

Cytotoxicity Evaluation of Self Adhesive Composite Resin Cements by Dentin Barrier Test on 3D Pulp Cells

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ABSTRACT

Objectives: The aim of this study was to evaluate the effects of five self-etch dental composite resin cements on the cell viability of bovine dental papilla-derived cells.

Methods: The cytotoxicity of composite resin cements (Rely X Unicem Clicker, 3M ESPE; MaxCem; KERR, Panavia F 2.0; Kuraray, BisCem; Bisco and Bistite II DC; Tokuyama) was analyzed in a dentin barrier test device using three-dimensional (3D) pulp cell cultures. A commercially available cell culture perfusion chamber was separated into two compartments by 500 µm dentin disc. The three dimensional cultures placed on a dentin disk held in place by a special biocompatible stainless-steel holder. Test materials were introduced into the upper compartment in direct contact with the cavity side of the dentin disks according to the manufacturer's instructions. Subsequently, the pulpal part of the perfusion chamber containing the cell cultures was perfused with medium (2 ml/h). After an exposure period of 24 h, the cell survival was determined by the MTT assay. Statistical analyses were performed using the Mann-Whitney U-test.

Results: In dentin barrier test, cell survival was similar with Maxcem and negative control group ($P>.05$), and all other tested materials were cytotoxic for the three dimensional cell cultures ($P>.05$).

Conclusions: The significance of composite resin cements is being more important in dentistry. The cytotoxic potencies demonstrated by these materials might be of clinical relevance. Some composite resin cements include biologically active ingredients and may modify pulp cell metabolism when the materials are used in deep cavities or directly contact pulp tissue. (Eur J Dent 2009;3:120-126)

Key words: Resin cement; Cytotoxicity; Dentin barrier test; Three dimensional cultures.

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INTRODUCTION

Resin cements are gaining popularity in the dental profession for a number of reasons. The vast majority of non-metallic restorations, which are also utilized more at the present time than before, can be cemented only with resin cements. These restorations include ceramic and resin composite inlays and onlays as well as ceramic crowns and porcelain veneers.¹ An ideal dental luting cement should be biocompatible, have little interaction with body tissues and fluids,² be

nontoxic, and have low allergic potential.³

It has been long demonstrated that different components of resinous materials can be released in an adjacent aqueous phase.^{4,5} In this way, when applied to a wet surface, such as dentin, uncured free monomers released from resin-based materials may diffuse across dentinal tubules to reach the pulpal space.⁶ Several investigations have shown that released monomers cause chemical damage to cultured cells.^{7,8} In addition, many *in vivo* studies have shown that uncured resin components which reach the pulpal space cause inflammatory response and tissue disorganization.^{9,10}

Dimethacrylate monomers are widely used in dentistry and the most widely used monomers for the preparation of resin cements are crosslinking dimethacrylates like bisphenol A glycol dimethacrylate (Bis-GMA), triethylene glycol dimethacrylate (TEGDMA), urethane dimethacrylate (UDMA), and bisphenol A ethoxylated dimethacrylate (Bis-EMA). Usually, mixtures of these monomers are used.^{11,12} Acrylates and mainly methacrylates were found to cause cytotoxic effects. Evaluation of the cytotoxicity of acrylates and methacrylates that had been used in dental resin materials showed a relationship between their structure and the degree of cytotoxicity.¹² TEGDMA, and mainly Bis-GMA and UDMA, are highly cytotoxic.¹²⁻¹⁵

Many *in vitro* studies have shown that the polymerization reaction that produces the cross-linked polymer matrix from the dimethacrylate resin monomer is never complete and adverse reactions are due to the release of nonpolymerized monomers such as TEGDMA or 2-hydroxy-ethyl-methacrylate (HEMA). Unbound free monomers seem to be directly responsible for the cytotoxicity of resin composites on pulp and gingival cells, and they are probably also implicated in the allergic potential of the material.^{16,17}

