Differentiation and characteristics of undifferentiated mesenchymal stem cells originating from adult premolar periodontal ligaments

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Objective: The purpose of this study was to investigate the isolation and characterization of multipotent human periodontal ligament (PDL) stem cells and to assess their ability to differentiate into bone, cartilage, and adipose tissue.

Methods: PDL stem cells were isolated from 7 extracted human premolar teeth. Human PDL cells were expanded in culture, stained using anti-CD29, -CD34, -CD44, and -STRO-1 antibodies, and sorted by fluorescent activated cell sorting (FACS). Gingival fibroblasts (GFs) served as a positive control. PDL stem cells and GFs were cultured using standard conditions conducive for osteogenic, chondrogenic, or adipogenic differentiation.

Results: An average of 152.8 ± 27.6 colony-forming units was present at day 7 in cultures of PDL stem cells. At day 4, PDL stem cells exhibited a significant increase in proliferation (\(p < 0.05\)), reaching nearly double the proliferation rate of GFs. About 5.6 ± 4.5\% of cells in human PDL tissues were strongly STRO-1-positive. In osteogenic cultures, calcium nodules were observed by day 21 in PDL stem cells, which showed more intense calcium staining than GF cultures. In adipogenic cultures, both cell populations showed positive Oil Red O staining by day 21. Additionally, in chondrogenic cultures, PDL stem cells expressed collagen type II by day 21.

Conclusions: The PDL contains multipotent stem cells that have the potential to differentiate into osteoblasts, chondrocytes, and adipocytes. This adult PDL stem cell population can be utilized as potential sources of PDL in tissue engineering applications.

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INTRODUCTION

Many researchers have shown that bone marrow-derived cells can be differentiated into various mesenchymal or connective tissues, leading to recent investigations focusing on the ability of stem cells to repair damaged tissues. Stem cell studies of the maxillofacial area have evaluated the effects of the regeneration of cartilage and bone tissues in temporomandibular joints, the regeneration of periodontal tissues, and bone regeneration in oral cancer defects using stem cell and gene therapy. Stem cells can be classified as mesenchymal cells (MCs; referring to mesenchymal stem cells at the prenatal stage) or mesenchymal stem cells (MSCs; referring to mesenchymal stem cells after birth). MCs are known to be the best for the regeneration of all human tissues; however, due to technical difficulties and bioethical arguments, MC research has been met with significant limitations; therefore, adult MSCs are frequently used for research. Adult MSCs play a major role in physiological tissue remodeling and are also able to induce the regeneration of damaged tissues. These processes can be applied in various fields. However, since MSCs have to be collected from the bone marrow, researchers have developed novel methods (albeit complicated and limited) that utilize MSCs from fascia and fat tissues.

Homeostasis between teeth and the alveolar bone is maintained through continuous remodeling of the periodontal ligament (PDL), even after orthodontic tooth movement. PDLs are connective tissues coupling the space between the tooth's cementum and the alveolar bone. They maintain the tooth at a regular position within the alveolar bone, supply nutrition to the surface of roots, and are key factors involved in the regeneration of damaged roots and alveolar bone. Replacement root resorption during childhood results in PDL loss, which can result in defective eruption and ankylosed teeth. These defective teeth may then have a lower occlusal surface than proximal teeth (infraclosure). PDLs also stabilize the position of teeth and participate in alveolar bone growth and development, remodeling themselves throughout their lifetime. Recent studies have suggested that the PDL space may contain MSCs that are able to differentiate into cementoblasts and osteoblasts, producing mineral tissues and fibroblasts that form fibrous connective tissues. These MSCs are thought to originate from capillaries in bone tissues and the PDL space, ultimately differentiating in the PDL space. Therefore, teeth and PDLs have emerged as potential important sources of undifferentiated MSCs, even in adults, where acquisition of MSCs is limited to the bone marrow and fascia.

In the dental field, Gronthos et al. reported the successful isolation and differentiation of undifferentiated MSCs from surgically extracted mandibular third molars. Since then, many researchers have also successfully isolated MSCs from uninfected impacted third molars and dental pulp tissues. However, all of these studies are restricted to fairly young, unerupted third molars, which are not easy to obtain from general dental clinics.

