Characterization of deciduous teeth stem cells isolated from crown dental pulp

Karacterizacija matičnih čelija izolovanih iz zubne pulpe mlečnih zuba dece

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Abstract

Background/Aim. The last decade has been profoundly marked by persistent attempts to use ex vivo expanded and manipulated mesenchymal stem cells (MSCs), as a tool in different types of regenerative therapy. In the present study we described immunophenotype and the proliferative and differentiation potential of cells isolated from pulp remnants of exfoliated deciduous teeth in the final phase of root resorption.

Methods. The initial adherent cell population from five donors was obtained by the outgrowth method. Colony forming unit–fibroblast (CFU-F) assay was performed in passage one. Cell expansion was performed until passage three and all tests were done until passage eight. Cells were labeled for early mesenchymal stem cells markers and analysis have been done using flow cytometry. The proliferative potential was assessed by cell counting in defined time points and population doubling time was calculated. Commercial media were used to induce osteoblastic, chondrogenic and adipogenic differentiation. Cytology and histology methods were used for analysis of differentiated cell morphology and extracellular matrix characteristics.

Results. According to immunophenotype analyses all undifferentiated cells were positive for the mesenchymal stem cell markers: CD29 and CD73. Some cells expressed CD146 and CD106. The hematopoietic cell marker, CD34, was not detected. In passage one, incidence of CFU-F was 4.7 ± 0.5/100. Population doubling time did not change significantly during cell subcultivation and was in average 25 h. After induction of differentiation, the multiclonal derived cell population had a tri-lineage differentiation potential, since mineralized matrix, cartilage-like tissue and adipocytes were successfully formed after three weeks of incubation.

Conclusion. Altogether, these data suggest that remnants of deciduous teeth dental pulp contained cell populations with mesenchymal stem cell-like features, with a high proliferation and tri-lineage differentiation potential and that these cultures are suitable for further in vitro evaluation of cell based therapies.

Key words:

dental pulp; stem cells; tooth, deciduous; child, preschool; cell differentiation; adipogenesis; chondrogenesis; osteogenesis.

Apstrakt

Uvod/Cilj. Prošla dekada je bila posebno obeležena napomina na polju korišćenja ex vivo razvijenih i usmeravanih mezenhimskih matičnih čelija (MSCs), kao sredstva za različite tipove regenerativne terapije. Cilj ove studije bio je da se utvrdi imunofenotip i potencijal za proliferaciju i diferencijaciju čelija izolovanih iz zubne pulpe mlečnih zuba dece eksfoliranih u periodu kada je koren zuba bio u poslednjoj fazi resorpcije.


Proliferativni potencijal i vreme udvajanja čelija (PDT) u kulturi je definisano na osnovu apsolutnog broja broja čelija na početku i na kraju svake pasaze. Posle tronodjelne kultivacije čelija u komercijalnim međijumima za stimulaciju osteogenze, kondrogeneze i adipogeneze, citometrijski i histoloskim metodama je određena morfologija čelija i karakteristike vancelijonskog matriksa. Rezultati. Antibij koji karakterišu mezenhimske matične čelije CD29 i CD73 su bili ekspimirani na svim nedeficiranim čelijama, dok su antiženi CD146 i CD106 bili ekspimirani na ograničenom broju čelija. Antižen CD34 (karakterističan za čelije hematopoetske loze) nije bio ekspimiran. Incidenca CFU-F bila je 4,7 ± 0,5/100 čelija. PDT se nije menjao tokom osam pasaža i u proseku je iznosio 25 h. Posle tronodne stimulacije diferencijacije u kulturama sa adipogenim međijumom došlo je
Introduction

The last decade has been profoundly marked by persistent attempts to use ex vivo expanded and manipulated mesenchymal stem cells (MSCs), as a tool in different types of regenerative therapy. Most of them were focused on healing sloughing and dental medicine.

Research targeting cell therapy and tissue engineering in regeneration of tooth structures, was stimulated after Grothos and coworkers described dental pulp stem cells (DPSC) isolated from impacted third molars of adult donors. Clinically interesting populations of cells have also been isolated from deciduous teeth. Thus, Miura et al. described stem cells from human exfoliated deciduous teeth (SHED) and Kerkis et al. obtained immature DPSC (IDPSC) from the same source. Subsequently, several more papers confirmed these findings and enlarged our knowledge about stem cells that could be isolated from deciduous dental pulp. In the review of Kerkis and Caplan all isolated cell populations were named deciduous teeth stem cells (DTSC) with the conclusion that they have a higher colony forming capacity and a higher proliferation rate than DPSC, and therefore are more primitive than their counterparts isolated from permanent teeth. The pluripotent nature of DTSC, and the fact that teeth develop from oral ectoderm and neural crest-derived mesenchyme, led some investigators to conclude that these cells display developmental potential similar to embryonic stem cells. In vivo, SHEDs generate a tissue with morphological and functional properties that closely resemble those of human dental pulp and strongly induce bone formation. Besides stem cells from dental pulp, periodontal ligament stem cells, dental follicle progenitor cells, stem cells from apical papilla and even stem cells from periapical lesions have been described. Development in the field of biomaterials and tissue engineering, together with stem cell research, has shown promising results for the development of optimal restorations to replace lost tooth structures.

