Effect of photoinitiator concentration on marginal and internal adaptation of experimental composite blends photocured by modulated methods

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ABSTRACT

Objective: The aim of this study was to evaluate the influence of photoinitiator concentration on marginal and internal adaptation of composites photocured by modulated methods. Materials and Methods: Composites based on BisGMA/triethylene glycol dimethacrylate and 65 wt% of filler were prepared with different concentrations of camphorquinone/amine (C1-0.5%, C2-1%, C3-1.5%). Cavities were prepared (3 mm × 3 mm × 2 mm) on the buccal surface of 120 bovine incisors and the adhesive system Adper Single Bond 2 was applied following manufacturers instruction. Specimens were then distributed according to type of composite (C1, C2, C3) and photoactivation method (high-intensity – 750 mW/cm² for 40 s; low intensity – 150 mW/cm² for 200 s; soft-start – 150 mW/cm² for 10 s + 750 mW/cm² for 38 s; pulse-delay – 150 mW/cm² for 10 s + 3 min dark + 750 mW/cm² for 38 s). Superficial and internal margins were analyzed by scanning electron microscopy, using the epoxy replica technique. The length of gaps was expressed as a percentage of the total length of the margins. Data were submitted to two-way analysis of variance and Tukey’s test (α =0.05). Results: Modulated curing methods did not influence gap formation regarding both superficial and internal adaptation. The composite with the lower initiator concentration (C1) presented higher gap formation when compared with those with higher concentrations (C2 and C3). Conclusion: Modulated photoactivation methods did not reduce gap formation for the experimental composite restorations evaluated. However, higher photoinitiator concentrations promote better marginal seal.

Key words: Camphorquinone, gap formation, modulated photoactivation methods, pulse-delay, resin composites, soft-start

INTRODUCTION

Resin composites have been widely used in direct esthetic restorative procedures. Despite the development of the resin composites, polymerization shrinkage still harms the interfacial quality. When the shrinkage takes place under confinement, due to bonding to cavity walls, stresses on the bonding interface will develop,¹ potentially leading to gap formation. The presence of gaps on the tooth/resin...
interface is considered as the first sign of restoration failure.²³

Although polymerization shrinkage is considered the main cause of the adaptation problems of composite restorations, it is an inherent characteristic of the composite and directly related to its composition.⁴‒⁶ Moreover, the photoinitiator concentration can influence the shrinkage stress⁷ and consequently, the gap formation. However, the effect of photoinitiator concentration in gap formation of composite restorations has not been evaluated.

The most frequently employed photoinitiator on dental composites is camphorquinone (CQ), which is a Type II photosensitizer (it needs a reducing agent to produce free radicals) with a light absorption peak near 468 nm. When CQ is photoactivated, in the excited form, a pair of free radicals is produced by proton abstraction. Such a process can be made highly efficient by the formation of a complex between the photoexcited sensitizer and a reducing agent, such as a tertiary amine.⁷

A higher concentration of CQ can induce a quick and high generation of free radicals, yielding the production of a faster polymerization kinetics reaction and higher degree of conversion.⁸ As a result, mechanical properties of the material may also be enhanced.⁹,¹⁰ However, studies have shown that there is an ideal level for the increase of CQ concentration and above this level, the increase in photoinitiator concentration does not benefit the final degree of conversion.¹¹

Conversely, if the concentration of photoinitiators were kept too low, resin composite could not be satisfactorily cured,¹⁰ which has been associated with poor biocompatibility,¹² color stability,¹³ physical and mechanical properties,¹⁴ wear resistance¹⁵ and potentially early failure of the restoration.

In addition, a substantial problem is that CQ is inherently yellow and has poor photobleaching, which means the yellow color remains after light irradiation. It may cause problems in color matching to natural dental substrate¹⁶ and limits the quantity of CQ that can be incorporated in dental composites.

