental bone formation over the TCP. Soft tissue interposition was observed between the TCP and the callus (arrowheads); C: BMSC/TCP constructs implanted into femurs. The TCP cylinders of the BMSC/TCP constructs were broken. Arrowheads indicate soft tissue interposition between the TCP and the autologous femur. BMSCs: Bone marrow stromal cells; TCP: Tricalcium phosphate.

DISCUSSION

The present study clearly demonstrated that the osteogenic potential of the BMSC/TCP/Sheet construct was significantly higher than that of the BMSC/TCP construct, namely a conventional engineered bone tissue structure. When the conventional engineered bone tissue structures were combined with an osteogenic matrix cell sheet, the newly formed bone area spread from the center to the surface of the TCP. We observed bridging bone formation over the TCP cylinder implanted between both ends of autologous bones. Moreover, biomechanical examination showed significantly higher strength against compressive stress in the BMSC/TCP/sheet construct with continuous bone formation to host bones, indicating that the BMSC/TCP/sheet construct may be useful for hard tissue reconstruction.

Some scaffolds such as TCP can be replaced with

autologous bone by bone remodeling after implantation into bone defects^[19]. Although osteogenesis does not occur in artificial bone itself, HA and TCP are common scaffolds for artificial bone in hard tissue reconstruction^[20]. Ohgushi *et al.*^[21] and Akahane *et al.*^[22] reported a transplantation technique using artificial bone combined with cells. The cells combined with artificial bone can be freshly prepared bone marrow cells or cultured mesenchymal stem cells (MSCs). In clinical cases, differentiated osteoblasts have been used in combination with scaffolds for the treatment of patients with osteoarthritis and osteonecrosis^[8,9,23]. When a BMSC suspension is combined with such scaffolds, bone formation is observed but only in the pores of the artificial bone. Therefore, when artificial bone is combined with BMSCs for transplantation into a bone defect, soft tissue interposition between the construct and the host bone might occur and result in non-union.

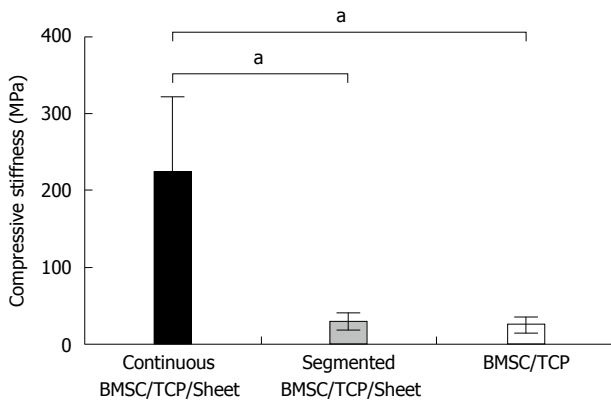


Figure 5 Results of biomechanical compression testing. The stiffness was highest in the BMSC/TCP/Sheet constructs with continuous bone formation (black column), while the stiffness of the BMSC/TCP/Sheet constructs with segmental bone formation (gray column) was similar to that of the BMSC/TCP constructs (white column). Values are means \pm SD. ^a $P < 0.05$. BMSCs: Bone marrow stromal cells; TCP: Tricalcium phosphate.

Several studies have reported tissue-engineered bone combined with BMSCs for reconstruction of bone defects^[24-26]. Kadiyala *et al.*^[26] showed that a critical-size segmental defect of the femur in a canine model can be healed by transplantation of culture-expanded cell-loaded artificial bone. However, defects that received no implant, cell-free artificial bone, or artificial bone loaded with fresh marrow failed to heal at 8 wk. Dallari *et al.*^[25] reported critical-size defect reconstruction using BMSCs, platelet-rich plasma (PRP), and freeze-dried bone allografts alone and in combination. They concluded that the combination of freeze-dried bone allografts, BMSCs, and PRP accelerates bone healing and remodeling processes. However, the bone healing was evaluated by radiology, histology, and histomorphometry in these previous studies. In contrast, we evaluated bone reconstruction by mechanical testing (compression testing) as well as micro-CT, radiology, and histology because the mechanical strength of the reconstructed site is an important clinical outcome.