The elution of unpolymerized resin components becomes significant when these materials diffuse across the permeable hybrid layer and dentinal tubules to reach the pulpal space⁴ from a fresh cavity preparation and are of concentrations high enough to produce a biological effect upon the dental pulp. It is evident from previous work that certain precursor components of resins are cytotoxic for fibroblastic cells in culture,¹⁸

and thus could contribute to tissue injury of the dental pulp following restorative procedures.¹⁹ In this specific situation, the odontoblasts that are underlying the circumpulpal dentin are the first cell line affected by the residual components leached from the dental materials used for cavity restoration. For this reason, odontoblasts are the appropriate as a target cells to evaluate the *in vitro* side effects of dental restorative materials.²⁰ In this connection, adequate contact between cells and test material is crucial to cell cytotoxicity testing. To simulate the *in vivo* situation for dental filling materials, *in vitro* pulp chambers have been designed, introducing dentin as a barrier between test materials and target cells. Schmalz et al²¹ described an *in vitro* dentin barrier test system, which is mainly based on commercially available components, thus fulfilling a basic requirement for standardized testing techniques. An additional advantage of this artificial pulp chamber was thought to be the possibility of perfusing the pulpal part with nutrition medium, thus simulating *in vivo* pulpal blood flow. The objective of the present study was to evaluate the cytotoxic effects of five self-etch dental composite resin cements by a dentin barrier test device using the three-dimensional cultures of transformed pulp-derived cells to simulate three-dimensional structure of the dental pulp tissue.

MATERIALS AND METHODS

Cell culture

Bovine dental papilla-derived cells,²² derived from calf dental papilla were maintained in growth medium (Minimum Essential Medium Alpha, Gibco Invitrogen, Germany) supplemented with 20% fetal bovine serum (FBS) (Biochrom AG, Berlin, Germany), 150 IU/ml penicillin, 150 mg/ml streptomycin (Biological Industries, Israel) and 0.1 mg/ml geneticin (Biological Industries, Israel) in a humidified atmosphere at 37°C, 5% CO₂. For all experiments, cells within passages 19 to 21 were used.

Preparation of three-dimensional cultures

Polyamide meshes (0.5 cm²; Reichelt Chemietechnik, Germany) were immersed in 0.1 M acetic acid for 30 min, washed three times with phosphate-buffered saline, air dried

and coated with 0.03 mg/ml fibronectin (Sigma Aldrich, Germany). Cell culture inserts (Millipore, Germany) were placed in 6-well plates with 1.25 ml medium (MEM α with 20% FBS) per well. The fibronectin-coated meshes were placed on the inserts and 8×10^4 cells in volume of 25 μ l/mesh were seeded on them. After 48 h incubation (37°C, 5% CO₂, 100% humidity), meshes were transferred to 24-well plates and incubated for up to 21 days. Culture medium (supplemented with 0.05 mg/ml ascorbic acid) was changed three times a week.

Cytotoxicity testing

After 14 \pm 2 days, three-dimensional cultures were introduced into a dentin barrier test system as described;²³ a commercially available cell culture perfusion chamber (Minucells & Minutissue GmbH, Bad Abbach, Germany) made of polycarbonate with a base of 40x40 mm and a height of 36 mm was modified. The three dimensional cultures placed on a dentin disc held in place by a special biocompatible stainless-steel holder, resulting in a dentin barrier test situation. The dentin disc (500 \pm 20 μ m thick) was cut from a bovine incisor, etched on pulpal side with 50% citric acid for 30 s and autoclaved as described.²¹ Thus, the cell culture chamber was separated into two compartments by the dentin disc. The cell culture tissues were placed in direct contact with the etched side of the dentin disk and held in place by the stainless-steel holder. All chambers were perfused with 0.3 ml assay medium (growth medium with 5.96 g/L HEPES buffer, Merck, Germany) per hour for 24 h at 37°C. Perfusion was switched off; test materials were introduced into the upper compartment in direct contact with the "cavity" side of the dentin disc.

Cytotoxicity of test materials was recorded after the pulpal part of the in vitro pulp chamber was perfused with cell culture medium (2 ml/h) for 24 h of incubation at 37°C. Each material was tested 6 times; after 24 h incubation, vitality of the cultures was determined using the MTT assay.

