In Vitro Biocompatibility and Oxidative Stress Profiles of Different Hydraulic Calcium Silicate Cements

Ashraf A. Eid, BDS, MSc, ‡ Johnny L. Gosier, DMD, ‡ Carolyn M. Primus, PhD, § Barry D. Hammond, DMD, ‡ Lisiane F. Susin, DDS, ‡ David H. Pasbley, DMD, PhD, || and Franklin R. Tay, BDSc (Hons), PhD||

Abstract

Introduction: MTA Plus is a new calcium silicate cement with unknown cytotoxicity characteristics. The objectives of this study were to examine the effect of MTA Plus on the viability, apoptosis/necrosis profile, and oxidative stress levels of rat odontoblast-like cells.

Methods: MDPC-23 cells were exposed to gray and white MTA Plus (GMTA, WMTA), gray and white Pro-Root MTA (MTA, WMTA) cements, or their eluents. The cells were evaluated for (1) cell viability by using XTT assay, (2) apoptosis/necrosis by using flow cytometry and confocal laser scanning microscopy, and (3) oxidative stress by measuring reactive oxygen species.

Results: XTT assay showed that all test cements exhibited marked initial cytotoxicity that decreased with time. By the end of the third week, GMTA and GMTA were comparable to untreated cells (negative control) in terms of cell viability, whereas WMTA and WMTA were significantly lower than the untreated cells. Apoptosis/necrosis profiles of cells exposed to WMTA and GMTA were not significantly different from untreated cells, whereas cells exposed to WMTA and GMTA showed significantly less viable cells. All experimental groups exhibited reduction of intracellular reactive oxygen species formation compared with untreated cells, although cells exposed to WMTA were not significantly different from untreated cells. Conclusions: Both the gray and white versions of MTA Plus possess negligible in vitro cytotoxicity that are time and dilution dependent. They enrich the spectrum of hydraulic calcium silicate cements currently available to clinicians for endodontic applications. (J Endod 2014;40:255–260)

Key Words

Apoptosis, cell viability, hydraulic calcium silicate cements, necrosis, reactive oxygen species

Hydraulic calcium silicate cements (HCSCs) have become an integral component of endodontists’ armamentarium because of their bioactivity (1), biocompatibility (2), and osteogenicity (3). These cements have successfully replaced traditional dental cements in endodontic surgeries, apexification, vital pulp therapy, perforation repair, and regenerative endodontic procedures (4). Despite the favorable biological properties of current HCSCs, high operational cost, suboptimal handling properties, relatively long setting times (5), initial cytotoxicity (6), incompatibility with other restorative materials (7), and potential staining of tooth structure (8) are some of the drawbacks of contemporary HCSCs that hinder their use in endodontics. Although some of those hurdles have been addressed in more recent formulations (9, 10), none of the currently available HCSCs address all of the above challenges.

MTA Plus (Avlon Biomed Inc, Bradenton, FL; gray and white versions) is an HCSC with finner particle size that mixes with a proprietary water-based gel for enhanced handling and application as well as improved washout resistance (11). Because of its variable stoichiometric properties (12), the powder/gel ratio of MTA Plus may be adjusted to enable more diverse applications, ranging from perforation repair, root-end filling, direct pulp capping (thicker consistency) to sealing of the cleaned-and-shaped root canal space (thinner consistency).

Because HCSCs are usually applied in intimate contact with pulpal or periradicular tissues, in vitro testing of biocompatibility and cellular responses is an important preliminary step in assessing the overall biocompatibility of such cements. Thus, the objective of the present study was to evaluate the cellular viability, apoptosis and necrosis profiles, and oxidative stress levels exhibited by a rat odontoblast-like cell line after their exposure to the gray and white versions of MTA Plus.