The aim of this research was to determine whether undifferentiated MSCs isolated from the root surface of adult premolars extracted for orthodontic treatment could be cultured to differentiate into bone, cartilage, and fat tissue.

MATERIALS AND METHODS

Cell culture

Maxillary first premolars were extracted from 7 patients (5 men and 2 women, ages 20.3 - 28.4 years) before orthodontic treatment. All procedures were approved by the ethics committee of Pusan National University Hospital Medical Institute (2008-2) and were performed after obtaining informed consent agreements from patients. Tissues from PDLs were carefully excised with a scalpel from the middle third of freshly extracted teeth. The PDL tissue was digested in 3 mg/mL collagenase type I (Invitrogen, Carlsbad, CA, USA) and 4 mg/mL dispase (Invitrogen) for 1 h at 37°C. Single-cell suspensions were obtained with a 70-μm cell strainer (Falcon BD, Franklin Lakes, NJ, USA). To separate cells expected to be stem cells, we allowed suspensions of 1 × 10⁴ cells to attach to 100-mm dishes (NUNC, Roskilde, Denmark) in alpha modification of Eagle's medium (alpha-MEM; Gibco BRL, Carlsbad, CA, USA) supplemented with 10% fetal bovine serum (FBS; Gibco BRL), 100 mM ascorbic acid 2-phosphate (Sigma, St. Louis, MO, USA), 2 mM glutamine, 100 U/mL penicillin, and 100 mg/mL streptomycin (Gibco BRL). Cells were then cultured for 10 days in 5% carbon dioxide at 37°C. Gingival fibroblasts (GFs) derived from the gingival tissue attached to the cementoenamel junctions of the extracted premolars were used as controls for cell proliferation assays. All cells used for further assays were second or third passage cells.

Colony formation assays

For colony formation assays, 1 × 10⁴ cells were plated in a 100-mm petri dish and cultured for 7 days. After fixation for 20 min with 5 mL of 4% paraformaldehyde, cells were stained with 0.05% crystal violet to assess colony formation efficiency. Fifty or more cells clustered together were considered a colony.

Fluorescent activated cell sorting (FACS)

Cell suspensions from single cells were acquired using a 70-μm cell strainer (Falcon BD). Next, 1 × 10⁴ cells were added to a FACS tube in blocking buffer (HBSS, 10% FBS, 1% streptomycin, 2 mM glutamine, 100 U/mL penicillin).
20 mM HEPES, 1% normal human serum, 1% bovine serum albumin, 5% FBS), and 10 μL of anti-STRO-1 immunoglobulin (Ig)M (Santa Cruz Biotechnology, Santa Cruz, CA, USA). Anti-human CD29 (integrin β-1 receptor marker for cell proliferation and adhesion; BD Bioscience, Sparks, MD, USA), anti-human CD34 (specific hematopoietic stem cell marker; BD Bioscience), and anti-human CD44 (hyaluronate receptor that facilitates adhesion to stromal progenitors during cell culture but not to hematopoietic progenitors; BD Bioscience) antibodies were also added. Cells were then cultured for 30 min at 4°C. Anti-mouse K antibodies were used as a negative control. After primary antibody incubation, goat anti-mouse IgM FITC secondary antibodies (2 μg/mL; Molecular Probes, Eugene, Carlsbad, CA, USA) were added, and cells were incubated for an additional 1 h. Positively stained cells were separated by flow cytometry and analyzed with the automated cell deposition unit FACScan (Becton Dickinson, Parsippany, NJ, USA) at 4°C. For comparison of MSC marker expression, all experiments were performed separately for each patient.

**Cell proliferation assay**

To estimate cell proliferation capacity, 1 × 10^4 cells were seeded and cultured in DMEM media containing 20% FBS and 1% antibiotics for 1 week. Coulter cell counters (Coulter Electronics Inc., Hialeah, FL, USA) were used to determine the number of cells.

