Cytotoxicity, Morphology and Chemical Composition of Two Luting Cements: An in Vitro Study

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Abstract

Objective: To assess the cytotoxicity, surface morphology, elemental compositions and chemical characterization of two commonly used luting cement. Material and Methods: The two luting types of cement used were Elite Cement® and Hy-Bond Resiglass®. Freshly mixed (n=6) and set form (n=6) of each cement was placed in medium to obtain extracts. The extract from each sample was exposed to L929 mouse fibroblasts (1x10^4 cells/well). Alamar Blue Assay assessed cell viability. Surface morphology and elemental composition were evaluated using scanning electron microscopy and energy dispersive spectroscopy. The chemical characterization was performed by Fourier Transform Infrared Spectroscopy. One-way ANOVA and post-hoc Tukey analysis were conducted to assess results. Results: Hy-Bond Resiglass® was the more cytotoxic of the two types of cement in both freshly mixed (68.10 ±5.16; p<0.05) and set state (87.58 ±4.86; p<0.05), compared to Elite Cement® both freshly mixed (77.01 ±5.45; p<0.05) and set state (89.39 ±5.66; p<0.05). Scanning electron microscopy revealed a more irregular and porous structure in Hy-Bond Resiglass® compared to Elite Cement®. Similarly, intense peaks of aluminium, tungsten and fluorine were observed in energy dispersive spectroscopy in Hy-Bond Resiglass. Conclusion: All these three elements (aluminium, tungsten and fluorine) have cytotoxic potential. The Fourier transform infrared spectroscopy revealed the presence of hydroxyethyl methacrylate in Hy-Bond Resiglass®, which has a cytotoxic potential.

Keywords: Dental Materials; Dental Cements; Zinc Phosphate Cement; Fibroblasts.
Introduction

In clinical dentistry, luting cement are used for the purpose of sealing, cementing or bonding objects or particles together. By definition, it is the cement that holds a fix prosthesis, orthodontic appliances, post and core static, stable for a long period in the oral cavity. The retention mechanisms for this prosthesis can be micromechanical, mechanical and chemical. In most of the cases, a combination of two or three mechanisms are used for successful retention of the prosthesis. Ideally, these cements should be nontoxic, nonirritant, good mechanical properties, strong adhesion, excellent esthetics, radiopaque, resistant to caries and plaque accumulation, less soluble, antimicrobial, average film thickness and should possess a good sealing ability [1].

The biological compatibility with tooth vitality is an important factor that needs to be fulfilled. Cytotoxicity of any dental cement is an essential consideration because it may evoke a certain adverse reaction of allergic and/or toxic nature [2]. These can be from the chemical irritation of the materials and pH changes during setting reactions [3]. The acidic content of the cement is also a factor that may contribute or aggravate cytotoxic response; therefore, it is necessary to know the acidic potential of a cement. Dental cement releases different elements during and after setting; therefore, different biological responses are to be expected from different types of cement. Numerous studies focus on the cytotoxicity of dental materials with special regards to contact with soft tissue [4].

One of the oldest luting materials still in use today is Zinc phosphate [5]. It remains quite popular despite a documented track record of disadvantages such as increased solubility and low adhesion. In comparison, the most widely used luting material in modern dentistry is resin-modified glass ionomer cement. Despite a lot of improvements, cytotoxicity of the resin-based dental materials is still remains unsatisfactory and a challenge to deal with. Due to incomplete polymerization, there are unreacted ingredients of resin-based cement in the oral cavity and they are genotoxic and cytotoxic [6]. The resin-based materials contain monomers such as hydroxyethyl methacrylate (HEMA), bisphenol A-glycidyl methacrylate (Bis-GMA), triethylene glycol dimethacrylate (TEGDMA) and other initiating components. Due to the presence of HEMA and TEGDMA, cytotoxicity remains a considerable concern in the resin-based materials. The concentration and chemical structure of the monomer determines the cytotoxic properties of resin [7].

In this research, a comparative study and analysis were performed to monitor the changes in cell viability and metabolic activity to determine the cytotoxicity of the luting cement, followed by scanning electron microscopy with energy dispersive x-ray spectroscopy was to study surface morphology and elemental mapping of luting cement. Fourier transform infrared spectroscopy was performed for chemical characterization of the cement to list different possible components of the cement that may be responsible for the cytotoxic response.

