Original Article

Cytotoxicity evaluation of five different dual-cured resin cements used for fiber posts cementation

M Dioguardi¹, D Perrone¹, G Troiano¹, L Laino¹, F Ardito¹, F Lauritano², M Cicciù³, L Lo Muzio¹

¹Department of Clinical and Experimental Medicine, University of Foggia, FO, Italy; ²Department of Odonto-Stomatology, University of Messina University of Messina School of Dentistry, ME, IT; ³Department of Human Pathology, School of Dentistry University of Messina School of Dentistry, ME, IT

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Abstract: Custom-cast posts and cores are usually used to treat endodontically treated teeth. However, several researches have underlined how these devices may be a much higher elastic modulus than the supporting dentine and the difference in the modulus could lead to stress concentrating in the cement lute, leading to failure. The role of the cement seems to play a fundamental role in order to transfer the strength during the chewing phases. Aim of this research is to record the rate of cytotoxicity of five different dual-cured resin cements used for fiber posts cementation. We tested the cytotoxicity of this five materials on MG63 osteoblast-like cells through two different methods: MTT ([3-4,5-dimethylthiazol-2-yl]-2,5-diphenyltetrazolium bromide succinate) assay which tests for mitochondrial enzyme activity and xCELLigence® system.

Keywords: Fiber post cementation cytotoxicity

Introduction

The aim of the endodontic treatment is represented by the morpho-functional restoration of the tooth, through the use of appropriate restorative materials. Before proceeding to the anatomical reconstruction of the tooth, is important to have obtained cleaning and filling of the endodontic system by shaping of channels, to reach the killing of bacteria responsible for any periapical pathology [1]. The endodontic space under therapy should not be considered only as a system of channels to clean, shape and fill, but as a space that interfaces with the adjacent tissues [2]. Endodontic materials should remain inside the root canal space and should not extrude in periapical tissues, but in reality the clinic shows us how these can be in contact for a long period with the periodontal tissues [3, 4]. For this reason it is important to study the direct cytotoxicity of these materials. This problem is also present during the cementation of endodontic fiber posts that may be in close contact with the periapical and lateral periodontium [5]. The purpose of this study was to evaluate the rate of cytotoxicity of five different dual-cured resin cements used for fiber posts cementation. We tested the cytotoxicity of this five materials on MG63 osteoblast-like cells through two different methods: MTT ([3-4, 5-dimethylthiazol-2-yl]-2,5-diphenyltetrazolium bromide succinate) assay which tests for mitochondrial enzyme activity and xCELLigence® system for evaluating the kinetics of cell adhesion [6, 7].

Materials and methods

Cell cultures

MG63 osteoblast-like cells, originally isolated from a human osteosarcoma, were used. MG63 were cultured in 200 ml culture flasks each containing a 15 ml mixture of 90% Ham’s F12/Dulbecco’s modified Eagle’s medium (DMEM-F12) complemented with 2 mM glutamine, 10% foetal bovine serum, 200 U mL-1 penicillin and 200 μg mL-1 streptomycin and then placed in standard cell culture conditions at 37°C in a 5% CO₂ humidified air environment until confluence had been achieved. Cells were trypsinized and then seeded, for the following assays, on...
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Preparation of test cements

The following materials were tested: Biscem, C & B, Duolink (Bisco Inc., Schaumburg, USA ®); Virage Plus and Virage dual (Sweden & Martina, Padova, IT ®). The materials were prepared according to the manufacturer's instructions under aseptic conditions. Materials were inserted into a round metal appliance designed for production of discs measuring 5 mm wide and 3 mm high. Materials were allowed to set for 24 h in a humid atmosphere and aseptic conditions. For cyto-compatibility assays, samples were sterilized by incubation for 2 h in 200 U ml⁻¹ penicillin, 200 μg ml⁻¹ streptomycin, 250 μg ml⁻¹ fungizin and 50 mg L⁻¹ gentamycin. Twelve samples of each material to be tested were disinfected by immersion in ethanol and left to dry in the flow cabinet [8-9]. These were added to the culture wells using sterile forceps. All the samples were subsequently incubated at 37°C with 5% CO₂ in DMEM following a 1 h ultraviolet exposure [8-10]. The elutes of materials were prepared separately and were incubated in DMEM-F12 at 37°C in a 5% CO₂ for 1, 2, 3 and 7 days [11]. Confluent MG63 were counted using a hematocytometer (4×10⁴ cell) seeded onto 96 well plates and then incubated with the supernatant of each cements and for each time-points. Cell viability was evaluated by MTT assay after 24 h. Negative controls were performed in empty (not cell-containing) wells and positive controls were performed in wells with unexposed MG63.

