Periodontal tissue regeneration by transplantation of adipose tissue-derived multi-lineage progenitor cells

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Several stem and progenitor cells are currently under investigation for their application in cell-based therapy for periodontal tissue regeneration. The present study evaluates periodontal tissue regeneration following transplantation of adipose tissue-derived multi-lineage progenitor cells (ADMPCs) into periodontal tissue defects in beagle dogs.

ADMPCs were isolated from the greater omentum and their characteristics were identified using in vitro studies. Flow cytometric analysis demonstrated that the isolated ADMPCs were CD29+, CD44+, and CD90+. When cultured in mineralization-inducing media, these cells upregulated osteogenic genes and formed calcified nodules. Gene expression of the periodontal ligament specific gene, PLAP-1, was also increased. In addition, culture in adipogenic media resulted in accumulation of intracellular lipid droplets, suggesting multi-lineage differentiation capability of ADMPCs.

The efficacy of ADMPC transplantation for periodontal regeneration was evaluated using a beagle dog model. The furcation bone defects were surgically created and autologous transplantation of ADMPCs with fibrin gel was performed. Six weeks after transplantation, periodontal regeneration was analyzed using micro-CT, which showed a significant increase in bone formation at sites where ADMPCs were applied compared with control sites. Histological analysis revealed new cementum formation on the instrumented root surface was significantly increased following ADMPC transplantation and connective tissue fibers were inserted vertically in newly formed bone and cementum. Importantly, no instances of undesirable healing, such as root resorption or ankylosis, were observed at any sites examined. These results indicate that transplantation of ADMPCs with fibrin gel promotes periodontal tissue regeneration.

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Introduction

Periodontitis is an inflammatory disease caused by bacterial biofilms formed on the dental root surface. This disease can lead to irreversible destruction of the tooth-supporting tissues, including the alveolar bone, periodontal ligament, root cementum, and gingiva. Although conventional periodontal treatments based on mechanically removing the cause of disease shows some success in controlling inflammation and suppressing the progression of periodontitis, unfortunately these treatments rarely regenerate lost periodontal tissue or its functionality to any statistically or clinically significant level. Thus, periodontal regeneration therapy is required for successful periodontal treatment, which can be accomplished by re-formation of all components of the periodontium, including cementogenesis and osteogenesis.

Periodontal regeneration can be achieved, to some extent, by a variety of periodontal regenerative therapies, such as bone grafting, guided-tissue regeneration, application of enamel matrix derivatives, and cytokine therapies. However, the results are variable and the indication is limited to mild-to-moderate periodontal tissue defects. This may be because these therapies depend on the activity of somatic stem cells that exist within periodontal ligament tissue. As reported previously, an in vitro study revealed that aging can affect the proliferation and mineralized nodule formation of periodontal ligament cells and the number of stem cells within periodontal ligament tissue\(^6\). Moreover, progress of periodontitis not only destroys periodontal tissue but also deprives of existing somatic stem cells. Thus, it is important to use stem cells isolated from other tissues for periodontal regeneration, with a view to enhancing the outcome predictability and expanding the indication of regeneration therapy to severe periodontal defects\(^5\).

Embryonic stem cells, induced pluripotent stem cells and somatic stem cells, are the three major stem cell sources. Owing to their safety and ethical profiles, somatic mesenchymal stem cells (MSCs) have been under intense investigation for clinical use in cell therapies. In terms of periodontal regeneration, transplantation of bone marrow-derived MSCs and periodontal ligament stem cells have been examined in preclinical\(^3,4\) and clinical studies\(^5,6\), highlighting their advantageous effects. However, difficulties associated with the isolation of these cells, including tissue harvesting, cell purifying from a heterogeneous cell population, and patient morbidity from associated harvesting pain, need to be overcome.

Adipose tissue-derived stem cells have been identified as a pluripotent cell population in adipose tissue, able to differentiate into several cell types\(^7\). These cells are also reported to secrete a variety of cytokines, including HGF, VEGF, TGF-β, IGF-1, and FGF-2, which are favorable factors for angiogenesis\(^8\). This cellular profile is thought to be important for the application of these cells in regenerative medicine. In addition, stem cells can be easily and safely obtained from adipose tissue without issues regarding tissue availability and ethics. To date, one group demonstrated that transplantation of adipose tissue-derived stem cells enhanced periodontal regeneration in the presence of platelet-rich plasma in rats and dogs\(^9,10\). However, several issues remain to be addressed, including the effects solely from the transplanted cells and the safe and optimal scaffold material for these cells.