The starting cell numbers were different for the three groups in this study. It would be very difficult to control the cell numbers because of the different modalities used for construct preparation. Considering clinical cases of bone defect reconstruction, tissue-engineered bone with a vigorous osteogenic potential should be used for better clinical outcomes. Therefore, we first evaluated the osteogenesis of the three types of constructs, BMSC/TCP, Sheet/TCP, and BMSC/TCP/Sheet, using the subcutaneous implantation model to determine the highest osteogenic construct, and found that the BMSC/Sheet/TCP construct had the most vigorous osteogenic potential. The number of cells that can be combined with artificial bone materials like TCP is limited because suspended cells remain in the pores of the TCP. In contrast, cell sheets can be wrapped around TCP, resulting in a large amount of loaded cells on the TCP. Moreover, the Sheet/TCP

constructs showed bone formation on the surface of the construct, which is greatly advantageous for bridging bone formation.

The bone union of conventionally engineered bone tissue structures was improved by combining the construct with an osteogenic matrix cell sheet. The BMSC/TCP construct with an osteogenic matrix cell sheet (BMSC/TCP/Sheet construct) had an excellent ability to form bone, as indicated by bone formation on the TCP surface as well as in the pores of the central area. Bone formation around the TCP was continuous with the host bone when BMSC/TCP constructs with osteogenic matrix cell sheets were implanted into bone defects of the femur. This finding may have occurred because the BMSC suspension entered the pores of the central area and the osteogenic matrix cell sheet was located on the surface of the TCP ceramic. The subsequent combination of osteogenesis by the BMSC suspension and osteogenic matrix cell sheet resulted in vigorous bone formation in the subcutaneous and bone defect sites. Prevention of soft tissue invasion and achieving early bone formation at the bone defect site is clinically important for reconstruction of bone tissue. Accordingly, our transplantation technique using constructs with osteogenic matrix cell sheets is useful because the defect is filled with the osteogenic matrix cell sheet to prevent the invasion of soft tissue, resulting in early bone formation.

MSCs have the potential to replace or regenerate damaged and diseased tissue by multipotent differentiation into osteogenic, chondrogenic, and adipogenic cells among others^[4,7,27]. Recently, although the detailed molecular mechanisms remain unclear, it was reported that MSCs, including BMSCs, can interact with cells of the immune system and modulate immune responses *in vitro* and *in vivo*^[28,29]. MSCs can drive immune responses toward immunosuppression and anti-inflammation. In the present study, we used TCP constructs to repair bone defects in the femur. Scaffolds such as TCP might cause inflammatory responses when they are transplanted *in vivo*^[30]. However, the transplantation model in the present study used a combination of TCP ceramics and cell sheets created from BMSCs, and therefore inflammatory responses could be reduced by the anti-inflammatory function of BMSCs. Further studies are necessary to clarify this point.

We used an osteogenic matrix cell sheet created in a 100-mm cell culture dish to prepare BMSC/TCP/Sheet constructs. When BMSC/TCP/Sheet constructs were transplanted into bone defects of the femur, half of the constructs (four of eight constructs) formed a segmental callus. Similar to the TCP cylinders, the mechanical properties of constructs with segmental bone were still fragile. This result indicates that coverage of the entire TCP cylinder with an osteogenic matrix cell sheet is important for continuous bone formation around the TCP cylinder. Two or more sheets

or a larger sheet can be used to cover the TCP surface and increase the compressive stiffness. Further studies will establish the numbers of sheets needed to cover certain sizes of TCP cylinders.

