Revitalization of frozen autologous bone graft using adipose-derived stem cells

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Abstract

Adipose-derived stem cells (ADSCs) are pluripotent stem cells and have multipotentiality equivalent to bone marrow-derived stem cells. We examined the ability of ADSCs to revitalize frozen bone toward clinical application. In a conditioned culture method, proliferation and signaling pathways (Smad and JNK) of osteoblasts affected by ADSCs were examined using cell proliferation and Western blotting assays, and they were significantly enhanced. In addition, in a coculture method, the gene expression levels (BMP2 and ALP1) of osteoblasts affected by ADSCs were determined using real-time reverse transcription polymerase chain reaction (RT-PCR) and they were significantly less. Therefore, both of the methods confirmed osteoblast differentiation and osteogenesis-promoting action of ADSCs. Next, the femoral shafts of 20 rats were excised, frozen in liquid nitrogen, and then reimplanted as an autologous bone transplant. The models were divided into groups with \( n = 10 \) and without ADSCs \( (n = 10) \). Histopathologic examination was performed to assess its revitalization ability, and bone revitalization was observed significantly more with than without ADSCs in week 8. The osteogenesis-promoting action of ADSCs for frozen bone was suggested. Using ADSCs to promote revitalization of frozen bone is expected in the future.

Introduction

Various methods exist for reconstruction after resection of malignant bone tumor, including recycling of the autograft, for example autoclaved, Pasteur, or irradiated autografts. Since 1999, we developed and have clinically applied a novel method of biological reconstruction using tumor-bearing autografts treated with liquid nitrogen [1]. Apart from autograft recycling, for example for simplicity and perfect fit, no propagation of pathogens, or no allergic reaction, the use of liquid nitrogen presents several advantages. It has good osteoinduction and osteoconduction potential, allows for easy attachment of soft tissue and cryoimmunology, and provides sufficient bone stock and mechanical hardness [1-4]. However, frozen bone is a useful alternative for biological reconstruction of bone defects post-tumor resection, as revitalization of bone following a transplant requires a long time. Delayed bone union was the main problem in biological reconstruction cases, and frozen autograft is used less often than autoclaved, irradiated, and Pasteur autografts. We reported a host bone-frozen bone union seen at a mean of 6.2 months postoperatively among the long-term outcomes of using frozen autografts [5]. However, we experienced some delayed bone unions after using frozen autograft, and some of these cases required a second surgery. We aimed to achieve bone revitalization and improvement of a host bone-frozen bone union to overcome delayed bone union. For example, we used a hemi cortical resection method to preserve more host bone in some cases if we could preserve the contralateral cortex of the lesion during bone tumor resection. Furthermore, we developed a new technique of reconstruction using the pedicle freezing method. These techniques allowed the autografts to obtain a greater amount of protein through blood [6,7]. Further treatments that would allow rapid bone revitalization are desirable, and we sought a nontecnological method to promote revitalization of frozen bone. We focused on stem cell therapy and particularly on adipose-derived stem cells (ADSCs).

Stem cell therapy uses pluripotent stem cells, such as embryonic (ES) and induced pluripotent (iPS) stem cells, and multipotent stem cells, such as hematopoietic and mesenchymal stem cells. Mesenchymal stem cells have different characteristics based on the tissue to be collected. They are also called bone marrow–derived stem cells (BMSCs), ADSCs, and so forth based on the tissue to be collected. BMSCs have been validated in the regenerative medicine area for a long time [8]. However, since fewer cells are obtained by one sampling and the sampling method is invasive, clinical application is not easy. ADSCs have multipotentiality equivalent to BMSCs. Additionally, as large volumes of adipose tissue can be collected easily, ADSCs can be obtained in large quantities [9]. The group of cells collected from adipose tissue with collagenase is called the stromal vascular fraction and includes endothelial cells, fibroblasts, white blood cells, and ADSCs. ADSCs can be readily harvested in large numbers from low donor sites. Seki et al. [10] reported that injected ADSCs can repair and restore the function of the impaired liver in cirrhosis. Furuichi et al. [11] reported that injection of ADSCs reduced acute tubular necrosis in the injured kidney. ADSCs are beneficial for regeneration and repair of various injured tissue. We are also conducting research on

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tissue regeneration using the stromal vascular fraction and ADSCs and have so far reported some studies, for example, studies on distraction osteogenesis [12], peripheral nerve [13], osteoarthritis [14], ADSCs sheet [15], meniscus [16], tendon-bone [17], and atypical lipomatous tumors [18]. Lee et al. [19] reported that ADSCs promote bone union after fracture. It was considered that ADSCs promote not only bone formation, but also revitalization. We examined the use of the stromal vascular fraction and ADSCs for tissue regeneration, specifically the ability of ADSCs to revitalize frozen bone in vitro and in vivo for clinical application. The humoral factors of growth and differentiation from ADSCs to osteoblasts were evaluated in vitro. Bone revitalization of frozen autograft–added ADSCs were evaluated in vivo.

