**Storage protocol of dental pulp cells from human exfoliated deciduous teeth**

**Introduction**

The development of cell cultures has been proposed since the 50s to evaluate microscopically normal cell events. Currently, this methodology is largely explored in researches on many knowledge areas because the obtained results are very close or equal to those observed in living organisms [1-2]. Moreover, laboratorial studies are reproducible, have excellent cost-benefit ratio, are pertinent and adequate to guide *in vivo* experiments [3-5].

In Dentistry, dental tissues for culture of primary cells as dentin, gingiva, and pulp can be obtained during routine procedures [6]. The primary cells exhibit a morphology identical to the tissues of origin, maintain the physiologic features, and can represent the natural conditions of an organism [1,7].

Recent researches demonstrate a great potential of the pulp tissue from exfoliating primary teeth because this tissue is rich in undifferentiated mesenchymal cells, which raise great scientific interest to evaluate the bioactive potential and possible clinical applications [8-11]. The easy access to pulp tissue, obtained from a non-vital organ that is normally discarded after extraction, are very attractive in research. Notwithstanding, limitations in obtaining, culturing and controlling of the cell proliferation emphasize the search for new sources, techniques, and applications [7,12-13].

Different methodologies can be employed to establish the culture of primary cells from pulp tissue. As far as we are concerned, the literature lacks a methodology specific for primary tooth. The techniques of pulp cells culture used recently are adaptations from methodologies used for obtaining pulp cells from the permanent teeth [6,12].

This study aimed to isolate the cells from the dental pulp tissue of human primary teeth, study the capacity of proliferation, characterize the cells and standardize the technique of culture and expansion to create a cell banking.

**Material and Methods**

This study was submitted and approved by the Institutional Review Board regarding the ethical aspects (CAAE 49806215.6.0000.5417). The children referred to the Pediatric Dentistry clinics for extraction of sound teeth due to orthodontic reasons were selected as potential donors for collecting the pulp tissue. Inclusion criteria were children of both genders, aged from 5 to 9 years, with good general health. The exclusion criteria comprised teeth with caries lesions and children taking any type of medicines.

After extraction, the teeth were placed into falcon tubes containing Dulbecco’s modified Eagle’s medium (DMEM) (Gibco, Thermo Fisher Scientific in Waltham, MA, USA) supplemented with 20% Fetal Bovine Serum (FBS) (Gibco, Thermo Fisher Scientific in Waltham, MA, USA). The pulp tissue was extracted through the apical foramen with the aid of sterile curettes and endodontic files. The collected pulp tissue was placed into a Petri plate (100 mm in diameter x10 mm height), cut at small pieces with the aid of 15c scalpel blade (Figure 1A), and then, immersed in the supplemented medium containing DMEM 20% FBS supplemented with 600 µL of penicillin (Gibco, Thermo Fisher Scientific in Waltham, MA, USA), 300µL of gentamicin (Gibco, Thermo Fisher Scientific in Waltham, MA, USA), and 100 µL of amphotericin B (Gibco, Thermo Fisher Scientific in Waltham, MA, USA) (Figure1B, 1C and 1D), and incubated at 37oC and 5% CO2 for 40 minutes (Figure 1E). Elapsed that period, all content of the plate was collected and dispensed into a falcon tube, centrifuged at 1200 rpm for 5 minutes at 200C (Figure 1F, 1G and 1H). The cell pellet was resuspended in new culture medium (DMEM 20% FBS), and stored in 25cm2 cell culture flasks incubated at 37oC and 5% CO2 (Figure 1I and 1J).

The cell cultures were maintained with periodic changes of the culture medium (DMEM 20% FBS) at every 48 hours until the cells reached confluence (Figure 1K), defining the passage zero (P0). Next, the cells were trypsinized, centrifuged, the pellet resuspended and transferred to 75 cm² culture flasks. After confluence, the procedure was repeated to obtain the subsequent passages.

The isolated pulp cells were characterized as fibroblasts by immunofluorescence technique. The cells were marked with primary monoclonal antibody and anti-human fibroblast surface protein (Anti-FSP) (ABCAM, Cambridge, United Kingdom) at final concentration of 2 µg/mL. Pulp cells from the 5th passage were seeded at 104 cells/well on 8-well chamber slide, left overnight to adhere on the well. Then, the cells were fixed in 4% paraformaldehyde for 15 minutes and incubated with PBS supplemented with 3% bovine serum albumin for 30 minutes at environment temperature. After that, the cells were marked with the primary antibody (1:1000), left overnight; followed by the secondary antibody (1:500) (ABCAM, Cambridge, United Kingdom). The slides were prepared for the analysis through laser confocal microscopy (Leica TCS SPE, Mannheim, Germany).