Special interest in characterization of stem cell populations that can be found in dental pulp of human exfoliated deciduous teeth is underlined by the fact that these cells are easily obtained. Instead of being discarded, they could be cryopreserved, and if necessary, expanded and used for autologous or allogeneous treatment.

The aim of this study was to test the proliferation and differentiation potential of cells isolated from dental pulp of human exfoliated deciduous teeth in the final phase of root resorption, and to describe their colony forming capacity, population doubling time, immunophenotype in the undifferentiated state and their tri-lineage differentiation capacity. The term DTSC will be used subsequently for the cell population isolated in this work.

Methods

Isolation of the initial cell population

Deciduous incisor teeth from children aged 6 and 7 years (5 patients) were obtained after extraction due to orthodontic reasons, under local anesthetic, with informed consent of their parents and ethical comity approval. Teeth roots with the mesial half of their crowns that contained gelatinous pinkish pulp tissue were used. Dental pulp was pulled out with a barbed Nuvbroach, washed twice with sterile phosphate buffered saline (PBS) supplemented with antibiotics (100 U/ml penicillin and 100 µg/ml streptomycin) and antimycotic (2.5 µg/ml amphotericin B) – AA solution (antibiotic/antimycotic). Pulp tissue was minced into 1–2 mm fragments, transferred to 35 mm Petri dishes and cultivated using Dulbecco’s modified Eagle’s medium (DMEM) / Ham’s F12 (1:1, Invitrogen, Carlsbad, California, USA) supplemented with 10% FBS and penicillin and streptomycin. Cultures were incubated at 37°C in a humified atmosphere with 5% CO2.

The growing culture of the initial cell population (passage 0) was maintained for 10 to 15 days, dissociated in a 0.05% TrypLE™ Express (Invitrogen, Carlsbad, California, USA), and seeded at 1 × 105 cells per 25 cm2 flasks. Cultures initiated as multicounty culture systems were maintained semi-confluent in order to prevent premature senescence. Thus, the cells were passed every 5 days, while the medium was replaced every 2–3 days. Cells were used from passages three to eight.

Colony forming unit fibroblast (CFU-F) assay

After harvesting cells from passage 0, single-cell suspensions (1 × 103 cells) designated passage 1, within DMEM/F12 containing 10% FBS were seeded into T 25 tissue culture flasks (BD Falcon, Becton, Dickinson and Company – BD, NJ, USA). After 10 days, cultures were fixed with 10% methanol, and then stained with Crystal violet solution. Aggregates containing 50 or more cells were counted as CFU-F under the microscope.

For assessment of colony-forming efficiency (CFE), cells in the fourth passage were plated at a density of 500
cells in six-well plates and colony formation was inspected under a microscope after 7 days of culture. The CFE index was calculated by dividing the number of colonies formed by the number of cells plated and multiplying with the factor 100.

**Population doubling time**

For analysis of population doubling time (PDT), cells were seeded at a density of $1 \times 10^4$ cells/well in six-well plates. The cell number was assessed after 4 days, with a hemocytometer, after collecting the wells by trypsinization (3 replicates for each time point). PDT was calculated by the formula: $PDT = \left[ \ln (N_t / N_0) / \ln (2) \right] / t$ ($t$ = the time period, $N_t$ = number of cells at time $t$ and $N_0$ = initial number of cells).

**Flow cytometry**

After harvesting, cells (third to sixth passage) were washed in cold PBS supplemented with 0.5% BSA (Sigma-Aldrich, Saint Louis, MO, USA). Aliquots of $5 \times 10^5$ cells were labeled (30 min in the dark at $4^\circ C$) with monoclonal antibodies specific for human markers associated with mesenchymal and hematopoietic lineages. Namely, mouse anti-human antibodies against the following antigens were used: CD34 (PE conjugated), CD29 PECy5 conjugated, CD73 and CD146 (PE conjugated) and CD106 (FITC conjugated), all purchased from BD Biosciences. To determine the level of nonspecific binding, fluorochrome conjugated isotype control antibodies were used. Flow cytometry was performed using a CyFlow CL (Partec, Münster, Germany).