In this study, we isolated adipose tissue-derived multilineage progenitor cells (ADNPCs) from the greater omentum of beagle dogs and evaluated the cell characteristics. We then examined the effects of autologous transplantation of ADNPCs with fibrin gel into a furcation periodontitis model.

Materials and Methods

1) Experimental animals

Female beagle dogs, 50-56 months old, weighing 9-11 kg, were used and analyzed for experiments. All protocols were approved by the Institutional Animal Care and Use Committees of Osaka University Graduate School of Dentistry.

2) Isolation of ADNPCs

After beagle dogs were subcutaneously and intravenously anesthetized with 1 mL xylazine (Bayer Yakuhin, Ltd., Osaka, Japan) and 10 mg/kg pentobarbital (Kyoritsuseiyaku Co., Tokyo, Japan), respectively, the greater omentum was resected from each subject. ADNPCs were prepared as described previously\(^11\). Briefly, the resected greater omentum was minced and digested at 37°C for 1 h in 0.075% collagenase (Wako Pure Chemical Industries, Osaka, Japan) in Hank’s balanced salt solution. Digested tissue was diluted by adding Dulbecco’s modified Eagle’s medium high glucose (DMEM-HG; Gibco Life Technologies, CA, USA) with 20% fetal bovine serum (FBS; Sigma-Aldrich, St. Louis, MO, USA) and centrifuged at \(\times 400\) g
for 10 min. The cell pellet was suspended with DMEM-HG with 10% FBS and red blood cells were excluded using density gradient centrifugation with Histopaque (d=1.077 g/ml; Sigma). Cells were then filtered through a 40 µm cell strainer and cultured in DMEM-HG with 10% FBS at 37°C for 24 h. Following incubation, cells were washed with phosphate buffered saline (PBS) and treated with 0.02% ethylenediaminetetraacetic acid (EDTA) solution (Nacalai Tesque, Kyoto, Japan). Floating cells, called ADMPCs, were collected and seeded onto a fibronectin-coated dish (BD Biosciences, San Jose, CA, USA) in 60% DMEM-low glucose (Gibco), 40% MCDB-201 medium (Sigma).

ADMPCs analyzed in vitro and in vivo were at passages 3 and 3.4, respectively.

3) Flow cytometric analysis

Isolated canine ADMPCs were characterized by flow cytometry. Single-cell suspensions were prepared by trypsinization by 0.05% trypsin / EDTA (Gibco) and incubated for 30 min at 4°C with fluorescein isothiocyanate (FITC)-conjugated antibodies; mouse anti-canine CD29 (EXBIO, Praha, Czech Republic), rat anti-canine CD44 (eBioscience, San Diego, CA, USA) and mouse anti-canine CD105 (BD Biosciences). Incubation with rat anti-canine CD90 (Thermo Scientific, Rockford, IL, USA) was followed by reaction with FITC-conjugated mouse anti-rat IgG antibody (BD Biosciences). Isotype-identical antibodies served as controls for each antigen. Data were collected with a FACSCalibur (BD Biosciences) and analyzed with CellQuest software.

4) Osteogenic and adipogenic differentiation procedure

For osteogenic differentiation, ADMPCs were cultured in 12-well plates until confluent and were then exposed to mineralization medium consisting of alpha modification of minimum essential medium; α-MEM supplemented with 10% FBS, 10 mM β-glycerophosphate, and 5 µM ascorbic acid, which was replaced every 3 days. Differentiation was examined by histochemical staining of calcified nodules with alizarin red S. Cell monolayers were washed twice with PBS and then fixed with dehydrated ethanol. After fixation, cell layers were stained with 1% alizarin red S (Wako) in 0.1% NH₄OH (pH 6.5) for 5 min, then washed with H₂O. For adipogenic differentiation, ADMPCs were cultured in 24-well plates until confluent and were then exposed to adipogenic-inducing medium consisting of α-MEM supplemented with 5% FBS, 500 µM 3-Isobutyl-1-methylnethane (Sigma), 10 µM Insulin (Sigma), 100 µM Indomethacin (Sigma), and 1 µM Dexamethasone (Sigma), which was replaced every 3 days. Differentiation was examined microscopically by observation of intracellular lipid droplets after staining with Oil Red O as follows; cell monolayers were fixed with 4% paraformaldehyde for 30 min, washed and stained with 0.16% Oil Red O for 20 min.