**Differentiation**

**Osteogenic induction (calcification in colony-forming assay, quantitative assay of calcium content)**

Seven dishes of 3 × 10^4 cells each were seeded and cultured for 3 days. Osteogenesis-inducing media (α-MEM containing 1% acetic acid, 10% FBS, 0.1 μM dexamethasone, 50 μM ascorbic acid 2-phosphate, and 10 mM β-glycerophosphate) was changed every 2 - 3 days for 3 weeks to induce osteogenesis. After 3 weeks, the culture was washed 3 times with phosphate buffered saline (PBS) and fixed for 15 min with 4% paraformaldehyde. One hundred microliters of substrate solution was added, and cells were cultured for 30 min at 37°C. Next, 100 μL stop solution (0.9 N NaOH) was added, and calcium formation was determined by 2% Alizarin red S staining (300 μL; Sigma-Aldrich) after a 15-min fixation with 70% ethanol and washing with PBS.

**Adipogenic induction (adipogenesis during colony formation assays)**

Cells were seeded into 7 dishes and cultured for 3 days. Adipogenesis-inducing media (α-MEM containing 1% acetic acid, 10% FBS, 0.5 mM isobutylmethylxanthine, 1 μM dexamethasone, 10 μM insulin, and 100 μM indomethacin) was changed every 2 - 3 days for 3 weeks to induce lipogenesis. After 3 weeks, cultures were washed 3 times with PBS, fixed for 15 min with 4% paraformaldehyde, and then washed 3 more times with PBS. The dishes were stained for 2 h at room temperature with 3% Oil Red O (Sigma-Aldrich). After washing the cultures 3 times with PBS, photographs were taken. Staining was quantified by the addition of 500 μL isopropanol after drying.

**Chondrogenic induction**

To induce chondrogenesis, 3 × 10^5 cultured cells were placed into a 15-mL tube for centrifugation. Chondrogenic inducing media (Cambrex Bio Science, Charles City, IA, USA) with 10 ng/mL transforming growth factor (TGF)-β3 (Cambrex Bio Science) was changed every 2 - 3 days for 3 weeks. Collagen type II immunohistochemistry was performed to check collagen formation. Culture tubes were frozen for 30 min and fixed with acetone for 10 min. After drying, the culture was washed twice with PBS and then primary antibodies (collagen type II, Sigma-Aldrich) were added. After 24 h, secondary antibodies were used.

**Statistical analysis**

Differences in the number of colony forming units (CFUs) between PDL stem cells and GFs were compared using t-tests. Sexual differences between donors were not shown to be significant due to the small sample sizes. Statistics were calculated using SPSS for Windows (version 16.0; SPSS Inc., Chicago, IL, USA).

**RESULTS**

PDL stem cells exhibited more rapid proliferation than control GFs

Fifty or more cells clustered together were considered a CFU. Control GFs exhibited an average of 26.3 ± 13.4 CFUs after 7 days, while undifferentiated MSCs exhibited an average of 152.8 ± 27.6 CFUs (Figures 1 and 2). As shown in Figure 1, there was a statistically significant difference in the total number of CFUs between PDL (n = 7) and GF (n = 7) cultures. In addition, we analyzed the cell proliferation rate over the course of 6 days and found that the proliferation rate of PDL stem cells was over 2 times higher than GFs at day 4 and nearly 3 times higher at day 6 (Figure 3).

**FACS analysis revealed expression of proliferation and stem cells markers in MSCs**

FACS analysis was performed using antibodies specific for undifferentiated MSCs. CD29 and CD44 were expressed in an average of 52.2 ± 21.4% and 99.6 ± 0.2% of cells, respectively. In contrast, CD34-positive cells were
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found at a very low rate of 0.5 ± 0.4%. An average of 5.6 ± 4.5% of cells were positive for the stem cell-specific marker, STRO-1 (Table 1 and Figure 4).

Figure 1. Cell culture colony-forming units (CFUs). At day 7 after plating, PDL cells were fixed with 10% formalin and stained with 4% crystal violet. Aggregates of 50 or more cells were considered colonies and counted. A, A representative GF colony (10x). B, A representative PDL cell colony (10x). The black arrows and circles represent the images of 7-day cell population (right panels) at 10x magnification using light microscopy.

Figure 2. Colony forming units (CFUs) in gingival fibroblasts (GFs) (n = 7) and periodontal ligament (PDL) cells (n = 7). Colony formation was measured as described in the Material and Methods section. PDL cells exhibited a statistically significant increase in CFUs as compared to GFs, the positive control. *p < 0.05.