Materials and methods

Cell culture of ADSCs

ADSCs were isolated according to a modification of a previously reported method as described below [16,20]. In brief, we harvested adipose tissue from the inguinal subcutaneous layer of 10-week-old Wistar rats (Japan SLC Co., Shizuoka, Japan), and the adipose tissue was washed with phosphate-buffered saline (PBS; Wako, Osaka, Japan). We pasted the adipose tissue after cutting it for 5 minutes using scissors. Type I collagenase (Wako) was dissolved in PBS to achieve 0.12% in 20 mL and was used for digesting adipose tissue in a water bath at 37°C for 45 minutes. The adipose tissue mixture was agitated every 15 minutes while digesting. Immediately after digestion was completed, we added 20 mL Dulbecco’s modified Eagle’s medium (DMEM; Wako) containing 10% fetal bovine serum (FBS; Nichirei Biosciences, Inc., Tokyo, Japan), 100 units/mL penicillin, and 100 μg/mL streptomycin (P/S; Wako), and neutralized the collagenase activity. We filtered the obtained aqueous solution twice (Cell Strainer, BD Falcon, Tewksbury, MA, USA) and collected it in a tube. We centrifuged the filtrate at 1500 rpm at 4°C for 10 minutes. The supernatant was dispensed quickly into a new tube that had been previously cooled.

Cell harvest and culture of osteoblasts

We purchased primary rat osteoblasts, isolated from Sprague-Dawley (SD) rat calvaria, from Primary Cell from Cosmo Bio Co. Ltd. (Sapporo, Hokkaido, Japan), and cultured them in phenol red-free α-minimal essential medium (αMEM; Wako) containing 10% FBS, 100 units/mL penicillin, and 100 μg/mL streptomycin (P/S) in a humidified atmosphere of 5% CO₂ at 37°C in a humified atmosphere of 5% CO₂ at 37°C in 100 mm tissue culture dishes (TPP, Trasadingen, Switzerland) and ADSCs were cultured with complete medium, same concentration as of DMEM, including FBS and P/S. After culture for 1 week, the cells were subcultured, harvested with 0.25% trypsin-ethylenediaminetetraacetic acid (EDTA; Wako), and were centrifuged at 1000 rpm for 5 minutes. These cells were cultured in the medium at 37°C in a humidified atmosphere of 5% CO₂, and the media were changed every 3 days in 1 week. These cells were subcultured every week, and the passage of 3 ADSCs after washing twice with PBS was used for the conditioned culture and coculture methods in vitro and for addition in vitro.

In vitro experiments

In the in vitro experimental model, to ascertain the paracrine effect of ADSCs for osteoblast cells, we used the conditioned culture and coculture methods. We performed cell proliferation and Western blotting assays using the conditioned cultured method, and real-time RT-PCR assays using the coculture method.

Conditioned cultured method

Conditioned medium of osteoblast cells and ADSCs was obtained as follows. Briefly, 1.0 × 10⁶ osteoblasts and ADSCs were plated on 100 mm tissue culture dishes and incubated in complete growth medium containing serum until the cells reached the appropriate cell density. The medium was removed and was washed gently with PBS twice. After PBS was removed and the medium without serum gently added, these cells were incubated for 1 day. Then, we transferred the medium to a centrifuge tube and centrifuged at 1500 rpm at 4°C for 10 minutes. The supernatant was dispensed quickly into a new tube that had been previously cooled.