5) Reverse transcription-PCR

Total RNA was extracted from cultured cells using RNA-Bee (TEL-TEST, Inc., Friendwood, CA, USA) in accordance with the manufacturer's protocol. Purified total RNA was reverse transcribed using M-MLV (Invitrogen, Carlsbad, CA, USA) reverse transcriptase with random hexamers. Real-time PCR analysis was performed using Power SYBR Green PCR Master Mix and a 7300 Fast Real-Time PCR system (Applied Biosystems, Foster City, USA). The primer sequences used for PCR are listed in Table 1.

6) Surgical procedure and transplantation of ADMPCs

Extensive bone defects around the molar furcation area were surgically created in mandibular right and left premolars (P3 and P4) of beagle dogs, as shown in Fig.1. All surgeries were performed under pentobarbital anesthesia and local infiltrated anesthesia with 2% lidocaine hydrochloride and 1/80,000 epinephrine. After elevation of the mucoperiosteal flaps, the buccal bone was removed to create the bone defects (4 mm inferior and 3 mm horizontal). The

Table 1. Nucleotide sequences of primers used for PCR

<table>
<thead>
<tr>
<th>Gene</th>
<th>Primer sequence</th>
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<tr>
<td>COL1A2</td>
<td>5'- GCACATGCGGAGACCTGAGA -3'</td>
</tr>
<tr>
<td></td>
<td>R: 5'- GCATCCAGAGTCATCTTGTTAG -3'</td>
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<tr>
<td>RUNX2</td>
<td>5'- TTCGGCCTGAAACCTGATGA -3'</td>
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<td></td>
<td>R: 5'- GCCGGCAACAAATCTCAG -3'</td>
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<tr>
<td>PLAP-1</td>
<td>5'- TGCGCGCCAGCAGATC -3'</td>
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<tr>
<td></td>
<td>R: 5'- GTGCCGATCATTCTTCT -3'</td>
</tr>
<tr>
<td>HPRT</td>
<td>5'- GCAGCTATACTCAAGATGTA -3'</td>
</tr>
<tr>
<td></td>
<td>R: 5'- TCAGGTTTATAGCCAACACTCGC -3'</td>
</tr>
<tr>
<td>LPL</td>
<td>5'- CTCACTGGTCTGTGATGGA -3'</td>
</tr>
<tr>
<td></td>
<td>R: 5'- GTGTGTAATGTCATTGC -3'</td>
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exposed cementum of the teeth was removed using Gracey Curettes (Hu-Friedy, Co. Chicago, IL, USA) and tooth planing bars. Vinyl polysiloxane impression material (Putty type: GC Corporation, Tokyo, Japan) was placed in all furcation defects to initiate inflammation. The wounds were closed by suturing the flaps over the furcation defects and stitches were removed after 1 week. Four weeks after the first surgery, the full thickness flap was raised to expose the inflamed furcation and granulation tissue was removed. After root planing using a root planing bar, a horizontal groove on each root was made by a small round bar to indicate the base of the defect. The furcation defect of the right and left side was then filled with fibrin gel (Boheal®; Teijin Pharma Limited, Tokyo, Japan) alone and the ADMPC-fibrin gel complex, respectively, and surgical closure was performed.

7) Micro-computed tomography (μCT) analysis
Under general anesthesia, beagle dogs were euthanized by exsanguination 6 weeks after transplantation. Tissues containing the transplanted region were extracted and fixed with 4% paraformaldehyde. Tissues were then scanned using a μCT apparatus (Scan Xmate-E090; Comscan Techno, Tokyo, Japan). Three-dimensional-CT (3D-CT) images were analyzed and quantified using image analysis software (TRI/3D-BON, Ratoc System Engineering, Tokyo, Japan).

8) Histological analysis
After μCT analysis, tissues were decalcified for 2 months in 10% formic acid, dehydrated, and embedded with paraffin. Serial sections (4 μm) were prepared in the mesial-distal plane. The mesio-distal plane in the center of the horizontal defect of each furcation involvement was then stained with Azan staining. Staining was performed as follows: deparaffinized sections were treated with dye mordant for 10 min and then stained with azocarmine G solution (Muto Pure Chemicals Co. Ltd., Tokyo, Japan.) for 30 min at 60°C. After differentiation in anilin-ethanol, reaction was stopped with 1% acetic ethanol. Then sections were treated with 5% phosphotungstic acid solution for 60 min, followed by anilin blue-orange G solution (Muto Pure Chemicals Co. Ltd.) for 30 min. And then dehydration and differentiation was performed with 100% ethanol. Histological measurements were performed using WinRoof® image analysis software (Mitani Corp., Tokyo, Japan). The new cementum formation rate was histomorphometrically calculated by dividing the length of newly formed cementum by the perimeter of intra-bony defect.

