Cytotoxicity of Endodontic Irrigants Containing Calcium Hydroxide and Sodium Lauryl Sulphate on Fibroblasts Derived From Mouse L929 Cell Line

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The aim of this study was to evaluate the cytotoxicity of root canal irrigating solutions containing calcium hydroxide and sodium lauryl sulphate on fibroblasts derived from L929 cell line. Saturated calcium hydroxide aqueous solution (CH), sodium lauryl sulphate (SLS) and SLS associated with calcium hydroxide (HCT20) were diluted with sterile distilled water at 50%, 20%, 10% and 5% concentrations. Minimum essential medium (MEM) served as the control group. The cytotoxicity of the solutions was evaluated on L929 mouse fibroblast cell line, at 4 and 24 h of contact time by the 51Cr radiotracer method. Data were compared and statistical inferences were made with the chi-square test. In all analysis, significance level was set at 5%. CH and HCT20 showed toxicity at 50% concentration, while at concentrations lower than 50% these solutions showed cell tolerance. SLS was cytotoxic at all concentrations. In conclusion, the association of calcium hydroxide and SLS (HCT20) combines the beneficial properties of these solutions and was not harmful to the fibroblast cell line, seeming to be a suitable endodontic irrigating solution.

Key Words: cytotoxicity, calcium hydroxide, sodium lauryl sulphate, endodontic irrigants.

INTRODUCTION

Dental pulp and periapical tissues are commonly affected by a widely varied microbial flora, including Gram-positive and Gram-negative microorganisms and their metabolite byproducts or toxins. Even after microbial death, components of the cellular wall, such as lipopolysaccharide (LPS), persist for prolonged periods and can cause biological reactions resulting in chronic inflammation (1).

Endodontic therapy requires use of irrigating solutions to eliminate microorganisms, remove debris and neutralize organic compounds. Because of the risk of leakage through the apical foramen, irrigants must be biocompatible and non-irritant to the periapical tissues (2,3). Sodium hypochlorite (NaOCl) is the most commonly used irrigating solution because it has outstanding antibacterial action, bleaches the dentin and dissolves organic material (4-6). It must be used with adequate irrigation and instrumentation technique, since may pass through apical foramen causing periapical damage and disturbing repair inhibiting adherence capacity of macrophages. As drawbacks, the high superficial tension of this chemical irrigant avoids its penetration into the canal system irregularities and its use increases the dentin hydraulic conductivity (2,6-9).

Calcium hydroxide has antimicrobial action, neutralizes microbial organic debris, induces repair with mineralized tissue deposition and hydrolyzes the lipid moiety of LPS (1,4,10-12). It has low solubility in water and high superficial tension, which is lowered with the association to sodium lauryl sulphate (SLS) (13). The toxicity of calcium hydroxide solutions has not yet been clearly elucidated.

The aim of this study was to evaluate in vitro the cytotoxicity of saturated calcium hydroxide solutions, sodium lauryl sulphate and HCT20 (sodium lauryl sulphate associated with calcium hydroxide solution) at 5%, 10%, 20% and 50% concentrations on fibroblasts derived from L929 cell line.

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MATERIAL AND METHODS

Three potential irrigating solutions were evaluated. Saturated calcium hydroxide aqueous solution composed of 0.2 g of calcium hydroxide in 100 mL distilled water, pH 12.2 (Calcium Hydroxide, Sigma, St. Louis, MO, USA), designated as CH, 0.125% sodium lauryl sulphate in distilled water, pH 7.8 (Tergensol-Inodon; Porto Alegre, RS, Brazil), designated as SLS, and 0.16 g calcium hydroxide plus 0.025% sodium lauryl sulphate in 100 mL distilled water, pH 11.0, named HCT20. The solutions were diluted with sterile distilled water at 50%, 20%, 10% and 5% concentrations. Minimum essential medium (MEM) served as a control group.

Fibroblasts derived from mouse L929 cell line (ATCC L929) were used due to their suitability for cytotoxicity evaluation. The culture medium was minimum MEM with Eagle’s salts (Flow Laboratories, McLean, VA, USA) supplemented with 10% (v/v) fetal bovine serum, 100 IU/mL penicillin, μg/mL of streptomycin, 2 mM/L glutamine and 2.2 mg/mL of sodium bicarbonate. Five- to 7-day cultures were used for the experiments. The culture medium was changed every other day, and the day before the experiments. The $^{51}$Cr was supplied as sodium chromate in sterile isotonic saline (Dupont NEN, Wilmington, DE, USA). Cell monolayer cultures were labelled by incubation with 1 μCi per $10^5$ cells, 20 to 24 h before the experiments (14). The labelled cells were harvested with 0.125% trypsin and washed 3 times in Ca$^{2+}$ free and Mg$^{2+}$ free phosphate buffered saline (PBS). The cells were resuspended in MEM at a concentration of $4 \times 10^4$ cells/mL.

One milliliter of the test solution was mixed with 1 mL of labeled cell suspension in multiwell cell culture plates (Costar, Cambridge, MA, USA), in such a way that the test concentration of each solution evaluated was 5%, 10%, 20% and 50%. Ten samples of each solution were used for each experimental period. In the control wells, the irrigating solution was replaced by MEM. During the dispensing of the cells, cell suspension was randomly distributed into 6 test tubes and used as reference samples. The experimental and control plates were incubated for 4 and 24 h at 37°C in an atmosphere of 5% CO$_2$ in air and 100% humidity.