**Results**

The pulp tissue obtained from the extracted primary teeth allowed the primary cell culture. After 72 hours, the fragments of pulp tissue adhered on the flask surface (Figure 2A). At seven days of cell culture, the pulp cells started loosening (Figure 2B) and continued that way at 14 days (Figure 2C). After 21 days, the cell confluence was reached and the microscopic analysis revealed the normal morphology of the cells compatible with fibroblasts (Figure 2D). During all this study period, neither the medium nor the cell culture showed contamination. Also, marked cell death was not present.

The microscopic analysis of the slides marked by the immunofluorescence reaction with the antibody specific for fibroblast demonstrated positive immunostaining. The immunofluorescent staining of the selected antibody revealed the cell nuclei (Figure 3A) and cytoplasm (Figure 3B) with different colors, evidencing the cell morphology of fibroblasts (Figure 3C).

**Discussion**

Researches on Dentistry field has investigated the potential of cell culture and tissue engineering, so this methodology can be used for investigating material’s cytotoxicity, cell viability, and proliferation after different stimuli, guiding further clinical trials with lower cost, shorter time and small number of individuals [2,4-5,14-15]. However, the literature reports lack of standard techniques to obtain the cells from specific tissues [2,14]. Thus, it is important to investigate and standardize the cell culture techniques for different tissue types by characterizing the cells, using specific reagents, and correct storing the cells for further studies [14,16-18].

Currently, the cell culture technique has been used frequently in dental researches [4-5,15]. The creation of cell banks either in study centers or for commercial purposes, resulted in great advances in the standardization of the technique for obtaining, culturing, and expanding many cell types, aiming at mutual collaboration between researchers and companies to achieve common guidelines on techniques and laboratorial supplies [17,19].

The guidelines on primary culture from tissue explants, mainly pulp and gingival tissue, have been the technique most adopted in dental researches [2,5-6]. The pulp tissue of primary teeth has some advantages, such as the cell pluripotency and explants easily obtained from the teeth extracted in pediatric dentistry clinics [12-13,17]. The studies of SIPERT et al. [5] and TORIUMI et al. [6] evidenced the need for standardizing the techniques of cell culture from the selection of the donor teeth or areas for obtaining the tissue explants to avoid culture contamination and accelerate the expansion of the cultured cells.

After obtaining the cells from primary culture, especially those from the pulp tissue of primary teeth, the characterization (i.e. determination of the cell type) is mandatory. For that purpose, many techniques may be employed: PCR, immunophenotyping, and fluorescence. The immunofluorescence through specific antigen-antibody systems is the most used technique at many research centers to determine cell types [2,12,17,20-21].

Although some authors stated that the pulp tissue explant is an atraumatic procedure relatively easy to obtain, the creation of cell banks is a viable option allowing the study of specific lineages, characterization and expansion through standard culture techniques, reducing the waste of laboratorial supplies, and eliminating the clinical step to obtaining those explants [17,19,22].

Considering these advantages and the increasing use of cell cultures in researches and medical therapies, cell banks have gained space and importance. Accordingly, cell banking can be supported by companies aiming at the correct storage and manufacturing of the cells for further commercialization of the lineages. Cell banking contributes to obtaining different cell lineages and directly collaborates with the standardization of the specific techniques and supplies to be used and optimizes laboratorial studies with cell culture [2,23-24].

**Conclusion**

The creation of cell banking using dental pulp cells from human primary teeth enables the easy application of cells in laboratorial studies, reducing the cost and time for obtaining the samples, avoid the involvement of new subjects and allow a fast reproducibility of the researches.

**Funding**

This study received financial support from the Sao Paulo Research Foundation (FAPESP) (grants #2013/18886-5).

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**FIGURE LEGENDS**

**Figure 1 –** Flowchart of the cell culture and isolation technique.

**Figure 2 –** (A) Fragment of pulp tissue at initial stage, (B) fragment of pulp tissue starting the cell loosening at 7 days, (C) fragment of pulp tissue at advanced stage of cell proliferation at 14 days and (D) cell confluence after 21 days.

**Figure 3 –** (A)Morphology of the nuclei stained by immunofluorescence, (B) morphology of the cell membrane and cytoplasm stained by immunofluorescence, (C) characterization of the fibroblast immunostained by the specific immunofluorescent antibody.