Cell proliferation assay

We analyzed cell proliferation and viability using the Cell Counting Kit-8 (Wako) as described previously [21,22]. A total of 100,000 osteoblasts were seeded in 96-well plates (TPP) and incubated in the prepared conditioned culture media of osteoblasts (control group) and ADSCs (ADSCs group). After 3 days, the cells were incubated with WST-8 (4-[3-(4-iodophenyl)-2-(4-nitrophenyl)-2H-5-tetrazolio]-1,3-benzene disulphonate; 10 μL WST-8 in 100 μL medium) for 3 hours at 37°C. We recorded the absorbance of the colored formazan product produced by mitochondrial dehydrogenases in metabolically active cells at 450 nm using iMark™ Microplate Absorbance Reader (Bio-Rad Laboratories, Hercules, CA, USA) as the background value. Osteoblast proliferation affected by ADSCs was compared between the control and ADSCs groups.

Western blotting assays

One million osteoblasts and ADSCs were plated in 10 cm tissue culture dishes and incubated in complete growth medium containing serum until the cells reached the appropriate cell density. The medium was removed and was gently washed with PBS twice. After PBS was removed and medium without serum gently added, these cells were incubated for 6 hours. The medium was removed and was gently washed with PBS twice. After PBS was removed, we gently added the prepared conditioned medium of osteoblast cells (control group) and ADSCs (ADSCs group). Furthermore, these cells were incubated for 20 minutes.

We performed Western blotting analyses as described previously [21,23]. Briefly, we used 10% sodium dodecyl sulfate-polyacrylamide gel electrophoresis (SDS-PAGE) to separate the same amounts of protein. We transferred separated proteins onto membranes of polyvinylidene fluoride, and they were subsequently incubated with primary antibodies (1:1000), followed by incubation with horseradish peroxidase (HRP)-linked secondary antibodies (1:2000). We used Immobilon Western Chemiluminescent HRP Substrate (Millipore, Billerica, MA, USA) to develop the blots.

Materials

Phospho-Smad1/5 (Ser463/465; 41D10) rabbit monoclonal antibodies (mAb), Smad1 rabbit mAb, phospho-SAPK/JNK (Thr183/Thr185; 81E11) rabbit mAb, SAPK/JNK rabbit mAb, and horseradish peroxidase-conjugated anti-rabbit IgG (#7074); they were purchased from Cell Signaling Technology, Inc. (Danvers, MA, USA).
Cocultured method

The noncontact coculture was established between ADSCs (1.0 × 10^4 cells/well) in Cell Culture Inserts with 0.4 μm pores (BD Falcon) and osteoblasts (1.0 × 10^4 cells/well) in six-well plates (Cell Culture Insert Companion Plates, BD Falcon) in αMEM (ADSCs group). On the other hand, noncocultured was not established in any cells in Cell Culture Inserts and was established in osteoblasts only in six-well plates (1.0 × 10^4 cells/well) in αMEM (control group). After 3 days, ribonucleic acid (RNA) was extracted from rat osteoblasts of the ADSCs group noncontact cocultured with ADSCs and with the control group noncocultured osteoblasts.

Total mRNA isolation

We used a NucleoSpin RNA II kit (Takara Bio, Otsu, Japan) to extract RNA. We used a syringe to disrupt and homogenize each sample. Thereafter, we performed treatment according to the manufacturer's protocol, and measured the absorbance of the extracted total RNA at 260 nm with a Thermo Fisher Scientific NanoDrop Lite (Thermo Fisher Scientific, Waltham, MA, USA), and the concentration was calculated.

cDNA synthesis

We used 5 × PrimeScript RT Master Mix (Perfect Real Time; Takara Bio) to synthesize cDNA. We performed treatment according to the manufacturer's protocol and adjusted it to a total volume of 20 μL. We synthesized cDNA using a T100 Thermal Cycler (Bio-Rad Laboratories) under the following conditions: 37°C for 15 minutes, 85°C for 5 minutes, and 4°C hold.

Real-time RT-PCR assays

We added 2 μL of the obtained cDNA sample, 10 μL SYBR® Premix Ex Taq II (Tli RNaseH Plus, Takara Bio), 0.8 μL forward primer, 0.8 μL reverse primer, 0.4 μL ROX Reference Dye (50×), and 6 μL RNAse free water (Takara Bio) to make a total volume of 20 μL. Primers for rat glyceraldehyde-3-phosphate dehydrogenase (GAPDH), bone morphogenetic protein-2 (BMP-2), and alkaline phosphatase-1 (ALP-1) were purchased from Takara Biosystems. We performed reactions for rat glyceraldehyde-3-phosphate dehydrogenase (GAPDH), bone morphogenetic protein-2 (BMP-2), and alkaline phosphatase-1 (ALP-1) were purchased from Takara Biosystems. We performed reactions and amplification using an Applied Biosystems® StepOne™ Real-Time PCR System (Thermo Fisher Scientific) with 40 cycles at 95°C for 2 seconds and at 60°C for 30 seconds. We calculated the expression ratios of BMP-2 and ALP-1 to GAPDH for each sample, comparing the ADSCs and coculture groups.