At the end of the exposure period, the pH of the solutions was obtained by 2 readings and 1 mL of medium from each well was transferred to test tubes and centrifuged at 500 g for 8 min. The supernatant (0.5 mL) was counted for 1 min in a gamma-particle counter (Gamma 5500; Beckman Instruments, Wakefield, MA, USA). The percentage of $^{51}$Cr released from the experimental and control samples was calculated according the equation: $^{51}$Cr release in percentage = $2 \times (T-b) \times 100$, $R-b$

where T = chromium released in test samples, b = background radiation and R = reference sample.

Sample frequencies were compared and statistical inferences were made using the chi-square test at 5% significance level.

RESULTS

All solutions showed some level of toxicity. CH and HCT20 at 50% concentration presented significant chromium release, being 2-3 times greater than that of the control group. At 20%, 10% and 5% concentrations, however, these solutions had similar behavior to the control group. SLS showed high toxicity at all evaluated concentrations. The different concentrations of the solutions did not show significant pH variation. Data are graphically presented in Figures 1 and 2.

Figure 1. Effects of irrigating solutions on L929 mouse fibroblast cell line in vitro. Release of radiochromium (%) after 4 h of exposure. Values are expressed as Mean ± SD (n=10).

Figure 2. pH of the irrigation solutions.
DISCUSSION

CH and HCT20 at concentrations below 50% presented similar results to those of the control group on L929 fibroblasts. These data reveal good biological behavior since MEM is not harmful to cell viability and therefore it is possible that the extrusion of those solutions do not cause damage to the periapical tissues. The pH of CH and HCT20 at 5%, 10% and 20% concentrations is not viable for the survival of various microorganisms of endodontic infections (Fig. 2) (12,13,15).

At the highest concentration (50%), Ca(OH)$_2$ and HCT20 were cytotoxic, which may be related to the superficial caustic action and the enhancement of tissue solubility inherent to calcium hydroxide (4,16). Some authors (5,17) have previously reported that calcium hydroxide can dissolve dental pulp remnants and affect the dental pulp stump, but its reparative action requires a high pH to occur (12,16).

SLS properties, such as low superficial tension and lubricant ability, certainly help the flushing action and cleaning of the root canal system (18). Though, it does not have antimicrobial action (13) and has some superficial caustic action and the enhancement of tissue solubility inherent to calcium hydroxide (4,16). Some authors (5,17) have previously reported that calcium hydroxide can dissolve dental pulp remnants and affect the dental pulp stump, but its reparative action requires a high pH to occur (12,16).

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SLS can increase LPS disaggregation (19) and inhibit bacterial coaggregation (20). In combination with calcium hydroxide, it may be a viable irrigating solution, since calcium hydroxide has antimicrobial action, detoxifies the bacterial LPS and is tolerated by the tissues (1,10,11). The association of these substances does not seem to have drawbacks to endodontic therapy.

The evaluation of irrigating solutions using $^{51}$Cr radiotracer assay is an accurate and well known method because it allows the contact liquid/cells (14). In vivo, differences may be understood as suitable and support the use of the irrigating solution because the contact time during an endodontic treatment is lesser. In this study the evaluation performed done for 4 h to warrant accurate readings.

Apical extrusion of debris and solutions during root canal instrumentation is an important aspect to be considered resulting in inflammation or flare-up (2,3). Therefore, it is recommended that endodontic irritants are biocompatible, present antimicrobial activity and stimulate or do not jeopardize the periapical repair (13).

Comparison of other irrigants and calcium hydroxide solutions, with or without SLS, indicates no better results considering the combination of effective antisepitic action and low toxicity (9,10). Due to its biological properties, calcium hydroxide solutions can extrude to the periapical region without causing tissue damage. In conclusion, the association of calcium hydroxide and SLS (HCT20) combines the beneficial properties of these solutions and was not harmful to the fibroblast cell line, seeming to be a suitable endodontic irrigating solution.

RESUMO

O objetivo desta pesquisa foi avaliar a citotoxicidade de soluções irrigadoras de canais radiculares contendo hidróxido de cálcio e laurel sulfato de sódio em linhagem de fibroblastos L929. Solução aquosa saturada de hidróxido de cálcio, laurel sulfato de sódio e HCT20 (lauril sulfato de sódio e hidróxido de cálcio) foram diluídos em água destilada em concentrações de 50%, 20%, 10% e 5%. O grupo controle foi representado por meio de cultura de células (MEM – minimum essential medium). A citotoxicidade das soluções sobre os fibroblastos foi avaliada em 4 e 24 h de contato, pelo método do cromo radioativo. Os resultados foram analisados estatisticamente pelo teste do qui-quadrado. Em todas as análises, o intervalo de confiança referente às médias entre os grupos foi estabelecido em 95%. As soluções saturadas de hidróxido de cálcio e o HCT20 apresentaram toxicidade nas concentrações de 50%. O laurel sulfato de sódio foi tóxico em todas as concentrações. As soluções de hidróxido de cálcio em concentrações menores que 50% apresentaram tolerância celular, assim como combinadas ao laurel sulfato de sódio. Tal comportamento não foi observado na solução pura de laurel sulfato de sódio em todas as concentrações.

REFERENCES