Figure 3. Cell proliferation rates. Periodontal ligament (PDL) cells and gingival fibroblasts (GFs) were cultured for 1, 2, 3, 4, 5, and 6 days in 6-well plates with media changes every other day. The cells were harvested by trypsinization and counted with a Coulter counter. The data are from 1 of 7 representative experiments, each yielding similar results. From day 4, PDL cell proliferation was significantly increased as compared to GFs, the positive control. Diamonds correspond to PDL cells, and squares correspond to GFs. *p < 0.05, †p < 0.01.
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PDL cells exhibited osteogenic, chondrogenic, and adipogenic differentiation potential

**Osteogenic induction**

After culturing PDL cells and GFs in osteogenesis-inducing media for 21 days, calcified crystals were observed in both cell lines. PDL cells in particular contained an abundance of calcified crystals. In contrast, both cell types showed no formation of calcified crystals when cultured with normal media (Figure 5).

**Chondrogenic induction**

Chondrogenesis was induced by culturing cells in specific differentiating media for 21 days. Immunohistochemical staining for collagen type II revealed positive staining in all cell groups. Furthermore, when cultured in chondrogenesis-inducing media, both PDL cells and GFs demonstrated chondrogenic induction with metachromasia and cell morphology corresponding to embryonic stages of cartilage formation (Figure 6).

**Adipogenic induction**

Following culture in adipogenesis-inducing media for 21 days, both PDL cells and GFs were positively stained for an adipose tissue marker. Staining was faint in GFs, and both cell types exhibited no evidence of adipogenic induction after 21 days of culture in normal media (Figure 7).

**DISCUSSION**

Many researchers have investigated the potential applications of tissue engineering for restoration of teeth and periodontal tissues after suffering disease or trauma. Indeed, there has been a huge focus on studies examining potential methods for grafting and successful biological substitutes. However, these methods have failed to result in successful tissue regeneration. To overcome this limitation, tissue regeneration using stem cells is now being studied. However, stem cell-related work is limited by ethical issues surrounding embryonic stem cell research as well as experimental challenges related to the limited supply of adult stem cells. Therefore, the discovery of new, robust sources of stem cells is a major focus of many researchers. While adult stem cells do not exhibit totipotency like embryonic stem cells, they still have regenerative, proliferative, and differentiating potential. Adult stem cells can be divided into hematopoietic cells and undifferentiated MSCs according to their differentiating potential. Undifferentiated MSCs have the ability to differentiate into specific cells and possess clonogenic potential, that is, the potential for a single cell to form a cluster of identical cells. Undifferentiated MSCs become progenitor cells, i.e., partially differentiated cells (in morphogenic terms), before finally progressing into mature cells. The PDL space contains massive amounts of fibroblasts, osteoblasts, and cementoblasts for regeneration of PDL tissues. Recent studies have shown that PDL spaces contain progenitor cells, which are assumed to differentiate into these various cells. However, it is still unclear whether these cells are progenitors or undifferentiated MSCs that have the potential to become any kind of cell. The purpose of this study was to determine whether undifferentiated MSCs exist on the adult premolar root surface and, furthermore, to determine whether these cells could differentiate into various progenitor cells in vitro.

In this study, we collected PDL cells from the maxillary first premolar root surface and cultured these cells for 2 passages. Seven days later, CFUs were counted following culture in stem cell-specific medium. Culturing cells at a low density (1 × 10⁴) for 7 days resulted in the growth of over 150 CFUs. In a previous study, Seo et al. also reported the isolation of over 170 single colonies assumed to be stem cells after isolation of PDLs from the third molar. Single cells exhibited morphologies similar to fibroblasts when visualized by light microscopy. Moreover, Pittenger et al. demonstrated that undifferentiated MSCs are