A thermosensitive cell culture plate is commonly used to fabricate cell sheets^[31]. Memon *et al.*^[32] used culture dishes coated with thermosensitive polymers to fabricate cell sheets for implantation into a heart disease model and for clinical treatment of corneal surface dysfunction^[33]. However, other methods have been reported, such as those using magnetite nanoparticles and magnetic force^[34]. Anil Kumar *et al.*^[35] reported a method for preparation of cell sheet/HA constructs. In their study, they showed rapid and complete cellularization of HA by wrapping it with a cell sheet. Gao *et al.*^[36] reported the results of osteogenic cell sheet transplantation using rabbit MSCs. They used osteogenic cell sheets to generate bone grafts, but did not find any bone formation on the outer HA surface in their model. We^[10,13] and others^[24,37] reported a method to prepare osteogenic cell sheets cultured in osteogenic medium. Our mechanical retrieval method only requires the use of regular cell culture equipment, including a cell scraper, to fabricate the osteogenic matrix cell sheet, in which cells form the sheet during culture with Dex and AscP. On the other hand, the methods of Anil Kumar *et al.*^[35] and Gao *et al.*^[36] require specialized culture dishes coated with thermosensitive polymers. The osteogenic matrix cell sheets used in this study were sufficiently robust to handle with forceps. The cell sheet could be easily handled for processes such as combination with artificial bone, and has osteogenic potential for osteoblastic differentiation induced by Dex and AscP. Therefore, we propose that this technique can be applied to bone defects that occur because of tumor resection or wounding.

In our study, there are a few limitations to be acknowledged. First, we only used TCP cylinders of 5 mm in length. We need to conduct further studies with longer TCP cylinders to assess their bone formation ability for cell sheet transplantation. Second, the experimental period in the present study was relatively short. Therefore, a further follow-up study is needed to determine whether the implanted TCP is replaced by autologous bone. Third, the TCP was transplanted into a bone defect site that was created by bone resection. We need to assess whether the cell sheet can form bone tissue when it is applied to a bone defect in a region with a poor blood supply, such as a large bone defect with soft tissue injury after severe trauma. Finally, we need to show the osteogenic ability using human BMSCs for future clinical application.

In conclusion, the present study clearly demonstrates that constructs of BMSCs and TCP combined with osteogenic matrix cell sheets show a vigorous osteogenic potential compared with conventionally engineered bone tissue structures consisting of a BMSC/TCP construct. Because bridging bone formation over the

implanted TCP can unite the ends of the cylindrical TCP to autologous bone, the construct will be useful for hard tissue reconstruction as new tissue-engineered bone.

ACKNOWLEDGMENTS

We thank F Kunda and M Matsumura (Nara Medical University) for their technical assistance. The TCP disks were kindly provided by Olympus Terumo Biomaterials Corp., Tokyo, Japan. We did not receive any monetary compensation or financial support from the company.

COMMENTS

Background

Bone defects are difficult to treat. Effective treatment options for bone defects have been reported that include tissue engineering of bone tissue structures using bone marrow stromal cells (BMSCs). Recently, many researchers have reported methods for creating cell sheets, such as thermo-responsive polymer-grafted culture dishes and mechanical retrieval.

Research frontiers

Tissue engineering for regenerative medicine using cell sheets are becoming much more common. The transplantation of cell sheets created from BMSCs will most likely be incorporated into clinical cases in the near future. Osteogenic matrix cell sheets have the potential to greatly advance the treatment of critical size bone defects.

Innovations and breakthroughs

Constructs comprised of BMSCs, tricalcium phosphate (TCP) and osteogenic matrix cell sheets show an increased osteogenic potential compared with conventionally engineered bone tissue structures consisting of a BMSC/TCP construct. Bridging bone formation by placing osteogenic matrix cell sheets over the implanted TCP can unite the ends of the cylindrical TCP to autologous bone.

Applications

The construct will be useful for hard tissue reconstruction, such as critical bone defects after tumor resection or traumatic injuries and avascular necrosis of femoral head, by acting as new tissue-engineered bone.

Terminology

Osteogenic matrix cell sheets are cell sheet created from BMSCs. The cell sheet is lifted by a mechanical retrieval method using a scraper. Beta-TCP is one of the most commonly used artificial bone in clinical treatments.

Peer-review

The article describes about the application of tricalcium phosphate and osteogenic matrix cell sheet for bone defect reconstruction. The efficacy of bone formation with matrix cell sheets, BMSC, and beta-tricalcium phosphate has been examined. Although the result section should be described more in detail, this study provides helpful insights for understating overall regenerative medicine technique.

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P- Reviewer: Yao CL, Tanabe S **S- Editor:** Song XX **L- Editor:** A
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