In vivo experiments

The Institute for Experimental Animals, Kanazawa University Advanced Science Research Center approved the in vivo (animal model) experimental protocol. We performed surgery in accordance with the Guide for the Care and Use of Laboratory Animals published by the United States National Institutes of Health (NIH publication no. 85-23, revised 1996; Bethesda, MD), using aseptic techniques.

Surgical procedure

Ten-week-old Wistar rats (Japan SLC Co, Shizuoka, Japan) were assigned randomly to two groups (ADSCs and control groups). Each animal was anesthetized by intraperitoneal administration of 6.5 mg sodium pentobarbital (Somnopentyl1; Intervet, Tokyo, Japan), and placed in a lateral recumbent position on the operating table. A longitudinal skin incision on the lateral thigh was made over the left femur, and the quadriceps femoris and hamstrings were separated. The femoral shaft (8 mm long) was removed by osteotomy using a manual saw. The excised bone was frozen using liquid nitrogen and reimplanted by autologous bone transplantation, and intramedullary fixation was performed with a Kirschner wire (1.4 mm diameter; DePuy Synthes; Johnson & Johnson, New Brunswick, NJ, USA). The ADSCs group received an addition of 0.1 mL Type I collagen gel (Atelocollagen; KOKEN, Tokyo, Japan) plus 5.0 × 10^4 ADSCs cells around the frozen bone. The control group was without ADSCs. Last, we sutured the fascia and skin and the rats were allowed movements without the limitation of weight bearing and external fixation in their cages after anesthesia.

Histological examination

Bone revitalization and union were evaluated through histological examination. Rats were euthanized 4 and 8 weeks postoperatively. The left total femurs were resected with the surrounding tissues and we fixed all specimens in 4% paraformaldehyde in PBS, pH 7.4. They were decalcified in 10% EDTA solution, were embedded in paraffin, were sectioned in the sagittal plane, and were stained with hematoxylin and eosin. We checked stained bone and the number of frozen bone revitalizations and host bone–frozen bone unions were compared. Furthermore, revitalization of frozen bone was assessed as the ratio of the number of lacunae with nucleated cells to that of whole lacunae according to a modification of a previously reported method, and this ratio was defined as the revitalization rate (Figure 1) [24]. Six fields at low magnification (objective lens ×10) on a KEYENCE BZ-9000 microscope (Keyence Corporation, Osaka, Japan) from two proximal, two central, and two distal sites in frozen bone were selected in each specimen.

Dil labeling

We labeled ADSCs for tracking cells with a fluorescent lipophilic tracer 1,10-dioctadecyl-3,3,30,30-tetramethylindocarbocyanine perchlorate (Dil; MolecularProbes; Vybrant1 Dil Cell Labeling Solution; Life Technologies, Carlsbad, CA, USA). For labeling, we resuspended the cells at 1 × 10^6 cells/mL in DMEM and added Dil at 5 μL/mL in a medium without serum. After incubation for 20 minutes at 37°C with 5% humidified CO₂, the cells were centrifuged at 1500 rpm for 5 minutes and were washed twice with PBS, then were resuspended in Type I collagen gel (Atelocollagen; KOKEN) for addition around the frozen bone. We performed this examination to determine the location of ADSCs post-addition of ADSCs. A fresh frozen sagittal section of the left total femur was prepared using Kawamoto's method.
Statistical analysis

We performed this as an observer-blinded study, and it was not
different between each group. The experimental results were assessed
as mean ± standard error. We performed a t-test to compare the control
and ADSCs groups in vitro and a Mann-Whitney’s U test in vivo.
Differences were considered significant when P-value was <0.05.

Results

Cell proliferation assays

To ascertain the paracrine effect of ADSCs on osteoblasts, their
cell viability was assessed using a conditioned culture method. The cell
viability of osteoblasts in the ADSCs group was found to be higher than
that in the control group (P = 0.034; Figure 2).