Material and Methods

Materials and Sample Preparation

Elite Cement® (ZnPO4) and Hy-Bond Resiglass® (RMGIC) were used as per the manufacturer’s instructions (Table 1). Both materials were mixed according to the instructions of manufacturers under a sterile condition. After mixing, cement was placed in a 5x5 mm cylindrical Teflon mold [8]. Two forms of samples were prepared, freshly mixed and set state [9].
Table 1. Manufacturers and compositions of luting materials.

| Materials       | Manufacturer               | Composition                                                                 | Working Time  | Setting Time | Excess Removal |
|-----------------|----------------------------|----------------------------------------------------------------------------|---------------|--------------|----------------|
| Elite Cement®   | GC Corp., Ltd., Tokyo Japan| Powder: Zinc oxide, barium oxide, magnesium oxide, calcium oxide, silica Liquid: Zinc phosphate, phosphoric acid, aluminum, aluminum phosphate, water. | 3 to 4 minutes | 7 minutes    | Easy           |
| Hy-Bond Resiglass® | Shofu Inc., Kyoto Japan    | Powder: Polymerizable resin, ion leachable glasses, chemical initiator, photoinitiator, silica (Sio). Liquid: Polyacrylic acid, methyl methacrylate (MMA), water, hydroxyethyl methacrylate (HEMA). | 3 minutes     | 7 minutes    | Difficult      |

Preparation of Luting Extract

Each fresh sample was subjected to Dulbecco's Modified Eagle’s Medium (DMEM) by a surface-to-volume ratio of 1.25 cm²/mL [10]. The tubes were stored for 24h at 37°C. The extracts were collected after 24 hours with a pipette (Eppendorf AG, Hamburg, Germany) and stored in sterile tubes (CO2 Incubator - Esco Micro Pte. Ltd., Singapore) [9].

Each set sample was subjected to Dulbecco’s Modified Eagle’s Medium (DMEM) as per the surface-to-volume ratio of 1.25 cm²/mL [10]. The sample stored in extraction media was incubated for 24h at 37°C [10]. After 24 hours, extracts were collected with a pipette and stored.

Assessment of Cytotoxicity

Cell Culture

L929 mouse fibroblasts from American Type Culture Collection (ATCC, VA, USA) were used for this study [11]. The cell was cultured using Dulbecco’s Eagle modified medium (DMEM) supplemented with 10% fetal bovine serum (FBS, Gibco, Grand Island, NY, USA), 100 ul/mL penicillin, 2.5 ug/mL streptomycin (Invitrogen, Thermo Fisher Scientific, Grand Island, NY, USA). The culture media was changed regularly twice a week and incubated in a humidified atmosphere with 5% CO2 in air at 37°C. After the cultured cells were allowed to proliferate and their adherence at a logarithmic phase was achieved, the cells were detached by trypsinization (0.02% trypsin in 0.25% EDTA) [12]. The cells were suspended in a cell medium to activate trypsin [13]. The cells (1x10⁴ cells/cm²) were seeded at in a 96 - well plate and cultured again for 2 days.

Alamar Blue Assay

The culture medium was drawn and the cells were subjected to 1 ul of extraction media per well and incubated for 1, 3 and 7 days. After Day 1 extraction media was drawn and alamar blue reagent 10% was added in each well. After 4 hours of the incubation period, the fluorescence of each well was measured at an excitation wavelength of 530 nm and an emission wavelength of 590 nm by a fluorescence well plate reader (PR 4100, Bio-Rad Laboratories, Inc., CA, USA) [14]. Three wells per sample were used, and wells without luting cement extracts were used as a control group. The same procedures were used for days 3 and 7.

Scanning Electron Microscope (SEM) and Energy Dispersive Spectroscopy (EDS)

The surface morphology of cement was analyzed by scanning electron microscope (TESCAN Brno s.r.o., Brno, Czech Republic) at an accelerating voltage of 20.0 kV. A working distance ranging from 14-15 mm. The magnification employed was between 500X to 1 KX [15]. Energy dispersive spectroscopy was also
performed for all the samples with VEGA3 SB (TESCAN Brno s.r.o., Brno, Czech Republic). The charged particles, which were bombarded from the emission of x-rays from the sample surface, were detected by energy dispersive spectroscopy. The aim of EDS was to evaluate the chemical composition of the sample. Thus making it possible to know elements that constitute more than 0.1% of the material [15].