9) Statistical analysis
Data were expressed as the mean ± standard deviation (SD). Statistical analyses were performed by the Student's t-test. p<0.05 was considered statistically significant.

Results
1) Surface marker expression of ADMPCs
To assess MSC marker expression on ADMPCs isolated from the greater omentum, flow cytometric analysis was
Periodontal regeneration by ADMPC transplantation

Fig.2 Characteristics of ADMPCs
(A) MSC surface marker expression on ADMPCs. Expression of each surface marker is shown with the shaded histogram. Staining with an isotype control antibody is shown with the black line. (B) Type I collagen, Runx2, and PLAP-1 mRNA expression in ADMPCs cultured with (gray bar) or without (white bar) mineralization media (Min-M) for indicated days. Data are expressed as relative expression to HPRT. (C) Mineral deposition by ADMPCs cultured with or without Min-M. Cells were stained with alizarin red S on the indicated days.

performed. Isolated ADMPCs were found positive for the markers characteristic of canine mesenchymal stem cells, including CD29, CD44 and CD90, and were negative for CD105 and CD140a (Fig.2A), as previously reported[12].

2) Differentiation capability of ADMPCs
To examine the differentiation capability of ADMPCs, cells were cultured in mineralization media to induce osteoblastic differentiation. Gene expression of type I collagen and Runx2 were significantly upregulated by induction (Fig.2B). Moreover, alizarin red S staining showed calcified nodule formation after 24 days of culture (Fig.2C). Notably, mRNA expression of PLAP-1, a specific molecule expressed in periodontal ligament tissue[13], was also increased (Fig.2B), suggesting not only osteoblastic capability but also potential to differentiate into periodontal ligament tissue. In addition, adipogenic differentiation was identified by gene expression of leptin and lipoprotein lipase (LPL) and oil droplet formation, after culture in an adipogenic-inducing medium (Fig.3). These results suggest the canine ADMPCs isolated from the greater omentum had multi-lineage differentiation capability.

3) Effects of ADMPC transplantation into the furcation periodontal defects
Inflamed furcation bony defects were artificially prepared and transplanted with either fibrin gel alone or autologous ADMPCs with fibrin gel. No severe inflammation or swelling was observed in any examined sites throughout the experimental period. Moreover, furcation areas in all experimental sites remained unexposed during the experimental period. As shown in Fig.4, μCT analysis revealed that transplantation of ADMPCs significantly enhanced alveolar bone regeneration compared with fibrin gel alone.
Fig. 4  Quantitative analysis of new bone formation by \( \mu \)CT

(A) \( \mu \)CT reconstruction of the periodontal tissue defects transplanted with fibrin gel alone (control) or with ADMPCs + fibrin gel (ADMPCs). Six weeks after transplantation, mandibular tissues were harvested and scanned with \( \mu \)CT. Yellow area shows newly formed alveolar bone. (B) Quantitative analysis of reconstructed newly formed alveolar bone. (n=5) Data are expressed as mean ± S.D. *p<0.05 compared with control.

Fig. 5  Histological analysis of periodontal tissue regeneration

Histological overview of furcation bone defects in the mesio-distal plane 6 weeks after transplantation. Histologic sections were stained with Azan. Representative image of fibrin gel applied site is shown in (A). Higher magnification of (A) is shown in (C). Representative image of fibrin gel + ADMPCs transplanted site is shown in (B). Higher magnification of (B) is shown in (D) and (E). Scale bar in (A) and (B)=1 mm. Scale bar in (C) and (D)=100 \( \mu \)m. Scale bar in (E)=50 \( \mu \)m. D: dentin, NB: new bone, P: periodontal ligament tissue, arrow head: newly formed cementum.