**Differentiation**

Complete commercial media (StemPro Osteogenesis, Chondrogenesis and Adipogenesis Kits, Gibco-Invitrogen, Carlsbad, CA, USA) were used to induce osteogenesis, chondrogenesis and adipogenesis of DTSCs from the third to sixth passage. Characteristic features of differentiated cells were visualized by cytochemical and/or histochemical methods.

After 3 weeks in complete medium (changed every 2 days) for osteogenesis, calcium depositions were demonstrated in the extracellular matrix. Cell layer was washed twice in PBS and fixed with 10% neutral buffered formalin (NBF) for 1 h at room temperature (RT). Cultures were then stained with 1% Alizarin red S solution (Sigma-Aldrich, Saint Louis, MO, USA), pH 4.2, for 20 min at RT, followed by rinsing three times with deionized water. After 3 weeks in complete medium for adipogenesis (changed every 2 days), cells were fixed in 4% paraformaldehyde for 8 h, rinsed twice with PBS, then treated with 60% isopropanol (until evaporation), stained with a fresh 0.35% Oil Red O solution for 10 min, followed by washing twice with deionized water. The chondrogenic differentiation potential of the expanded cells was investigated by micromass culture. The cell solution of $2 \times 10^5$ viable cells was prepared in chondrogenic or control medium. Tubes were centrifuged at 1,000 rpm for 6 min allowing cells to aggregate at the tube bottom. Pellets were formed after 24 h. After 2 and 3 weeks (medium changed every 2 days), pellets were fixed in 4% non-buffered formaldehyde for 24 h, embedded in paraffin and 5 μm thick sections were prepared. Sulfated glycosaminoglycans (GAG) were demonstrated with 0.1% Alcian blue (Sigma-Aldrich, Saint Louis, MO, USA) counterstained with 0.1% Fast nuclear Red (Sigma-Aldrich). The presence of collagen type II was detected immunohistochemically using rabbit polyclonal antibodies to collagen type II (Abcam, Cambridge, MA, USA).

All the quantitative data are presented as mean ± standard deviation. Data were processed in Excel for Windows program.

**Results**

Our study demonstrated that after adhesion of dental pulp explants to plastic, initial cell migration was obtained in 2 to 3 days, followed by rapid cell proliferation. Initial cell growth was designated passage 0. The number of cells harvested after passage 0 was $0.3 \times 10^5$ to $3 \times 10^5$. In passage 1, the incidence of CFU-F was 4.7 ± 0.5 per 100 cells (Figure 1).

In our experiment, in all time points, the PDT was approximately 25 h (Figure 2a). Colony forming efficiency (CFE) in passage four was 80.4 ± 7.5% on average (Figure 2b). Using flow cytometry, we demonstrated that all cells expressed CD29 and CD73 (Figures 3a and 3b), 88% of cells expressed CD146 (Figure 3c), and 5% of cells expressed CD106 (Figure 3d). We also confirmed that CD34 was not expressed on the cell population examined (Figure 3e). Figure 3f demonstrate mean expression of the analyzed markers from five donors.

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The differentiation potential of isolated cells is important when considering their potential to regenerate specified tissues, like bone, cartilage or adipose tissue. After 3 weeks of cultivation in adipogenic medium, the cells became more round and filled with fat droplets (Figure 4a) while cells...
in the control media had scarce and small fat droplets (Figure 4b). Highly induced calcium deposition in ECM was demonstrated with Alizarin red staining after 3 weeks of culture (Figure 4c). In control media no Alizarin red staining was noted (Figure 4d). After 2 and 3 weeks in chondrogenic medium, small, compact pellets rich in cells were formed. Cells produced the ECM with positive GAGs (Figure 4e) and collagen type II (Figure 4f) staining. Due to the loose tissue structure, pellets in control medium were decomposed during paraffin embedding protocols.
Discussion