| Sex     | Age  | Negative (%) | CD29 (%) | CD34 (%) | CD44 (%) | STRO-1 (%) |
|---------|------|--------------|----------|----------|----------|------------|
| Donor 1 | Female | 28.4         | 0.4      | 75.4     | 0.4      | 99.7       | 3.4        |
| Donor 2 | Male  | 24.7         | 1        | 45.8     | 0.5      | 99.5       | 2.6        |
| Donor 3 | Male  | 20.3         | 0.7      | 65.3     | 0.8      | 99.5       | 14.7       |
| Donor 4 | Male  | 22.4         | 0.8      | 78.7     | 1.3      | 99.6       | 8.6        |
| Donor 5 | Female | 23.1         | 0.6      | 43.6     | 0.3      | 99.2       | 3.1        |
| Donor 6 | Male  | 24.5         | 0.5      | 22.1     | 0.2      | 99.8       | 2.9        |
| Donor 7 | Male  | 22.3         | 0.2      | 34.7     | 0.2      | 99.8       | 4          |
| Mean±SD |       | 23.7 ± 2.6   | 0.6 ± 0.3| 52.2 ± 21.4 | 0.5 ± 0.4| 99.6 ± 0.2 | 5.6 ± 4.5  |

FACS, Fluorescent activated cell sorting; SD, standard deviation.
Figure 4. Primary digested pooled adult human periodontal ligament cells were immunostained with various antibodies, and cells were analyzed on a FACScan with an automated cell deposition unit equipped with an argon laser. The data were analyzed with Cell Quest software. These representative FACScan cell distributions were collected from a passage 2 cell population from Donor 1. Percentages of CD29-positive (A), CD34-positive (B), CD44-positive (C), and STRO-1-positive (D) cells were identified and are indicated by the circle in each corresponding graph.
similar to fibroblasts in terms of morphology, but exhibit differences in their replicating or proliferative potential. In this study, the proliferative potential of cultured PDL cells from day 3 showed a nearly 2.5-fold increase in proliferation when compared to fibroblasts on day 6, indicating that the isolated PDL cells contained undifferentiated MSCs.

The most common way to collect undifferentiated MSCs is from the bone marrow.\textsuperscript{[12]} However, cells from the bone marrow are mostly hematopoietic stem cells, and only approximately 0.1% of these cells are undifferentiated MSCs. Therefore, separating these 2 cell populations is a major challenge.\textsuperscript{[3]} A common method to separate undifferentiated MSCs is flow cytometry using FACS. FACS uses undifferentiated MSC-specific surface antibodies to mark cells with fluorescence or differential light absorbance for isolation (“sorting”).\textsuperscript{[28]} Therefore, it is important to identify and utilize appropriate markers for undifferentiated MSCs during FACS analysis. A commonly used undifferentiated MSC-specific marker is STRO-1. Mouse-raised IgM STRO-1 antibodies do not respond to hematopoietic stem cells, but instead attach to a surface antigen expressed on progenitor MSCs in human bone marrow. According to experiments performed by Simmons and Torok-Storb,\textsuperscript{[29]} STRO-1-positive cells are morphologically similar to fibroblast clonogenic CFUs. However, erythrocyte progenitor cells can also stain positive for STRO-1; therefore, most tests are conducted using several surface antibodies simultaneously. In this study, we also used anti-CD29 (integrin β-1 receptor),\textsuperscript{[30]} -CD44 (hyaluronate receptor),\textsuperscript{[30]} and -CD34 (hematopoietic stem cell marker)\textsuperscript{[31,32]} antibodies simultaneously. Baddoo et al.\textsuperscript{[30]}