The curing method also manages shrinkage stress and gap formation.¹⁷,¹⁹ Modulated photoactivation methods have shown some beneficial effects.²⁰‒²² A slow polymerization reaction allows a slow development of the composite stiffness and leads to better flow that can reduce the shrinkage stress.¹⁸,¹⁹,²³ Studies have demonstrated that the marginal adaptation of resin composites can be improved by light curing with low power density.¹⁷,²⁰,²⁴,²⁵ Conversely, the application of an adequate energy dose for composite curing is necessary to achieve deep and complete polymerization of the material.²⁶ Thus, modulated curing methods, such as soft-start (SS) and pulse-delay (PD) have been proposed.¹⁷,¹⁹,²¹,²⁴,²⁵,²⁷,²⁸

The SS curing method enhances marginal sealing of cavo-surface margins of composite restoratives,²⁸ especially due to initial lower viscosity of the composite, allowing a better material flow during the earlier stages of curing. SS also leads to shrinkage, surface hardness and residual monomer concentration similar to conventional photoactivation, in situations with similar radiant exposure.²¹ Moreover, the PD photocuring method, which is a variation of the SS method, can also decrease the intensity of polymerization stress.¹⁸,¹⁹ This curing method consists of an initial low irradiance exposure followed by a lag period in the absence of the curing light from 10 s to 5 min before a final high irradiance exposure.²⁷,²⁹ The initial low energy density is enough to allow the start of the polymerization reaction of the composite. The lag period increases composite viscous flow. The higher irradiance performed on the last stage of the curing method is responsible for ensuring similar physical and mechanical properties to the composite polymerized using the continuous light method.¹⁹ Some studies have associated this method with reduced gap formation, caused by the reduction of shrinkage stress.¹⁷,²⁷ However, it was also showed that PD method yields the formation of polymers with increased susceptibility to softening in ethanol, even though with similar monomer conversion in comparison with that produced using the continuous light curing technique.²¹,³⁰

The effects of modulated photoactivation method have been related to the composition of the resin material.¹,³¹ However, it is important to consider that most studies evaluating modulated photoactivation methods were conducted using commercial composites where fillers, resin matrix and photo initiation system differ considerably. Therefore, it seems to be more appropriate to use a resin composite model in which the variables could be controlled.

Thus, the aim of this study was to determine the influence of photoinitiator concentration on superficial and internal adaptation of experimental composites
photocured by different methods. The first tested hypothesis was that a higher concentration of photoinitiator can increase gap formation. The second tested hypothesis was that modulated curing methods reduce gap formation.

MATERIALS AND METHODS

Formulation of experimental composites
Monomer mixtures consisting of 65 wt% of BisGMA (bisphenol-a-glycidyl methacrylate) and 35 wt% of triethylene glycol dimethacrylate (TEGDMA) were prepared. The light-curing initiator system selected was CQ as photoinitiator and dimethylaminoethyl methacrylate (DMAEMA) as the co-initiator (proportion 1:1 by weight). The light-curing initiator system (CQ/DMAEMA) was thoroughly dissolved in the monomer matrix in the following concentrations: 0.5 wt% (C1-0.25% CQ/0.25% DMAEMA), 1.0 wt% (C2-0.5% CQ/0.5% DMAEMA) and 1.5 wt% (C3-0.75% CQ/0.75% DMAEMA). Furthermore, the inhibitor butylated hydroxytoluene was added to the organic matrix in a concentration of 0.1 wt% to avoid spontaneous polymerization of the monomers. The organic matrix was reinforced with silanized barium aluminum silicate glass fillers (BaAlSi – average size: 0.04 µm) and silica (SiO₂ – average size: 0.04 µm). The fillers were gradually added in the resin matrix and mixed homogeneously to a 65 wt% loading. Considering this filler content, 80 wt% were BaAlSi and 20 wt% were SiO₂. The manipulation of the experimental composites was carried out under filtered orange light. The formulation of the experimental composites used in this study was selected based on previous studies. [3,32]

Specimen preparation
A total of 120 bovine incisors were selected, cleaned and stored in a 0.5% chloramine T solution at 4°C for no more than a week. Roots were sectioned 1 mm below the cement-enamel junction using a double-face diamond saw (K. G. Sorensen, Sao Paulo, SP, Brazil). The buccal surface was ground on a water-cooled diamond saw (K. G. Sorensen, Sao Paulo, SP, Brazil). This reduction of irradiance was obtained using a standard separator. The manipulation of the experimental composites was carried out under filtered orange light. The formulation of the experimental composites used in this study was selected based on previous studies. [3,32]

A cavity with 3 mm width, 3 mm length and 2 mm deep was prepared on the flattened surface using cylindrical diamond tips (#56, KG. Sorensen, São Paulo, SP, Brazil) mounted in a standard preparation device. In this condition, cavity volume and C-factor were respectively 18 mm³ and 3, 6. Diamond tips were replaced after every fifth preparation. At the superficial margin, the cavity walls had a 90° angle with the dentin surface plane, while the internal cavity angles were rounded due to the design of the diamond tip used. If any sign of pulp exposure was noticed during the cavity preparation, the specimen was discarded.