Western blotting assays

A comparison between the phosphorylation ratios of Smad1 to
total Smad1 showed the phosphorylation ratio to be higher in the
ADSCs than in the control group (P = 0.027). Furthermore, comparison
of c-Jun N-terminal kinase (JNK) and total JNK showed a higher
phosphorylation ratio in the ADSCs than in the control group (P = 0.021).
This suggested that the conditioned culture of ADSCs enhanced
the Smad and JNK pathways of the differentiation and osteogenesis
signal of osteoblasts (Figure 3).

Real-time RT-PCR assays

The ADSCs group had lower BMP2 and ALP1 expression levels
compared to the control group (expression ratios 0.184 and 1,
respectively; P = 0.039). The expression ratio of ALP1 was 1 in the
control group and 0.399 in the ADSCs group (P = 0.041; Figure 4).

Histological examination

In week 4, bone revitalization was observed in two of five cases
in the control group, and one of five in the ADSCs group. In week 8,
bone revitalization was observed in both of five and five of five cases,
respectively, suggesting an osteogenesis-promoting action of ADSCs in
frozen bone. Revitalization rate was significantly higher in the ADSCs
group than the control group. In week 4, the control group had a
higher revitalization rate (4.2%) compared to the ADSCs group (1.7%),
however, the difference was not significant (P = 0.602). In week 8, the

Figure 2. Cell proliferation assays for osteoblasts cultured with conditioned medium of
osteoblasts (control group) and adipose-derived stem cells (ADSCs) (ADSCs group). The
cell viability of osteoblasts in the ADSCs group was higher than that of osteoblasts in the
control group (P = 0.034).

Figure 3. Western blotting assays for Smad1 and c-Jun N-terminal kinase (JNK) signal of
osteoblasts cultured with conditioned medium of osteoblasts (control group) and adipose-
derived stem cells (ADSCs) (ADSCs group). A comparison between the phosphorylation
ratios of Smad1 showed the phosphorylation rate to be higher in the ADSCs than in the
control group (P = 0.027). Similarly, a comparison between the phosphorylation ratios of
JNK showed a higher ratio in the ADSCs group than in the control groups (P = 0.021). This
suggested that the conditional culture of ADSCs enhanced the Smad and JNK pathways of
the differentiation and osteogenesis signal of osteoblasts.

Figure 4. Real-time reverse transcription polymerase chain reaction (RT-PCR) assays for
BMP2 and ALP1 osteoblast cocultured with rat adipose-derived stem cells (ADSCs) in
noncontact coculture (ADSCs group) and noncocultured rat osteoblasts (control group).
The ADSCs group had lower BMP2 and ALP1 expression levels compared with the control
group using coculture method (expression ratio of BMP2 0.184 versus 1, respectively; P = 0.039).
The expression ratio of ALP1 was 1 in the control group and 0.399 in the ADSCs group (P = 0.041)
ADSCs group had a higher revitalization rate (24.8%) compared to the
control group (5.1%). This difference was significant (P = 0.0283; Figure
5 and 6). In week 4, bone union was not observed in the control and
ADSCs groups. Bone union was observed in both groups in week 8;
however, the difference was not significant.

DiI labeling

DiI labeling indicated that the transplanted cells had survived
for 4 weeks after transplantation of ADSCs. DiI-positive region were
detected during the 4 weeks after transplantation. Locally distributed
transplanted cells were observed in soft tissue surrounding the frozen
bone and not in the frozen or normal bone (Figure 7).
ADSCs secrete cytokines, such as BMP2, that contribute to the osteogenic activity of ADSCs [26]. There are two signaling pathways for BMP signaling. The main BMP signaling pathway is through receptor I-mediated phosphorylation of Smad1, Smad5, or Smad8 (R-Smad). The other BMP pathway is mediated by transforming growth factor-β1 (TGFβ1) activated tyrosine kinase 1 (TAK1), and TAK1 can activate JNK signaling pathway by BMP led to negative feedback resulting in reduction of BMP2 osteoblast production. The conditioned culture and coculture method studies suggested that ADSCs have paracrine effects of differentiation and osteogenesis of osteoblasts.

Although differentiation of mesenchymal stem cells (MSCs) transplanted locally contribute to part of the mechanism for the therapeutic effects, such as repair or regeneration, the survival and differentiation of MSCs is limited, and most of the therapeutic effect is paracrine signaling [32]. Dil labeling suggested that the transplanted ADSCs did not differentiate into bone but were focally distributed in soft tissue surrounding the frozen bone. ADSCs did not expand into autologous frozen bone but remained around the frozen bone by week 8. From the results of an in vitro study and Dil labeling, transplanted ADSCs had a paracrine effect that led to frozen autograft revitalization.