Materials and Methods

The main constituents and primary phases of HCSCs included in the present study are summarized in Table 1. White and gray MTA Plus (WMTA, GMTA) were mixed with the proprietary hydrogel by using a liquid/powder ratio of 0.3. For comparison, white and gray ProRoot MTA (WMTA, GMTA) (Dentsply Tulsa Dental Specialties, Tulsa, OK) were also examined; these HCSCs were mixed with deionized water by using the same liquid/powder ratio. The mixed materials were placed in pre-sterilized Teflon molds (5-mm diameter and 3-mm thick), covered with pre-sterilized Mylar sheets, and allowed to set in a 100% humidity chamber for 24 hours. Untreated cells were used as the negative control. Disks of similar dimensions to the test cements and prepared from a zinc oxide–eugenol cement (Intermediate Restorative Material...
[IRM]; Dentsply Caulk, Milford, DE) were assigned as the positive control. All set materials were sterilized with ultraviolet light for 4 hours before testing.

**Cell Culture**

Rat odontoblast-like cells derived from the apical papilla (MDPC-23) were used (13). The cells were plated in complete growth medium and incubated at 37°C in a humidified 5% CO₂ atmosphere for 24 hours until fully established. The growth medium consisted of Dulbecco modified Eagle medium (Lonza, Walkersville, MD) and 10% fetal bovine serum (Invitrogen Corp, Carlsbad, CA) supplemented with 2 mmol/L L-glutamine and 100 U/mL penicillin/streptomycin.

**Cell Viability**

An XTT Cell Viability Assay Kit (Biotium Inc, Hayward, CA) was used to determine cell viability on the basis of the cleavage of the yellow tetrazolium salt 2,3-bis-(2-methoxy-4-nitro-5-sulfophenyl)-2H-tetrazolium-5-carboxanilide (XTT) by mitochondrial enzymes in metabolically active cells to form a soluble orange formazan product. Production of formazan is directly proportional to the number of vital cells and is quantified by measuring its absorbance at 490 nm. The viability of untreated cells was used as control, and absorbance of the control was adjusted to 100%, with which the relative dehydrogenase activities of the other groups were compared (N = 12). The XTT assay was performed on cells that had been directly exposed to test cements and on eluents derived from those materials.

Direct evaluation of the test materials was performed on a weekly basis according to a cycling regimen (14, 15). A weekly cycle consisted of direct evaluation of the toxicity of the cement disks over the plated cells for 3 days and indirect evaluation of the effect of eluents derived from the set cements on the plated cells. The latter was achieved by immersion of the disks in complete growth medium for 4 days to collect eluents. Accordingly, during the first part of each weekly cycle, cement and control disks were placed individually in Transwell inserts with a 3-mm pore size (BD Falcon, Franklin Lakes, NJ) to prevent direct contact of the cells by the specimen. After the inserts were placed over the plated cells, an additional 2 mL growth medium was added to each well to ensure that the level of the culture medium was above the sides of the Transwell insert. The disks were exposed to the plated cells for 3 days, without further change in culture medium, before testing for mitochondrial dehydrogenase activity. During the second part of each weekly cycle, the disks were retrieved and incubated at 37°C for 4 days to collect the eluents from the set cement before using the same disks for the next cycle. For each disk, the same growth medium was used for eluent collection throughout the entire testing period. This cycling regimen was repeated weekly for 3 weeks (ie, 3 cycles) until the material disks were rendered noncytotoxic (ie, >85% of the mean dehydrogenase activity exhibited by the untreated control).

For indirect evaluation of the eluents, each eluent concentrate collected after the 2-week aging period was diluted with fresh growth medium to 1:1, 1:5, and 1:10 of its original concentration to achieve a final volume of 2 mL (N = 12). Each diluted, eluent-containing growth medium was then used as the respective culture medium for freshly plated rat dental papilla–derived odontoblast-like cell line (MPDC-23) cells for testing cell viability.

**Apoptosis/Necrosis**

**Flow Cytometry.** After MDPC-23 cells were exposed to test materials for 3 days, the cells were detached, centrifuged, and resuspended at 1 × 10⁵ cells/mL in 1X binding buffer (Biotium Inc). The cells were...
stained with fluorescein isothiocyanate (FITC)–annexin V (AnV) \((\lambda_{\text{abs}}/\lambda_{\text{em}} = 492/514 \text{ nm}, \text{green fluorescence})\) and ethidium homodimer-III (Etd) \((\lambda_{\text{abs}}/\lambda_{\text{em}} = 528/617 \text{ nm}, \text{red fluorescence})\) and incubated for 15 minutes in the dark. The stained cells were subjected to fluorescence-activated cell sorting by using a FACSCalibur flow cytometer (BD Biosciences, San Jose, CA) to determine the percentage distribution of vital (AnV/Etd negative), early apoptotic (AnV positive, Etd negative), late apoptotic (secondary necrosis, AnV/Etd positive), and necrotic (AnV negative, Etd positive) cell populations. Experiments were performed in triplicates.