After cavity preparation, each specimen was restored using an etch-and-rinse adhesive system (Adper Single Bond 2, batch #3HR, 3M/ESPE, St. Paul, MN, USA), applied in accordance with manufacturer’s instructions: Cavity was etched with 35% phosphoric acid gel (Scotchbond Etchant, 3M/ESPE, St. Paul, MN, USA) for 15 s, rinsed for 10 s and blot-dried. The adhesive system was applied twice with a 5 s interval in between, dried carefully with air for 15 s in order to remove residual solvent (observing a glossy surface) and light cured for 20 s using a quartz tungsten halogen (QTH) light (XL 2500, 3M/ESPE, St. Paul, MN, USA) with an irradiance of 700 mW/cm². Afterward, the teeth were randomly assigned into six groups (n = 10), according to the type of composite (C1, C2 and C3) and curing method (high-intensity; low-intensity; SS; PD). Cavities were bulk filled with one of the tested experimental composites. Curing methods are described in Table 1. The same QTH curing unit (XL 2500, 3M/ESPE, St. Paul, MN, USA) was used for all curing methods using radiant exposure around 28 J/cm². The irradiance was frequently checked by a radiometer (Demetron Research Corp., Danbury, USA).

After the light curing procedures, the specimens were stored in distilled water at 37°C for 24 h. After that, the specimens were finished under running water using 600- and 1200-grit SiC abrasive paper (Carbimet Disc Set, #305178180, Buehler, Lake Bluff, IL, USA) to

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The reduction of irradiance was obtained using a standard separator.
expose and polished with 1 µm and 0.5 µm diamond pastes in a polish cloth under water. Specimens were ultrasonically cleaned for 5 min between finishing and polishing steps.

**Evaluation of marginal and internal adaptation by scanning electron microscopy**

In order to evaluate the marginal adaptation of the restorations, impressions were taken with a low viscosity polyvinyl siloxane material (Aquasil, Dentsply DeTrey, Konstanz, Germany) and poured using an epoxy resin (Buehler, Lake Buff, IL, EUA) to obtain replicas. Afterward, the replicas were gold-sputter coated (Balzers-SCD 050 Sputter Coater, Liechtenstein) and observed by SEM (JEOL, JSM-5600 LV, SEM, Japan) for evaluation, measurement and classification of the cavity margins. Replicas were visualized at ×25 and ×200 magnifications. The classification of the margins was made at ×200 magnification directly on the microscope monitor and the measurements were made by using a multi-point measuring device that allowed the observation of the entire perimeter of the restoration at ×25. The perimeter of the restorations was measured and margins were recorded and classified according to the morphologically defined parameters previously described by Kemp-Scholte and Davidson.[33] (1) Perfect margin: Defined as a continuous, gap-free transition between filling and tooth substrate and (2) marginal gap: Observed as gap formation with the loss of interfacing adhesion. Marginal gap formation was calculated and expressed as a percentage of the cavity perimeter of each specimen. The length of the gap formed was calculated as a percentage of the entire margin length.

After the evaluation of superficial adaptation, the restorations were cut in slices (1 mm thick) in the buccal-lingual direction, using a cutting machine (ISOMET 1000, Buehler, Lake Buff, IL, USA) to obtain three slices of each restoration for internal adaptation measurements. Thus, the sectioned slices were replicated and the internal margins of the restorations evaluated according to the procedures described for the superficial adaptation protocol.

**Statistical analysis**

Original gap formation data didn’t show normal distribution. For this reason, data were transformed ($x = \text{arc} \sqrt{x} / 100$). After transformation, data showed normal distribution according to Kolmogorov-Smirnov test. Transformed data (gap %) were subjected to two-way analysis of variance (ANOVA), considering the factors “composite” and “photoactivation method” and Tukey’s test at a significance level of 5%. All statistical analysis was executed in Assistat Beta 7.5 software (ASSISTAT – Statistic Assistance by Prof. Dr. Francisco de Assis Santos e Silva, DEAG - CTRN – University of Campina Grande – PB – Brazil).