Test materials

The materials are listed in Table 1. They were prepared according to the manufacturers' instructions.

Maxcem, is a two-part paste/paste dual-cure, self-etching and self-adhering resin cement. Its

unique formulation combines the etchant, primer/adhesive and resin cement into one material. Using the automix tip, the cement syringed into an insulin injector. Then applied 2 mm directly onto the dentin disc and light-cured (380-515 nm. LED light curing unit, Bluephase, Ivoclar Vivadent, Liechtenstein, Austria) for 20 s and continues to cure fully in 5 minutes.

Rely X Unicem Clicker, is a two-part paste/paste dual cure, self-adhesive universal resin cement. The clicker used to extrude adequate amount of base paste and catalyst paste onto the mixing pad and mixed evenly for 20 seconds with a plastic spatula and put in an insulin injector. Then applied 2 mm directly onto the dentin disc and light-cured (380-515 nm. LED light curing unit, Bluephase, Ivoclar Vivadent, Liechtenstein, Austria) for 20 s and continues to cure fully in 5 minutes.

Panavia F 2.0, is a two-part paste/paste dual-cure, self-adhesive resin cement. The pastes syringed the same amount of turn and mixed for 20 seconds on the mixing pad with a plastic spatula and put in an insulin injector. Then applied 2 mm directly onto the dentin disc and light-cured for 20 s (380-515 nm. LED light curing unit, Bluephase, Ivoclar Vivadent, Liechtenstein, Austria) and continues to cure fully in 5 minutes.

Biscem, is a two-part dual cure, self adhesive resin cement. Using the automix tip, the cement syringed into an insulin injector. Then applied directly onto the dentin disc and light-cured for 20 s (380-515 nm. LED light curing unit, Bluephase, Ivoclar Vivadent, Liechtenstein, Austria) and continues to cure fully in 6 minutes.

Bistite, is a two part paste/paste dual cure self adhesive resin cement. The syringe used to extrude adequate amount of pastes A and B onto the mixing pad and mixed evenly for 10 seconds with a plastic spatula and put in an insulin injector. Then applied 2 mm directly onto the dentin disc and light-cured for 20 s (380-515 nm. LED light curing unit, Bluephase, Ivoclar Vivadent, Liechtenstein, Austria) and continues to cure fully in 3 minutes.

A-Silicon Impression Material (President Coltene AG, Altstätten, Sweden) is used as a negative control (100% cell viability). The pastes syringed the same amount of base and catalyst and mixed for 20 s on the mixing pad and put in an

insulin injector. Then applied 2 mm directly onto the dentin disc and self-cured.

MTT assay

Cell viability of three-dimensional cultures was determined by enzyme activity (MTT assay). The tissues were removed from the pulp chambers, respectively, placed into 48-well plates containing 500 µl of prewarmed MTT (3-(4,5-dimethylthiazol-2-yl)-2,5-diphenyltetrazolium bromide), (MTT, Sigma Aldrich, Germany) solution (0.5 mg/ml in MEMα) and incubated for 2 h at 37°C. Then the tissues were washed two times with phosphate-buffered saline. The blue formazan precipitate was extracted from the mitochondria using 250 µl dimethyl sulfoxide on a shaker at room temperature for 30 min.

200 µl of this solution were transferred to a 96-well plate and the absorption at 540 nm was determined spectrophotometrically (µQuant, Bio-Tek Instruments, Inc, Winooski, VT, USA). The mean values of control tissues (cell cultures exposed to silicone impression material) were set to represent 100% viability. Results of cytotoxicity experiments were expressed as a percentage of control tissues. Statistical analysis between control and test materials was performed applying the nonparametric Mann-Whitney U test (P>.05).

RESULTS

The results of dentin barrier test with composite resin cements are summarized in Figure 1. A vinyl poly siloxane material (President

Table 1. Test materials, compounds and components.