Figure 5. Expression of mineral deposits within periodontal ligament (PDL) cells and gingival fibroblasts (GFs) after long-term culture in osteogenesis-inducing media. A, C, No calcium deposits were seen in PDL cells and GFs cultured in normal media (20×). B, GFs cultured with osteogenesis-inducing media exhibited relatively small calcium deposits at day 21; the distribution was observed as weak foci associated with cell clusters (20×). D, At day 21, PDL cells cultured with osteogenesis-inducing media showed mineral deposits as evidenced by Alizarin red S staining (20×), with deposits distributed throughout the tissue culture well.
reported that undifferentiated MSCs from bone marrow express both CD29 and CD44. Therefore, MSCs can be collected by screening out CD29- and CD44-negative cells. Additional reports have shown that CD34 is expressed on hematopoietic stem cells during culture, but not on MSCs, making it a useful screening tool. Using flow cytometry analysis, we found that between 2.6% and 14.7% of second-passage PDL-derived cells were positive for STRO-1. Moreover, these cells exhibited an average of 52.2%, 99.6%, and 0.5% cells that were positive for CD29, CD44, and CD34, respectively. In this analysis, however, staining with CD29 antibodies resulted in a larger standard deviation (i.e., 52.2 ± 21.4%) than staining with other antibodies. Thus, it is possible that CD29 is expressed in all proliferating cells within PDL cell pools. Gay et al. reported that an average of 27% of PDL-derived cells were STRO-1-positive, this was higher than the percentage of positively stained cells in our study. These differences may be related to the sources of MSCs as Gay’s study used the PDL and dental follicles of an
impacted third molar with an immature root end. In addition, Sawa et al.\textsuperscript{33} insisted that differences in the PDL cell cycle of an immature permanent tooth and a fully matured one were the result of aging. This supports our data because all of our sources of PDL cells were from fully matured maxillary premolars from adults older than 20 years. Unfortunately, the correlation between age and the number of undifferentiated MSCs was not examined in this study. Further studies with a larger data pool would be required to investigate this.

To determine if PDL cells differentiate successfully, calcium production was tested after culturing with osteogenesis inducing media for three weeks. This study resulted in overall high production of calcium in PDL cells from osteogenesis inducing media and from some GF cells as well. However, cultured in normal media, PDL cells and GF cells did not produce calcium. This result coincides with Gronthos et al.,\textsuperscript{17} Seo et al.,\textsuperscript{18} Gay et al.,\textsuperscript{19} Shi et al.\textsuperscript{20} Many various factors are required for an undifferentiated MSC to differentiate into osteoblast or cementoblast cells. Studies show that dexamethasone and ascorbic acid have a close relationship with the differentiation of osteoblasts. Our osteogenesis inducing media contained these substances as well. However, experiments carried out in vivo used bone morphogenic protein 2 (BMP2) more frequently.\textsuperscript{34,35} For clinical use, studies of growth factors like BMP2 are required.

To determine whether PDL cells have the potential to differentiate into various mature tissues, we first investigated whether they could form adipose tissue using Oil red O staining of PDL cells grown for 3 weeks in lipogenesis-inducing growth media. The main lipogenesis-inducing component was indomethacin, which acts as an antagonist to the peroxisome proliferator-activator receptor alpha (PPAR-\alpha).
activator receptor (PPAR) gamma on the surface of undifferentiated MSCs, thereby activating lipoprotein lipase within adipose-forming cells. Lecka-Czernik also demonstrated that application of PPAR gamma 2 to odontoblasts allowed these cells to differentiate into adipose tissue. However, in this study, we were unable to isolate and identify adipose tissues. Thus, further studies are required in order to shed more light on this process.

Cartilage tissues generally do not exist in GF cells. However, it is important to confirm whether cartilage tissues could form from undifferentiated MSCs of PDL origin since MSC-derived cartilage tissues could be used in temporomandibular joint disc regeneration. By using collagen type II staining, we demonstrated that both GFs and PDL cells could form cartilage tissues when cultured in TGFβ3-containing, cartilage-inducing media. TGFβ3 is known to be a major factor involved in cartilage formation in MSCs. Preliminary studies have demonstrated that these cells do not exhibit cartilage formation in normal media. However, through aggregation of cells by centrifugation and culturing with appropriate inducing media and TGFβ3, cartilage formation was induced. We expect that this is a result of the interaction between proteoglycans secreted from cells of connective tissues. Further studies are required to develop and adapt techniques for tissue engineering of cartilage cells.

CONCLUSION

In this study, we sought to identify whether PDL cells could be classified as undifferentiated MSCs and to test their potential to differentiate into bone, cartilage, and fat tissues. From this study, we found that PDL cells, collected and isolated from the maxillary first premolars of 7 patients, exhibited a 2-fold greater proliferation rate than GFs. We also successfully identified a certain amount of undifferentiated MSCs in the pool of PDL cells and confirmed that these cell populations could form bone, adipose, and cartilage tissues. These findings suggested that periodontal ligament cells from extracted adult premolars may be useful sources of stem cells for oral and maxillofacial tissue engineering in the future.

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