**RESULTS**

Superficial gap formation (%) means and standard deviations for all photoactivation methods are listed in Table 2. According to two-way ANOVA, for superficial marginal adaptation, only the factor “composite” had statistical differences. The factor “photoactivation method” as well as the interaction between the two factors was not statistically significant. There was no difference between the curing methods, regarding the composite used. In addition, the composite C1 presented higher gap formation when compared to C2 or C3.

Internal gap formation (%) means and standard deviations for all photoactivation methods are listed in Table 3. As observed for superficial marginal adaptation, only the factor composite showed a significant difference, regardless of the photoactivation method. In the accompanying results of marginal gap formation, there was no difference between the curing methods, regarding the composite used. In addition, the composite C1 presented higher gap formation when compared to C2 or C3.

| Table 2: Superficial gap formation (%) means and standard deviations for all photoactivation methods |
|-------------------------------------------------|-------------------------------------------------|-------------------------------------------------|
| Photoactivation methods | C1 (0.5%) | C2 (1%) | C3 (1.5%) |
| High intensity | 45.99 (21.94) | 37.89 (23.83) | 35.16 (17.76) |
| Low intensity | 41.64 (16.44) | 32.20 (19.61) | 31.55 (21.54) |
| Soft start | 48.14 (16.71) | 22.80 (20.18) | 31.73 (21.50) |
| Pulse delay | 38.75 (17.64) | 30.35 (22.04) | 29.29 (17.25) |

| Table 3: Internal gap formation (%) means and standard deviations for all photoactivation methods |
|-------------------------------------------------|-------------------------------------------------|-------------------------------------------------|
| Photoactivation methods | C1 (0.5%) | C2 (1%) | C3 (1.5%) |
| High intensity | 22.6 (12.54) | 1.5 (2.46) | 0.0 (0.00) |
| Low intensity | 04.0 (05.87) | 0.0 (0.00) | 0.0 (0.00) |
| Soft start | 16.7 (10.12) | 0.0 (0.00) | 0.0 (0.00) |
| Pulse delay | 10.2 (07.45) | 0.0 (0.00) | 0.0 (0.00) |

*Mean values followed by different small letters in the column and capital letters in the row differ statistically among themselves for the Tukey test at the level of 5%*.
DISCUSSION

Photoinitiator concentration is a fundamental parameter that determines the polymerization characteristics of a resin composite. This concentration varies among the commercial composites and its effects on marginal and internal adaptation are not still completely elucidated. Taira et al. found CQ concentrations that range from 0.17 to 1.03 wt% of the resin and Shintani et al. reported a range of 0.03-0.09 wt% of the dental composite. In this study, the concentration of CQ ranged from 0.25 to 0.75 wt% of the resin matrix (which corresponds to 0.087-0.26 wt% of the composite) and it has affected superficial and internal gap formation. However, this effect was opposite that expected: The composite with lower concentration of initiators (C1) presented a higher gap percentage than composites with a higher concentration (C2 and C3), rejecting the first hypothesis. Alonso et al. using the same composite formulation as the present work, found a significant reduction of monomer conversion when C1 (0.5% of CQ) was compared to C3 (1.5% of initiator), showing that the higher the initiator concentration, the higher the conversion degree. In this sense, it was expected that C3 would present higher gap formation, since a higher degree of conversion means higher shrinkage, higher elastic modulus and higher stress at the bonding interface. However as stated, C1 restorations showed the highest incidence of gaps. This condition was attributed to the formation of a deficient polymer network. The poorly polymerized composite would not adequately bond to the adhesive resin, allowing gap formation, as observed in Figure 1.

Photoinitiator concentration of a resin composite must be sufficient to allow an adequate polymerization. An insufficient polymerization would affect biocompatibility of the composite, due to residual monomer release. Geurtsen et al. showed that CQ and TEGDMA are released to the aqueous substrate by unsatisfactory polymerized resin materials and can damage oral tissue when presented in high concentration. In addition, bonding means higher shrinkage, higher elastic modulus and higher stress at the bonding interface. However as stated, C1 restorations showed the highest incidence of gaps. This condition was attributed to the formation of a deficient polymer network. The poorly polymerized composite would not adequately bond to the adhesive resin, allowing gap formation, as observed in Figure 1.