**Confocal Laser Scanning Microscopy.** Cells were plated at 2500 cells/cm\(^2\) onto coverslips in 6-well plates and exposed to the test materials for 3 days. The cells were double-stained with Hoechst 33342 \((\lambda_{\text{abs}}/\lambda_{\text{em}} = 350/461 \text{ nm}, \text{blue fluorescence})\), Etd (red fluorescence), and FITC-AnV (green fluorescence). The coverslips were mounted on slides for qualitative evaluation of intracellular ROS distribution.

**Oxidative Stress**

Detection of oxidative stress in MDPC-23 cells was performed by measuring intracellular reactive oxygen species (ROS) formation by using the CellROX Orange Oxidative Stress Reagent (Invitrogen). After the cells were exposed to test materials for 3 days, they were detached, centrifuged, and resuspended in 1% phosphate-buffered saline. CellROX Orange (a fluorescent redox cytoplasmic stain, \(\lambda_{\text{abs}}/\lambda_{\text{em}} = 545/565 \text{ nm}\) was added to the cells at a final concentration of 5 \(\mu\text{mol/L}\) and incubated at 37°C for 30 minutes. The FACSCalibur flow cytometer was used to detect the percentage of ROS-positive cells in each group \((N = 6)\). The experiment was run in triplicates; untreated cells were used for comparisons with the results derived from cells exposed to different materials. Additional cells were plated on coverslips, double-stained with CellROX Orange and Hoechst 33342, and examined with a fluorescence microscope (Axioplan 2 Imaging; Carl Zeiss) for qualitative evaluation of intracellular ROS distribution.

**Statistical Analysis**

The IRM positive control group was excluded from all statistical analyses to increase the robustness of the respective test. For XTT assay of the effect of materials on cell viability, results derived from each weekly cycle were analyzed separately by using one-factor analysis of variance (ANOVA) and post hoc Tukey test or their nonparametric equivalents. Similar tests were used for analyzing the effect of dilution of the eluents derived from the test materials on cell viability. Parametric versions of these tests were used after evaluation of the normality (Shapiro–Wilk test) and equal variance assumptions (modified Levene test) of the individual data tests. If those assumptions were violated, the data were nonparametrically transformed to satisfy those assumptions before using parametric testing methods. If those assumptions remained violated after nonlinear transformation, the original data set was analyzed by using Kruskal–Wallis ANOVA and Dunn multiple comparison tests. For apoptosis/necrosis, the numbers of vital, non-apoptotic, nonnecrotic cells in each group were analyzed by using one-factor ANOVA and post hoc Tukey test. For oxidative stress evaluation, the numbers of ROS-positive cells in each group were analyzed by using one-factor ANOVA and post hoc Tukey test. Statistical significances for all analyses were set at \(\alpha = 0.05\).
from different HSCSs were significantly more cytotoxic than cells cultured in the eluent-free medium ($P < .001$). At 1:5 dilution, only WMTA was significantly more cytotoxic than the untreated control ($P < .05$). At 1:10 dilution, there was no difference among the HSCSs and untreated control ($P = .565$).

**Apoptosis/Necrosis**

A representative CLSM image of vital, apoptotic, and necrotic cells is illustrated in Figure 2A. A 2-dimensional plot of the distribution of vital, early apoptotic, late apoptotic, and necrotic cells in a typical cell-sorting procedure is shown in Figure 2B. Cells exposed to WMTAP and GMTAP were not significantly different from unexposed (untreated) cells in the percentage of vital cells, whereas cells exposed to WMTA and GMTA had a significantly lower percentage of vital cells ($P < .001$ and $P = .002$, respectively). There was no difference between WMTAP and GMTAP ($P = .992$), as well as between WMTA and GMTA ($P = .962$) (Fig. 2C). The flow cytometry results were qualitatively confirmed by CLSM imaging. Cells exposed to the HSCSs were mostly vital and exhibited blue fluorescent nuclei with minimal signs of apoptosis/necrosis. These cells exhibited comparable fluorescence characteristics as the untreated cells. By contrast, cells exposed to IRM were mostly apoptotic, with prevalence of green fluorescent cytoplasm and occasional necrotic pink nuclei (merging of blue and red fluorescence) that are characteristic of late apoptosis and necrosis.