MTT assay

The cytotoxic effects of each materials were assessed by measuring the reduction in cell metabolic activity using the colorimetric assay as initially described [12]. After 1, 2, 3 and 7 days of growth of MG63 cells on the cement disk surfaces, the cells were analysed. Unexposed control cultures were maintained under the same conditions. Briefly, the cells were washed once with PBS, and MTT solution (0.5 mg/mL) was added to each well. After incubation at 37°C under 5% CO₂ for 3 h, the blue formazan formed by the reduction of the MTT was dissolved in 150 μl of IPA/HCl (0.4 N) by vigourously pipetting. The amount of formazan was determined by measuring absorbance at 570 and 650 nm for background using a microplate reader (model Multiscan Go UV-Vis, Thermo Scientific).

For each point, three wells were analysed, and three independent experiments were performed, so for every materials we tested 9 samples. Results were expressed as a percentage of viability, namely the ratio of absorbance of exposed to unexposed cells, according to the following equation:

Viability (%) = \( \frac{(\text{Ast570} - \text{Ast650})}{(\text{Asc570} - \text{Asc650})} \times 100 \)

Where Ast570 is the absorbance (at 570 nm) of the extract from incubated cells with tested material, Asc570 the absorbance (at 570 nm) of the extract from control cells and Asc650/Ast650 the absorbance at the reference wavelength for control and treated cells, respectively. Cytotoxicity responses were rated as severe (<30%), moderate (30-60%), slight (60-90%) or non-cytotoxic (>90%) based on the activity relative to values obtained for the controls [13].

xCELLigence

The RTCA instrument is composed of a RTCA impedance analyzer, a computer with RTCA software for controlling the system operation, a 48-well electronic microtiter plate (E-Plate) and the RTCA station, which accommodates the E-Plate and it is placed inside the tissue culture incubator. The presence of cell culture medium or buffer and the application of a low voltage create an electric field between the electrodes, which can be impeded by cell presence. The electronic readout of cell-sensor impedance is displayed in real-time as CI, a value directly influenced by cell attachment, spreading, and/ or cell proliferation. The CI value at each time point is defined as Rn-Rb/Rb, where Rn is the cell-electrode impedance of the well with the cells and Rb is the background impedance of the well with only medium. Fifty microliters of cell culture medium was added to each well for the impedance background measurement. After adding 4×10⁴ cells and extract, the final volume was 200 μL. The E-Plates were incubated at 37°C with 5% CO₂ and monitored on the RTCA system at 2-minute time intervals for up to 72 hours with or without treatment. To facilitate the statistical evaluation of the results, we performed at least two repeats of each experimental condition as recommended in the technical manual of xCELLigence [14, 15].
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Statistical analysis

Data were analyzed using GraphPad Prism software version 6.00 for Windows (GraphPad Prism Software, San Diego, CA, USA). Average values were expressed as mean ± s.d. Statistical significance between different groups was determined by repeated-measures ANOVA test and Tukey test. A p value <0.05 was accepted as statistically significant.

Results

The cell viability of human osteosarcoma cells in contact with the dishes was evaluated by means of MTT assay (Figure 1A).

Discussions

Cytotoxicity assays are the initial screening tests used to evaluate the biocompatibility of materials [17]. The combination of these tests with: genotoxicity, mutagenicity, carcinogeticity and microbial analysis allow obtaining parameters that characterize biocompatibility [18]. The aim of this study was to evaluate the cytotoxicity activity of five dual-cured sealers used for fiber posts cementation through two different tests: MTT and Xcelligence® assays. We evaluated the rate of cytotoxicity of Dual Virage, Virage Plus, Biscem, C & B and Duolink on MG63 osteoblast-like cells both on dry sam-
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Figure 2. Cell Index of control group (A) and with MG63 exposed to Biscem (B), c & b (C), Duolink (D) Virage Plus (E) and Virage Dual (F) eluates at different timepoint 24 h (-), 48 h (-), 72 h (-), 7 days (-) Results obtained with xCelligence analysis confirm the biocompatibility rates of the above-mentioned materials.
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amples of cements (set group), that on the eluates derived from them (fresh group). At 24 hours, results from set group revealed a low rates of cytotoxicity for all sealers except Duolink. However, this rate tended quickly to decrease and, after 48 h all sealers results highly biocompatibility (Figure 2). Furthermore, the cytotoxicity remained low even after a week of contact with cells. Trials on eluates, with MTT and Xcelligence®, revealed a very low rate of cytotoxicity for Virage and Virage Plus dual, whereas C & B, Biscem and Duolink shown to be more cytotoxic. Biscem, Duolink rates of cytotoxicity could be due to the presence of TEGDMA as reported in previously studies [19, 20]. Results appear to be steady after a week of observation.

Conclusions

Virage Dual and Virage Plus revealed very low cytotoxicity, but further studies needs to understand effects of these cements on different types of cells culture.

Disclosure of conflict of interest

None.

Address correspondence to: Dr. Marco Cicciù, Department of Human Pathology, Dental School, Messina University, Via Consolare Valeria, Messina 98100, Italy. Tel: 0039-090 221 6911; E-mail: acmarco@yahoo.it

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