Fig. 6  Histomorphometric analysis of new cementum formation

(A) Schematic drawing of histometric analysis of cementum formation 6 weeks after transplantation. Newly formed cementum rate=(a-b + a’-c)/a-a’ x 100 (%). (B) Quantitative analysis of newly formed cementum rate. Data are expressed as mean ± S.D. *p<0.05 compared with control.
Histological observations of the furcation defects demonstrated periodontal regeneration, including new alveolar bone, and periodontal ligament and cementum formation with vertically inserted fibers in ADMPC transplanted sites (Fig.5). No undesirable healing occurred at ADMPC-transplanted sites, including root resorption or ankylosis. Histomorphometric analysis was performed and the new cementum formation rate was determined as described in Fig.6A. The new cementum formation rate following ADMPCs transplantation was significantly higher than those at control sites (Fig.6B).

**Discussion**

Current regenerative procedures to address periodontal tissue damaged following periodontitis have limitations in achieving complete and predictable regeneration. Based on tissue engineering, stem cell transplantation therapy is expected to overcome these issues. The present study demonstrates for the first time, the positive effects of ADMPCs, in the absence of extrinsic signaling molecules, in the treatment of periodontal defects in beagle dogs.

ADMPCs are readily extracted, can be expanded efficiently in vitro, and have the capacity to differentiate into multiple cell lineages, all of which are criteria in regenerative medicine. In fact, reports about the clinical application of ADMPCs have increased in a wide variety of fields. In most cases, ADMPCs are isolated from subcutaneous adipose tissue harvested by liposuction. Because there is insufficient adipose tissue under canine skin, the greater omentum was used as an ADMPC source in this study. Although it has been reported that ADMPCs from visceral fat have less proliferation and differentiation capacity compared with those from subcutaneous adipose tissue\(^\text{14,15}\), ADMPCs used in this study significantly promoted periodontal regeneration. This finding promises greater impact of ADMPCs on future clinical trials, because subcutaneous adipose tissue is abundant and more readily accessible in humans.

During research into periodontal regeneration, a variety of animal models have been used. In this study, we employed the furcation periodontitis model in beagle dogs to evaluate the effects of ADMPC transplantation. Although we artificially induced inflammation in the defect site before ADMPC transplantation, the majority of periodontal regeneration preclinical studies have applied various biomaterials or cells just after surgical creation of defects. Our preclinical model is close to the clinical situation and our results enhance the utility value of ADMPCs for periodontal regeneration. In addition, we utilized clinical-grade fibrin gel (Bolheal\(^\text{16}\)) as a scaffold for ADMPCs. Our preliminary study demonstrates that application of Bolheal\(^\text{16}\) into periodontal defects does not impair the healing of periodontal tissue compared with sham procedure (data not shown).

ADMPCs possess the capacity to undergo osteogenic, adipogenic, chondrogenic, and myogenic differentiation. Notably, we showed not only the osteogenic and adipogenic differentiation of isolated ADMPCs, but also the induction of PLAP-1 gene expression, a specific marker of periodontal ligament tissue (Fig.2), suggesting the differentiation capacity of these cells into periodontal ligament tissue. ADMPCs had not been induced to undergo differentiation prior to transplantation. It could therefore be hypothesized that transplanted ADMPCs home to and differentiate into specialized cells in the transplanted sites adequately. Although this hypothesis needs to be demonstrated in future studies, this is supported by previous studies in which transplanted stem cells in periodontal bone defects, at least in part, differentiated into cementoblasts, osteoblasts, osteocytes and fibroblasts in regenerated periodontal tissue\(^\text{10}\).\(^\text{16}\).

Recent work, on the other hand, demonstrates that the mechanism by which stem cells participate in tissue repair and regeneration is related to their trophic factors\(^\text{17,18}\). In fact, our pilot studies show that ADMPCs secrete a number of growth factors, and conditioned medium from ADMPCs can stimulate differentiation of periodontal ligament cells to hard tissue-forming cells (data not shown). These results suggest a possible mechanism that transplanted ADMPCs secrete a number of factors, which can activate the tissue surrounding the periodontal defect, resulting in promotion of the regeneration process.

In conclusion, transplantation of ADMPCs with fibrin gel into periodontal defects has a positive effect on periodontal regeneration. Further investigations will be required to reveal the molecular mechanisms of this therapy and the true potential of this procedure, including the extent of periodontal destruction that can be rescued. It also needs to be determined which signaling molecules or scaffold materials or a combination of these give ADMPCs the optimal environment to regenerate periodontal tissue. As a consequence, we are confident that periodontal regeneration
therapy, achieving the ultimate goal of complete periodontal tissue regeneration following periodontal disease, will be developed.

Source of Funding and Conflict of Interest
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