**Oxidative Stress**

A fluorescent microscopic image of MDPC-23 cells with Hoechst 33342–stained, blue fluorescent nuclei and diffuse orange cytoplasmic fluorescence that is indicative of the production of ROS is shown in Figure 3A. Cells exposed to IRM had very high levels of ROS (Fig. 3A). Untreated cells exhibited significantly higher ROS levels compared with cells that were exposed to GMTA, GMTAP, and WMTAP ($P < .001$), whereas there was no difference between the oxidative stress levels of untreated cells and cells that were exposed to WMTA ($P = .315$) (Fig. 3B).

**Discussion**

The use of MDPC-23 cells for the present work was based on their potential clinical significance with the use of HSCSs for direct pulp capping and for their superior sensitivity, while maintaining the same cytotoxicity ranking, when compared with transformed fibroblast and osteoblast cell lines (16, 17). The XTT assay evaluates cell viability through colorimetric quantification of formazan produced via reduction of tetrazolium salts by mitochondrial dehydrogenases (18). These enzymes are expressed only in vital cells and are inactivated shortly after cell death. Accordingly, formation of highly colored formazan dyes is indicative of a metabolically active cell population. In the present work, a modified cycling regimen (14, 15) was used to test the direct effects of test materials on cell viability at different time points as well as the indirect cytotoxic effects through eluents derived from the HSCSs (19). Results from direct and indirect evaluations were in agreement and suggest that under the experimental conditions, gray versions of both cements appear to be less cytotoxic than the white versions. The results further support previously reported findings that gray MTA is more biocompatible than white MTA (20, 21).

Analysis of the mode of cell death (apoptosis/necrosis) via flow cytometry and fluorescent microscopy is helpful in better understanding
the cytotoxic effects of the test materials on cell permeability. Exposure to MTA Plus (both gray and white versions) resulted in more vital cells than ProRoot MTA (both gray and white versions). Judging by the apoptosis/necrosis profile of all test cements, it seems that the initial cytotoxicity of MTA Plus and ProRoot MTA is more likely to be attributed to apoptosis rather than necrosis, as evidenced by detection of phosphatidylserine expression on the cell surfaces via the use of AnV (22). This suggests that the initial sites of irreversible damage by the cytotoxic agents are extracellular, whereas the nuclear membranes of those cells still remain intact.

Evaluation of intracellular ROS formation provides another perspective toward understanding of cellular responses to the test materials. ROS is a natural by-product of normal oxygen metabolism and has been found to play important roles in cell signaling, proliferation, and survival (23). Increase in ROS levels occurs during stress, results in significant damages to cell structures, and has been implicated in diseases such as cancer, aging, neurodegenerative diseases, and diabetes (24–27). Interestingly, exposure to HCSCs resulted in decreasing the ROS levels in GMTA, WMTAP, and GMP when compared with levels identified from unexposed cells. A plausible explanation is that the pH elevation caused by release of calcium hydroxide from HCSCs may decrease ROS formation. These findings support earlier reports relating the production of ROS to decreases in extracellular pH (28–30). It is of interest to see whether the decreased level of oxidative stress when MDPC-23 cells are exposed to MTA Plus may enhance the ability of these cells to differentiate and deposit mineralized matrix when they are cultured in osteogenic differentiation medium. Research in this direction is in order.

Within the limits of the present study, it may be concluded that the cytotoxic effects imposed by MTA Plus on MDPC-23 cells are both time and concentration dependent, and that they possess negligible cytotoxic risks after the elution of their cytotoxic components. Because these risks are significantly lower than those imposed by zinc oxide-eugenol–based cement, a favorable risk/benefit perspective toward understanding of cellular responses to the test materials is in order. The authors thank Mrs Petra Lockwood for her assistance with cell culture.

Dr Primus is the inventor of MTA Plus. The remaining authors deny any conflicts of interest related to this study.

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