Nather et al. [33] reported that the addition of autologous BMSCs to frozen allograft improved bone union, new bone formation, and osteocyte cell counts. An in vivo study suggested that autologous frozen ADSCs and BMSCs had a faster cell proliferation rate and higher bone differentiation ability than BMSCs [25]. A cell proliferation assay using the conditioned culture method suggested that osteoblasts had increased cell proliferation capacity in the ADSCs proliferative environment, many ADSCs may produce much protein, and it was effective for osteoblasts of paracrine signaling. In this study, the humoral factors of ADSCs had a good effect on the bone environment; however, the paracrine effects of BMSCs were not confirmed for osteoblasts.
bone grafting of rat, in the control and ADSCs groups, showed bone revitalization ability at week 4 postoperatively. In our study, bone revitalization was significantly higher in the ADSCs group than in the control group at week 8 postoperatively. ADSCs have high bone forming and bone differentiation abilities at week 8; however, ADSCs did not demonstrate significant improvement of the host bone–frozen bone union. Nather et al. [33] used rabbits larger than the animals used in our study and they used plates for fixation of host bone–frozen bone. On the other hand, we used intramedullary fixation with a Kirschner wire, and the rats were allowed movements without the limitation of weight bearing and external fixation in their cages postoperatively. Therefore, postoperative load at the site of operation was not allowed. Intramedullary fixation of bone in our study was a bit weak and no-load limit increased the bone junction’s force point. If we used a much stronger fixation implant or load limit condition, the revitalization rate in week 4 would be changed. Kloe ters et al. [34] reported on new vascularization and osteocyte proliferation of bone allograft by application of vascularized scaffolds seeded with osteogenic-induced ADSCs (O-ADSCs) induce. We have not confirmed O-ADSCs. Revitalization of frozen bone may be different from adding ADSCs and O-ADSCs. In addition, we used fewer animals in this study. Additional study using larger and more numbers of animals is required before clinical examination can be performed.

Recently, there were many reports of regeneration medicine using cell sheets. Uchihara et al. [35] reported that osteogenic matrix cell sheets’ transplantation facilitated osteogenesis in irradiated rat bone. We reported that ADSCs sheets have better capability than separate ADSCs, which are not sheets, to be induced to differentiate into osteoblasts [15]. We used Type I collagen gel plus ADSCs cells around the frozen bone, and the transplanted cells were focally distributed in soft tissue surrounding the frozen bone. If we used the cell sheets, much transplanted cells might be distributed. We will try to study the ADSCs sheets.

Some studies have reported malignant tumor and ADSCs. ADSCs promote tumor growth in epithelial ovarian and breast cancers [36,37] or inhibit tumor growth in hepatocellular carcinoma and glioblastoma [38,39]. A relatively low proportion of ADSCs suppressed the growth of osteosarcoma, whereas high proportions showed a tumor-promoting effect [40]. Stromal ADSCs promote osteosarcoma progression and metastasis [41]. ADSC-secreted factors may not be involved in the risk of local recurrence [42]. It is unclear how ADSCs affect each tumor. There is no unified view. It is also necessary to evaluate the influence of ADSCs on malignant cells; for example, osteosarcoma or metastatic tumor cells to determine potential clinical applications. We examined the necrotic bone model using frozen autograft; however, we need to study the interaction of ADSCs with the tumor by the tumor bone model.

Various clinical trials have shown the regenerative capacity of stromal vascular fraction or ADSCs in medical subspecialties, such as cardiovascular disease, gastroenterology, plastic surgery, urology, orthopedic surgery, and so forth [43]. Some clinical trials were related to malignant tumor, especially breast cancer [44,45]. As reconstruction option after breast conservation therapy, adipose-derived regenerative cells were used for a prospective clinical trial, and no recurrences and no serious adverse events were reported [44]. Therefore, using ADSCs with frozen autograft after resection of malignant bone tumor may be a new reconstruction option.

Conclusion

In vitro, promotion of osteoblast proliferation and osteogenesis-promoting action of ADSCs was confirmed. In vivo, the possibility of promoting frozen bone revitalization was also suggested and clinical application for reconstruction of bone defects after resection of malignant bone tumors is expected in the future.

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Conflicts of interest

The authors indicate no potential conflicts of interest.

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