Fourier Transform Infrared Spectroscopy (FTIR)

Nicolet 6700 FT-IR spectrometer (Thermo Fisher Scientific, Grand Island, NY, USA) was used to collect FTIR spectra of all samples. It was equipped with attenuated total reflectance (ATR) and a photoacoustic sampling cell (PAS cell, Mtech Corporation). The PAS cell's sample chamber was cleaned by dry helium gas. In the case of ATR, samples were placed directly in contact with ATR diamond crystal. The spectra for all the samples were collected between spectral ranges of 4,000–400 cm⁻¹. The measurements of all the spectra were at 8 cm⁻¹ resolution accumulating a total of 256 numbers of scans [16].

Data Analysis

The data were inserted in SPSS 20 software (IBM Corp., Armonk, NY, USA). In order to assess two different cement groups, one-way ANOVA and post-hoc Tukey analysis were conducted. The level of significance was set at 5%.

Results

The control group maintained a survival rate of 100% across all days. Set cement had a higher survival rate, in comparison to their fresh state. On day 1, a fresh state of Hy-Bond Resiglass (75.738 ± 6.915) had a higher cell survival rate as compared with Elite Cement (44.88 ± 2.690). Similarly, on day 3, the higher cell survival rate was observed in Hy-Bond Resiglass (75.93 ± 4.77) than Elite cement (68.30 ± 5.984). Cytotoxicity analysis of the fresh luting cement on day 7 revealed a significant difference between the control group and both types of cement. The cell survival rate was better in Elite Cement (77.01 ± 5.456) then Hy-Bond Resiglass (68.10 ± 5.161).

![Cytotoxicity Analysis](image)

Figure 1. Cytotoxic analysis at days 1, 3, and 7.

Both types of cement had a greater survival rate than the control group in their set state on day 1. The cell survival rate observed was higher in Elite Cement (127.65 ± 22.21), followed by Hy-Bond Resiglass
(117.70 ± 13.30). However, the order was reversed on day 3, and the cell survival rate was in following order Hy-Bond Resiglass (91.45 ± 5.629) and Elite Cement (88.40 ± 5.22). On day 7 in their set state, both types of cement had significantly lesser survival rates than the Control group. The cell survival rate was in the following order Elite Cement (89.39 ± 5.669) and Hy-Bond Resiglass (87.58 ± 4.866).

SEM images of Elite cement revealed an irregular structure with evenly distributed particles (Figure 2A). Pores of variable sizes were observed at higher magnifications. At higher magnifications, cracks were observed in the structure of Elite Cement (Figure 2A and B). The EDS analysis of Elite Cement revealed an intense peak of zinc due to the presence of zinc oxide in its composition. Similarly, an intense peak of phosphorus was observed due to the presence of phosphoric acid, zinc phosphate and aluminium phosphate (Figure 2C).

Irregularly shaped particles were observed throughout the structure of Hy-Bond Resiglass (Figure 2D). At higher magnifications, pores of variable sizes were also found in the cement (Figure 2E). Intense peaks of tungsten, fluorine and aluminium were observed in the EDS spectrum of Hy-Bond Resiglass (Figure 2F).

FTIR Interpretations

The spectrum of freshly mixed Elite Cement revealed a stretching vibration of P-O at a wavelength of 1042 cm⁻¹. In the set form of Elite Cement spectrum, a strong, broad stretching peak at 1047 cm⁻¹ and 1044 cm⁻¹ were noticed (Figure 3A). In the spectrum of freshly mixed Hy-bond Resiglass a medium, sharp peak with an O-H bond was observed. However, in a set form of Hy-Bond Resiglass, a sharp peak with the O-H bond was observed at 3700 cm⁻¹ (Figure 3B). Table 2 shows the presence of chemical groups in luting cement along with their wavelength and absorbance.

| Peaks | Freshly Mixed | | Set Form |
|---|---|---|---|
| | Wavenumber cm⁻¹ | Absorbance | Wavenumber cm⁻¹ | Absorbance |
| Elite Cement | | | |
| O-H | 3735 cm⁻¹ | -0.010 | 3735 cm⁻¹ | -0.021 |
| H-O-H | 1717 cm⁻¹ | -0.035 | 1717 cm⁻¹ | -0.020 |
| P-O | 1047 cm⁻¹ | -0.011 | 1047 cm⁻¹ | 0.025 |
| CO-O-CO | 1044 cm⁻¹ | -0.010 | 1044 cm⁻¹ | 0.026 |

