Cytotoxicity of solutions recommended for the storage of avulsed teeth in cultures with periodontal ligament cells

Galia Cruz-Durán¹, Raúl Ortiz-Daza Raúl¹, Jacinto Armando Díaz-Acevedo¹, Benjamín Sánchez-Trocino¹, Ma. Concepción Arenas-Arrocena² and René García-Contreras²

¹Oral and Maxillofacial Surgery Area; ²Nanostructures and Biomaterials Area, National School of Higher Education, León Unit, Universidad Nacional Autónoma de México, León, Guanajuato, Mexico

Abstract

Introduction: The medium for avulsed teeth storage until their reimplantation is key to the preservation of human periodontal ligament fibroblasts (HPLF). Objective: Our purpose was to compare the cytotoxic effect of milk and isotonic solution, used for the storage of avulsed teeth, on the preservation of HPLF. Method: A subculture of periodontal ligament fibroblasts was carried out with a density of 1:2 (3 x 10⁵ cells/mL) and was incubated for 48 hours. The cells were divided in two groups, which were placed either in milk or isotonic solution for 24 hours at 5% CO₂, 37 ºC and 95% humidity. The number of viable cells was determined with a colorimetric fast assay by the reduction of MTT and mitochondrial activity. Data were processed with the Shapiro-Wilk normality test, Student’s t-test and paired Student’s t-test (with significance set at 0.05). Results: The cells exposed to milk for 24 hours showed statistically significant cytotoxicity at concentrations of 0.09, 0.39, 0.78, 1.56, 3.125, 6.25 and 50%. HPLFs exposed to isotonic solution showed no significant reduction in the number of cells at concentrations of 25 and 50%. Conclusion: Isotonic solution appears to be better for HPLF 24-hour storage in comparison with whole milk.


Introduction

According to the World Health Organization classification of dental trauma, dental avulsion is the complete displacement of a tooth from its socket in alveolar bone.¹ An incidence of 16 % has been reported among all permanent teeth dental traumas and 7 to 21 % for deciduous teeth. Avulsion is a severe dental injury and its prognosis varies depending on the immediate procedure after the accident. Replanting the tooth in the corresponding place is one of the treatments of choice, even when carrying this out immediately is not always possible.²

Different studies concur that tooth replanting has a better prognosis the less time the tooth remains outside the alveolus and if the periodontal ligament is fully preserved. The prognosis considerably improves (41 %) when the teeth still have an open apex, which increases the incidence for pulp revascularization.³ Periodontal ligament cells have constant blood supply, pH of 7.2, 280 to 300 mOsm osmolality, so that when the tooth has been displaced from its alveolus, these cells start dying in less than 15 minutes, and in less than 1-2 hour, a sufficient number of periodontal ligament cells will have died, which drastically reduces replantation endurance and success. The result of ligament necrosis is observed with tooth radicular resorption and ankylosis in the long term.

Replantation survival depends on preserving the cells that compose the periodontal ligament unharmed. For this, it is crucial to maintain the tooth immerse in a fluid medium; milk, water or saliva have been reported to be optimal for preservation; storage in these fluids can increase the waiting time to up to 24 hours...
after avulsion, for subsequent manipulation and replantation. When the avulsed tooth is replanted after 15 minutes, periodontal ligament damaged cells cause partial resorption. Furthermore, replantation with a delay longer than 30 minutes can cause irreversible cell damage, replantation with a delay longer than 60 minutes in dry conditions can cause periodontal ligament (PL) necrosis, leading to extended root resorption. The best prognoses are obtained when extra-alveolar time does not exceed 5 minutes.

The storage medium should be able to preserve cell vitality and adherence capacity, as well as to be readily available at the moment of avulsion. Both physiological osmolality and pH are crucial to preserve PL cells vitality. Cell growth occurs within a range of 230 to 400 mOsm; however, optimal growth ranges from 290 to 330 mOsm; pH ranges from 6.6 to 7.8 and the ideal range is 7.2 to 7.4.