A satisfactory marginal seal of composite restorations is important for its clinical success. Esthetics and bond durability of these restorations are directly influenced by the marginal seal. In general, the origin of gaps has been related to three main factors: Contraction stress of composite restoratives, bonding failures, and external mechanical or thermal stress. In the present study, specimens were not submitted to any mechanical or thermal stress and bonding procedures were standardized for all groups. This condition enabled the evaluation of gap formation as a result of contraction stress of the experimental composites with different photoinitiator concentrations and different curing methods. Therefore, it is important to state that besides the care taken during specimen preparation, gap formation studies usually show high variability of the data, as inside each group some specimens present a perfect seal while others present a high percentage of gaps. Data variability in gap formation studies can be attributed to the great variability of the dental substrate, especially considering bonding to dentin. For this reason, it is important to use a minimum of 10 specimens per group.

The majority of the specimens showed a perfect seal of the superficial enamel margins in the present study, as observed in Figure 2. Enamel presents high mineral content, allowing good and stable bond strength when
convenient by reducing the polymerization rate, which favors a slower contraction stress development. On the other hand, this effect was not able to reduce gap formation in the present study. This suggests that the effects of reduced curing rates on contraction stress are limited and significant reductions in stress can be verified only after the curing rate drops below a certain threshold, as stated by Braga et al. In this sense, it could be speculated that the determinant factor for gap formation would be the energy dose applied, so that composites photocured with similar energy density presents similar gap formation. This speculation is supported by others studies considering that the total volumetric contraction of a composite and maximum stress of this system depend on monomer conversion, which is energy-dose-dependent. Alonso et al. showed that the degree of conversion of the composites tested here was similar for all photoactivation methods as the same energy dose was applied.

Another studied factor was the curing protocol. It was hypothesized that modulated photoactivation methods could reduce gap formation (second hypothesis). However, this hypothesis was not confirmed in this study. Modulated curing methods did not improve marginal or internal adaptation of composite restorations, corroborating Lopes et al. and Amaral et al. Irradiance can directly influence the polymerization rate, which is related to shrinkage stress development. The higher polymerization rate decreases composite flow, leading to more frequent gap formation on the tooth/restoration interface due to the fast development of rigidity, reducing the viscoelastic period. Thus, the increase of elastic modulus added to polymerization shrinkage yields stress development at the bonding interface, which has been related to marginal adaptation failure of composite restorations. In this sense, the low irradiance method or the modulated ones can be other studies considering that the total volumetric contraction of a composite and maximum stress of this system depend on monomer conversion, which is energy-dose-dependent. Alonso et al. showed that the degree of conversion of the composites tested here was similar for all photoactivation methods as the same energy dose was applied.

Some studies have shown that the efficacy of modulated methods vary according to composite formulation. In this way, modulated methods would show beneficial effects only for specific formulations. Another disadvantage of these methods is the decrease of cross-link density on polymer network, caused by the lower irradiance application, yielding the composite matrix to be more susceptible to degradation.

In accordance with the results of the present study, resin composites with different photoinitiator concentration show different marginal and internal adaptation features, perhaps due to the polymerization behavior. However, it is not affected by modulation of light intensity; composite restoration photocured by modulated methods presents similar marginal quality as the energy dose was kept constant. Thus, only the understanding of the problem statement, allied to the development of techniques to reduce their consequences, would help clinicians to obtain

![Figure 2: Photomicrography of restoration with perfect marginal seal. Between the white arrows, it can be observed the sealed enamel margins. Most of specimens showed perfect marginal seal. E: Enamel, RC: Restorative composite](image)

![Figure 3: Photomicrography of internal gap formation. (a) Presence of gap at the axiopulpal angle and pulp wall. The loss of interfacial adhesion can be observed. (b) Gap between dentin and restorative composite. D: Dentin, RC: Restorative composite](image)
the maximum benefits of resin composite restorations in clinical practice.

CONCLUSION

Modulated photoactivation methods did not reduce gap formation of composite restorations. Therefore, higher photoinitiator concentrations in composite allow better marginal seal for the tested conditions. The composite with 0.5% of photoinitiators yielded increased superficial and internal gap formation.

ACKNOWLEDGMENT

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