**Figure 2.** SEM with EDS for luting cements.
Discussion

The main goal of our study was to compare the cytotoxicity of two luting cement. These cements were compared with a control group. The control group maintained a survival rate of 100% across all days. Set cement in both cases had a greater survival rate in comparison to the cement in their fresh state. The alamar blue assay was used for cell survival rate analysis. The readings were taken on days 1, 3, and 7 for the freshly mixed form and set the state of the materials.

On days 1 and 3, there was a significant difference in cell survival rate for both types of cement in their fresh state. Elite cement was found more cytotoxic as compared with Hy-Bond Resiglass cement. However, the trend was opposite on Day 7, as the cell survival rate was less in Hy-Bond Resiglass as compared with Elite cement. In their set state, there was a decreasing trend in both types of cement as the days progressed. At the end of Day 7, the cell survival rate was less in Hy-Bond Resiglass in comparison with Elite cement.

This study concluded that both luting cement has the potential to trigger adverse biological responses. The gingival and pulpal cells can be affected by certain elements released from the cement. The amount of zinc released from zinc phosphate released will determine its cytotoxicity. On the other hand, factors such as BIS- GMA, TEG-DMA, unbound free monomer and HEMA release during and post-polymerization, are also contributing factors for cytotoxicity.

The scanning electron microscopy results revealed that Hy-Bond Resiglass exhibit more irregular structure as compared with Elite Cement. This irregular structure may create cracks and provide a larger surface area then a spherical structure. Both factors will not cause only accelerated degradation but may also
cause the quick release of cytotoxic elements from the cement [17]. A high peak of zinc and phosphorus was observed in the EDS spectrum of Elite cement. In a previous study on different dental materials for the evaluation of chemical elements composition using a scanning electron microscope and energy-dispersive x-ray spectroscopy. It was observed that zinc has a cytotoxic potential [18]. Intense peaks of tungsten (W), fluorine (F) and aluminium (Al) were observed in the EDS spectrum of Hy-Bond Resiglass. An earlier study conducted on the elemental composition of different dental materials suggested that aluminium and tungsten both have the potential of cytotoxicity [18].

The FTIR spectrum of Elite Cement revealed two peaks at a wavelength of 1044 cm\(^{-1}\) and 1047 cm\(^{-1}\). In a previous study, FTIR spectrum of zinc phosphate cement revealed peaks of P-O between 1040 cm\(^{-1}\)-1110 cm\(^{-1}\) [16]. These peaks revealed the presence of stretching vibrations of P-O in zinc phosphate cement. The presence of P-O may be due to phosphoric acid, and aluminium phosphate, both phosphoric acid and aluminium phosphate have a cytotoxic potential [18,19]. In another study, a peak of P-O bond at 1050 cm\(^{-1}\) in a spectrum of zinc phosphate cement was observed, revealing the presence of phosphoric acid or aluminium phosphate [20]. In Fourier transform infrared spectrum of Hy-Bond Resiglass a medium, sharp peak at 3700 cm\(^{-1}\) with O-H bond was observed. The presence of O-H peaks between 3300 cm\(^{-1}\) to 3700 cm\(^{-1}\) indicates hydrogen bonding in the material, which confirms the presence of HEMA in the material that is a cytotoxic material. In a previous study FTIR spectrum of HEMA revealed an O-H peak between 3300 cm\(^{-1}\) - 3700 cm\(^{-1}\) as well [21].

**Conclusion**

Both cements exhibit cytotoxic potential in freshly mixed as well as the set state. However, the cell survival rate was higher in the set state then freshly mixed form. Hy-Bond Resiglass had had less cell survival in both freshly mixed and set state in comparison with Elite Cement. Hy-Bond Resiglass had a more irregular structure as well as more cytotoxic elements in it. Similarly, FTIR confirms certain bonds in Hy-Bond Resiglass that are potentially cytotoxic.

**Authors’ Contributions**

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|---------|----------------------|---------------|
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All authors declare that they contributed to critical review of intellectual content and approval of the final version to be published.

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Conflict of Interest
The authors declare no conflicts of interest.

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