Resin Cements	Lot Number	Manufacturer	Compound	Components
Rely X Unicem Clicker	279830	3M ESPE, Seefeld, Germany	Base, Catalyst	Powder: silanized glass powder, silanized silica, calcium hydroxide, substituted pyrimidine, sodium persulfate Liquid: methacrylated phosphoric acid esters, triethylene glycol dimethacrylate, substituted dimethacrylate
Maxcem	2712954	Kerr, Orange, CA, USA	Paste / Paste	Resin Matrix: Glyceroldimethacrylate dihydrogen phosphate (GPDM) –self-etching/self-adhering acidic monomer, Comonomers including mono-, di-, and tri-functional methacrylate monomers, Proprietary self-cure redox initiator, Photoinitiator, Stabilizer Fillers: Barium glass filler, Fluoroaluminosilicate glass filler, Fumed silica
Panavia F 2.0	Paste A: 00185A Paste B: 00027A	Kuraray, Okuyama, Japan	Paste A, Paste B	Paste A: Silica, Dibenzoyl peroxide Paste B: Silanated Barium Glass, silanated titanium oxide, sodium fluoride, colloidal silica, bisphenol A, polyethoxy dimethacrylate, hydrophilic dimethacrylate, hydrophobic dimethacrylate, N,N'-diethanol-P-toluidine, sodium 2,4,6-triisopropyl benzene sulfinate
BisCem	700004155	Bisco, Schaumburg, IL, USA	Base, Catalyst	Bis (Hydroxyethyl methacrylate) phosphate (Base), Tetraethylene glycol dimethacrylate, Dental Glass
Bistite II DC	50R-90C	Tokuyama Dental Corporation, Tokyo, Japan	Paste A, Paste B	Filler: Silica Zirconia Matrix: Dimethacrylate, MAC-10 (an adhesive promoter), initiator

was used as a negative control material. Biscem composite resin cement caused considerably cell damage only 52% of the cells survived when compared with cell cultures exposed to control (President) ($P>.05$). Exposure of the cell cultures to Rely X Unicem Clicker, Panavia F 2.0, Bistite II DC lead to 70%, 62% and 73% cell survival, respectively. Statistically, Rely X Unicem Clicker, Panavia F 2.0, Biscem and Bistite II DC groups were different from control material ($P>.05$). Maxcem composite resin cement lead to 79% cell survival, and Maxcem was statistically not different from control ($P>.05$).

DISCUSSION

When composite resin cement materials are placed onto prepared dentin surfaces, their direct toxic effects will be most likely on the pulp cells beneath the dentin.¹⁰ According to Schmalz et al,^{23,24} the in vitro dentin barrier test system for cytotoxicity tests might mimic a clinical situation and it is better than direct cell-material contact in vitro methods, and it has the potential to, at least partially, replace animal experimentation. This goal was achieved when a dentin disc, which functioned as a barrier, a test material, and three-dimensional cultures of pulp-derived cells were combined in a cell culture perfusion chamber.^{23,25} It was consistently demonstrated that dentin was an effective barrier, preventing cell damage from a great variety of materials and chemicals. For instance, cytotoxicity of dental adhesives decreased when the thickness of the dentin disc was gradually increased from 100 to 500 μm .²⁶ This effect was apparently selective and depended on the chemical nature of the dentin contacting material.^{23,25,26} In the present study the composite resin cements reduced cell viability in pulp-derived three-dimensional cell cultures even though under a 500 μm dentin thickness. In this paper it was demonstrated that only Maxcem composite resin cement is not cytotoxic and all other tested composite resin cements are cytotoxic on the three dimensional bovine dental papilla-derived cells when compared to control group.