Our study demonstrates that remnants of crown dental pulp of exfoliated deciduous teeth contain a population of cells that migrate in vitro, form CFU-F and have high colony forming efficiency. The CFU-F assay is a useful tool to demonstrate, among primary isolated cells, single ones with sufficient proliferative potential to form colonies of several hundred to a thousand cells. In analogy with the hematopoietic system hierarchy, those cells may form a population of progenitor cells with tri-lineage, bi-lineage or uni-lineage potential. The CFU-F frequency could represent the tissue potential for generating enough cells for tissue engineering or cell therapy. In our study the CFU-F frequency was comparable with earlier published data about culture of deciduous teeth pulp cells and those isolated from other tissues connected with tooth development. Calculating on the basis of $10^5$ cells, dental pulp contains at least 10 times more CFU-F than bone marrow (BM), but the total number of CFU-F in one digested remnant of crown dental pulp is about 12 to $20^4$, much below the total number of CFU-F that can be obtained after BM aspiration. From the clinical point of view, the low initial number of CFU-F is a disadvantage. However, DTSC can exert three times more population doublings than BM MSCs, so their proliferation potential is higher and they are naturally more primitive. Short PDT reveals a high proliferative activity of cells isolated in our experiment. This is consistent with similar findings of other authors, but much shorter than the average PDT reported by Suchaneck et al. This inconsistency could be explained by diverse culture conditions in different laboratory protocols, which could lead to isolation or expansion of different cell populations. The other possibility is that PDT could be influenced by different FBS lots containing different amounts of stimulators or inhibitors of cell proliferation. Besides fundamental stem cell biology, our data concerning the proliferative potential of DTSCs are important for cell therapy protocols. Namely, a small number of cells harvested from a primary source is a limitation for therapeutic use. We showed that, although the initial number of $0.3 \times 10^5$ to $3 \times 10^5$ cells harvested from dental pulp tissue explants was insufficient for clinical use, expansion was fast and the final number of cells after the fourth passage (calculating PDT ~ 25 h) was around $100 \times 10^6$. Also, their CFE was high, leading to the conclusion that most cells have the important proliferative potential necessary for tissue engineering strategies.

It is known that remnants of dental pulp contain extracellular matrix, odontoblasts, fibroblasts, endothelial cells, pericytes and MSCs. Among them, MSCs, endothelial cells and pericytes are migratory cells that at the same time have high proliferative potential. A heterogeneous phenotype for CD146 and CD106 antigens in multicolony culture of MSCs is a common finding and not all of the markers are specific for putative mesenchymal stem cells. CD146 is expressed in pericytes and endothelial cells in culture, and CD106 is a vascular cell adhesion molecule (VCAM) expressed in endothelial cells and also in smooth muscle cells and proliferating pericytes. DPSC and SHED were found positive for CD106 but less strongly than BM MSC. Since CD146 and CD106 molecules are expressed on endothelial cells and dental pulp stem cells easily differentiate...
into endothelial cells, we cannot exclude that a small portion of cells positive for CD106 in our cultures could be endothelial cells. Based on markers expression, the majority of cultivated cells could be pericytes. Indeed, multiple studies have recognized pericytes as MSCs. Therefore, we can conclude that using the outgrowth method to yield cells from remnants of deciduous teeth dental pulp, results in isolation of cells that do not belong to hematopoietic cell lineage but have markers indicative for pericytes that are also indicated as markers for MSCs.

The differentiation potential of harvested cells is important when considering their potential to regenerate specified tissues, like bone, cartilage and adipose tissue. We demonstrated that cells isolated in our multicollagen culture system are able to differentiate in cells that from large lipid droplets, deposit ECM with calcium salts and from cartilage like tissue that contains GAGs and collagen II. It was previously shown that SHED, obtained by enzymatic digestion of dental pulp, underwent adipogenic, osteogenic, dentinogenic and neurogenic differentiation in vitro, while chondrogenic potential and embryonic stem cell markers were demonstrated later. IDPSC obtained by the outgrowth method formed adipocytes, osteoblasts, chondrocytes, skeletal and smooth muscles as well as neurons. OCT-4 and other embryonic stem cell markers were also detected pointing to the very primitive nature of these cells. Considering all these findings, it seems that SHED and IDPSC, first claimed to have separate characteristics, do not differ from each other. The spectrum of the differentiation potential of DTSC was enlarged when these cells were found to differentiate into a pancreatic cell lineage resembling islet-like cell aggregates.

Previously, important results about the differentiation potential of DTSC multicollagen and clonal cell cultures were collected from experiments in vivo, mostly using immunocompromised mice. Thus, one quarter of SHED clones generated dentine-like tissue. Also, multicollony derived cells generated ectopic dentine-like tissue equivalent to that produced by clonal cells indicating that multicollony derived cell populations have the same differentiation capacity as clonal cells but are more convenient to use in a clinical setting.

**Conclusion**

In our experimental conditions, after 10 to 15 days of explant culture, the harvested cell population was able to expand for up to 1 month, when the cultures were stopped. Cells were positive for mesenchymal cell markers typically found on expanded stem cell populations, produced CFU-F and successfully formed a mineralized matrix, cartilage-like tissue and adipocytes, so showing multipotency. Taking together, the approach using dental pulp tissue explants yield high number of cells with MSC properties and is convenient for further investigations in vitro and work on tissue engineering protocols.

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