Milk has been recommended as a medium to store an avulsed tooth because it has an osmolality (220 mOsm) that is compatible with that of periodontal ligament cells, in addition to being widely available; however it lacks the necessary metabolites and glucose for normal cell physiology. Milk keeps periodontal ligament cells alive for 1 to 3 hours. Cells that had been dried for 20 to 30 minutes and then placed in milk for 45 minutes have been shown to have much less vitality than those that were dried for only 10 minutes. PL cells that were dried with air for 60 minutes and then placed in milk showed an almost inexistent vitality, which demonstrated milk capability to maintain osmotic pressure for PL cells, but showed no capability to reconstruct cell metabolism or to restore its vitality. Only cold milk can preserve PL cells capability to proliferate.

The isotonic solution used was a rehydrating, energizing commonly used isotonic beverage, which contains different components that might be able to preserve human periodontal ligament fibroblasts (HPLF): sodium chloride, potassium chloride, calcium chloride, magnesium chloride, sodium lactate and glucose; however, its 4.5 pH might represent an important disadvantage in comparison with other solutions such as milk.

The purpose of this study was to compare the cytotoxic effect of milk and an isotonic solution, both recommended for avulsed teeth storage, in the preservation of HPLF, by means of cell-viability tests with a rapid colorimetric assay by MTT and mitochondrial activity reduction.

Method

Primary cell culture

The periodontal ligament cells (HPLF) were obtained by means of biopsy of the periodontal tissue from a procedure of caries or infection-free erupted third molars dental extraction in two 18 and 25-year-old patients, after informed consent was signed and authorization was granted by the bioethics committee of the National School of Higher Education, Leon Unit, Universidad Nacional Autónoma de México, León, Guanajuato, Mexico.

The extracted teeth with the obtained tissue were suspended in Alpha-MEM medium (Life Technologies, Gibco, Carlsbad, Ca, USA), with 20 % fetal bovine serum added (FBS, Life Technologies, Gibco, Carlsbad, Ca, USA), as well as 100,000 IU/mL of penicillin G and 100 µg/mL streptomycin sulfate (Life Technologies, Gibco, Carlsbad, Ca, USA).

The tissues were sectioned into small portions with a number 15 scalpel blade. The explants were placed in 100-mm culture boxes and incubated at 37 °C with an atmosphere of 5 % CO₂ for 2 weeks for exponential growth, with culture medium changes first at seventh day, and subsequently every other day. Cell growth was obtained as a primary culture with a population doubling level (PDL) of zero. When cellular density was 80 %, sub-cultures were carried out by detaching the cells from the culture plate with 1 mL of 0.25 % trypsin-0.025 % EDTA-2Na (Life Technologies, Gibco, Carlsbad, Ca, USA) in PBS(-) and incubating them for 5 minutes at 37 °C; detachment was verified by observing the culture box circulating cells under the microscope. The subcultures were carried out at a 1:3 concentration and the culture medium (DMEM + 10% FBS) was replaced every other day. Periodontal ligament fibroblasts have a lifespan of 30 PDL in vitro (accumulated number).

Cytotoxicity assay

Subcultures were carried out at a density of 1:2 (3 x 10⁵ cells/mL) in 96-well plates (Ultra Cruz®, Santa Cruz Biotechnology, Hamburg, Germany) and were incubated for 48 hours with 5 % CO₂ at 37 °C and 95 % humidity. The entire culture medium was removed and the cells were placed in new, fresh culture medium. The whole milk (Pasteurizadora León, León, Guanajuato, Mexico) and the isotonic solution
(Electrolit, Laboratorios Pisa, Guadalajara, Jalisco, Mexico) were inoculated at different concentrations, from 0 to 50%. The cells were incubated for 24 hours at the same temperature and under the same conditions. Cell viability was determined with a rapid colorimetry assay by MTT® formazan reduction (Sigma-Aldrich, Toluca, Mexico).

Dissolved in 0.2 mg/mL of DMEM culture medium, the cells were incubated for 4 hours at 37 °C and then the culture medium was removed until crystals were dissolved with dimethyl sulfoxide (DMOS, Karal, León, Mexico). Dose-response curve cell viability was determined and analyzed in a microplate spectrophotometer at 570 nm. The assays were performed in duplicate, 16 for each solution.