Current investigations have also reported the cytotoxic effects of some resin monomers, such as BIS-GMA (Bis glycidyl methacrylate), UDMA (urethane dimethacrylate), and TEGDMA

(tri ethylene glycol dimethacrylate).^{7,27} These resin monomers are able to deplete intracellular glutathione as well as interfere with the expression of some proteins, such as collagen I, osteonectin, and dentin sialoprotein, which play a fundamental role in the pulp repair.^{20,28,29}

Resin monomers and other components are released from dental composite restorative materials even after polymerization. The TEGDMA and the hydrophilic monomer 2-hydroxyethyl methacrylate (HEMA) were detected among the various chemicals in hydrophilic as well as hydrophobic solvents.^{7,30,31} Resin monomers have been identified as cytotoxic by a variety of different methods, all indicating changes in basic cell structures such as cell membrane integrity and cell functions like enzyme activities or the synthesis of macromolecules.^{13,18,32} RelyX Unicem is a BIS-GMA/TEGDMA-based resin and many toxic resin monomers and chemical agents, such as TEGDMA, methacrylated phosphoric acid esters, and dimethacrylates, are incorporated in the formulation of the RelyX Unicem, it should be reasonable to expect a very high cytotoxicity of this dental cement to the culture of cells. De Mendonça et al²⁰ evaluated the cytotoxic effects of calcium hydroxide, Vitrebond, RelyX Luting, and RelyX Unicem cements, applied on the odontoblast like cells MDPC-23. The most intense cytotoxic effects were caused by calcium hydroxide, Vitrebond and RelyX Luting. On the other hand, RelyX Unicem caused low cytotoxic effects to the odontoblast cell line MDPC-23 but the results cannot be directly extrapolated to clinical situations in which the dental cements are applied on sound- or caries-affected dentin,

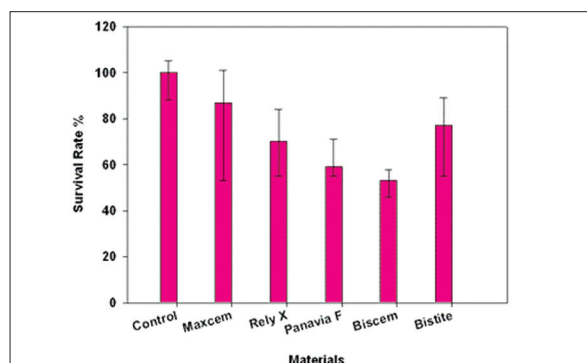


Figure 1. Cell survival of three-dimensional cultures in the dentin barrier test device after exposure to composite resin cements. Data are expressed as percentage of the negative-control cultures (President=silicone impression material). The indicated values are medians, 25th and 75th percentiles.

and the pulp-dentin complex presents intrinsic mechanisms of defense. In the present study RelyX Unicem showed significant toxic effects on the three dimensional cell cultures.

Maxcem composite resin cement is compound of a combination of adhesive monomers, including GPDM (glyceroldimethacrylate dihydrogen phosphate) and the patented Redox Initiator System that provides an efficient curing mechanism and it was demonstrated in the present investigation that Maxcem was caused the lowest cytotoxic effects.

Biscem composite resin cement contains TEEGDMA and HEMA and in the present study the most intense cytotoxic effects were caused by Biscem for the three dimensional pulp cell cultures. Chang et al³³ have shown that HEMA induces cell growth inhibition and cycle perturbation. The glutathione depletion and ROS production are key factors leading to cell apoptosis. On the other hand, we could not reach any data in the literature about cytotoxicity of TEEGDMA.

Composite resin materials may contain rather 'unknown' monomers and generally these monomers protect by patents. Patents may also hinder objective research.³⁴ Only available composition of the resin cements tested in this study showed in Table 1. They may also contain such unknown monomers.

It was considered that such a test design was appropriate for the purpose of this study, because it was more similar to the in vivo situation, where the composite resin cements covers the dentinal walls and only indirectly interacts with the pulp, by means of the tubular fluid. In this connection, Costa et al¹⁰ investigated the pulp response following cementation of inlays using two different composite resin cements. They showed that luting cements may cause specific pulpal damage. Variolink II associated with the adhesive system Excite cause more aggressive effects to the pulp-dentin complex than Rely X Unicem cement when both are used the cement inlay restorations.

CONCLUSIONS

In this study, it was concluded that the most intense cytotoxic effects were caused by Biscem> Panavia F 2.0> Rely X Unicem Clicker> Bistite II

DC. On the other hand, Maxcem caused lowest cytotoxic effects to the three dimensional cell cultures in this study.

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