### Statistical analysis

Average, standard deviation, percentage and mean cytotoxicity concentration (CC$_{50}$) were calculated based on the dose-response curve. The data underwent Shapiro-Wilk, Student’s t and paired Student’s t normality tests. Statistical significance was set at $p < 0.05$, with a reliability coefficient of 95%.

### Results

HPLF behavior was observed in both solutions, with cytotoxicity being assessed at different concentrations. The cells exposed to milk for 24 hours showed statistically significant cytotoxicity, with $p < 0.01$ at 0.09%, 0.39%, 0.78%, 1.56%, 3.125%, 6.25% and 50% concentrations, with CC$_{50}$ = 46.82% standing out.

The HPLF group exposed to the isotonic solution showed no significant differences, with $p > 0.05$ at 50% and 25% concentrations; however, these concentrations showed no cell-viability decrease below 80%, showing cytostable effects on the culture. When compared, statistically significant differences were found for HLPF: at 12.5 and 25% concentrations, a p-value < 0.05 was recorded, and in the rest, $p < 0.01$ (Table 1 and Fig. 1).

### Discussion

An ideal storage medium has been described to have to possess biological properties to preserve cell vitality and adherence capacity, as well as to be available at the moment of avulsion. Both physiological osmolality and pH are crucial to preserve PL cells viability. Cell growth occurs within a range of 230 to 400 mOsm; however, it is optimal within a range of 290 to 330 mOsm; pH ranges from 6.6 to 7.8, with the ideal range being 7.2 to 7.4.

Different solutions that might show favorable characteristics for HLPF storage have been investigated; those that have shown better capability for cell-viability preservation are Custodiol®, a fluid for organ transportation, and Hank balanced saline solution (HBSS). However, owing to their low availability and difficulty to find them, other solutions that might be good options for avulsed teeth storage have been investigated. According to Krasner et al., Sigalas et al., and Özcan, milk is an adequate solution for periodontal ligament fibroblasts (HPLF) storage owing to its osmolality and pH, in addition to its ready availability in the place of injury. Other solutions have been tested, such as Gatorade®, since its acidic pH showed high capability to produce cell apoptosis, and contact lens solution, which showed no significant differences with HBSS and milk; green tea extract, which is efficacious to preserve periodontal ligament cell viability for 24 hours and that also showed anti-inflammatory, antioxidant and anticarcinogenic effects; coconut water demonstrated to be better for PL preservation in comparison with HBSS and milk.

The used isotonic solution has a pH of 4.5 and osmolality of 116 mOsm. When the capacity to produce cytotoxicity at 24 hours at 0 to 50% concentrations was assessed, no significant difference was found in the reduction of cell viability. However, it is necessary...
for other investigations to be carried out, since to this moment no other author has reported the potential application of this solution. As a reference, physiological saline has been tested, by means of which, according to Caglar, similar results to those of milk and HBSS were observed at 30 and 45 minutes.

In the present study, when the isotonic solution was compared with milk, the former was found to be significantly better in cell preservation at any concentration; significant differences were observed with regard to the control group only at 50 % and 25 % concentrations, probably owing to the electrolytes present in the isotonic solution, which are necessary for HPLF preservation; however, this is controversial, since its osmolality and pH are lower than those recommended in the literature. It is important pointing out that the isotonic solution is of low cost and is available in various stores and accessible to the entire population.

Since the results regarding milk and the isotonic solution are not consistent with those of previous investigations, or with those expected based on both solutions’ biological characteristics, it is necessary for investigations aimed at testing the effects of both these fluids to be conducted with variability not only in concentrations but also in HPLF storage time before suffering irreversible cell death or expression of cytokines that may interfere in periodontal cell vitality, which is a fundamental aspect after dental avulsion.

**Conclusion**

Milk has been considered to be the ideal solution for the storage of avulsed teeth, owing to its biological characteristics and availability in the market. In this study, an isotonic solution that is highly available in Mexico was shown to apparently offer better results with regard to HPLF 24-hour storage.

**References**


