Effect of Curing Method of a Dual-cure Resin Cement on Monkey Pulpal Reaction after Bonding of Tooth-colored Inlay

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INTRODUCTION

Advances in dental adhesives have led to the development of an indirect adhesive procedure for esthetic restorations. Tooth-colored inlays, bonded to the tooth structure using a resin cement, have fairly widespread clinical applications, such that they have become a viable and aesthetic alternative to amalgam or metal inlays for posterior tooth restorations¹-⁴. Leveraging on the successful improvements in the adhesives for direct composite restorations, the bonding capacity of resin cements has also been improved by employing similar functional monomers used in the adhesives for direct composites. Formation of a hybrid layer that consists of a molecular-level mixture of adhesive polymers and hard dental tissues is recognized as one of the attributes that contributes to the high bond strength and minimum leakage of adhesive resin cements⁵.

Nonetheless, evidence of insufficient marginal sealing of bonded resin inlays has also been shown in both in vivo and in vitro studies, especially if resin cements were self-cured without any photoirradiation⁶. While adhesively luted inlays are believed to reduce the side effects of polymerization contraction of composites over direct restorations, lower bond strength values of self-cured resin cements due to the lower degree of conversion of resinous materials have been reported⁷-⁸. It has been demonstrated that the amount of unconverted monomers within the resinous materials is significantly influenced by the curing strategy used for the material.

Once these residual monomers reach the pulp tissue, a foreign body reaction may be elicited, causing some inflammatory effects⁹-¹¹. Monomers like Bis-GMA, HEMA, UDMA, TEGDMA, initiators, and solvents have already been associated with cytotoxic effects on fibroblast and odontoblast-like cells¹²-¹³. Several reports have indicated that these components individually or in combination ultimately compromise the pulp healing process¹⁰,¹²,¹³. Moreover, in deep cavity preparations, the high water content may prevent adequate polymerization of resinous materials, which in turn may release high levels of unconverted components into an adjacent aqueous phase that can diffuse through dentin to the pulp¹⁰,¹¹. On the contrary, interactions between the different resinous materials were reported to be less toxic than the sum of the individual toxicities¹⁴. Additionally, although several in vitro studies have been developed for the testing of new adhesive materials, the complex morphological and physiological characteristics of dentin and pulp are yet to be duplicated in vitro¹⁵. For example, intrapulpal pressure could also reduce the cumulative diffusion of unconverted monomers through dentin¹⁶.

The aim of this study, therefore, was to investigate the effects of the curing method of dual-cure resin cement on the reaction of monkey pulpal tissue after the bonding of tooth-colored resin inlays. In vivo bacterial leakage after the setting of resin inlays with two curing methods was also examined by means of histostaining of bacteria.
MATERIALS AND METHODS

The treatment and use of animals in this study were approved by the Institutional Animal Care and Use Committee at Tokyo Medical and Dental University and Hokkaido University, in conformance to the protocol and facilities approved by the Committee on Ethical Guidelines for Animal Care of Tokyo Medical and Dental University. As for all the materials used in this study, they were handled according to the manufacturers’ instructions.

Preparation of composite resin inlays

Resin inlays were prefabricated prior to cavity preparation, as described below. Impressions of approximately 2-mm end of diamond stones (ISO # SB2, GC, Tokyo, Japan) were obtained using the wash technique with an addition silicone (Exafine, GC, Tokyo, Japan), which in turn were used as a mold for the composite resin inlays (Fig. 1). A resin composite for indirect restorations (Estenia, Shade DA2, Lot # 00209B, Kuraray Medical Inc., Kurashiki, Japan) was placed into the impressions and irradiated for 60 seconds using a visible light curing unit for the laboratory (D-Light II, J Morita Co., Tokyo, Japan), prior to heat curing at 110°C for 15 minutes in air (KL 100, Kuraray Medical Inc.) (Fig. 2). Very small cylinders of resin inlay, 2 mm high with the same size as the diamond stone, were removed from the impression (Fig. 2) and silanized with the application of a silane agent (Ceramic Primer, Lot # 4WA, 3M ESPE, St. Paul, MN 55144-1000, USA) for five seconds.

Cavity preparation and bonding procedure

Two monkeys (Macaca fuscata, estimated age: 3-4 years, weight: 5-6 kg) were anesthetized by intramuscular injection of 2 mg/kg ketamine (Ketaral, Sankyo Co., Tokyo Japan) and intravenous injection of 2 mg/kg pentobarbital sodium (Nembutal Sodium Solution, Abbott Laboratories, Abbott Park, IL).

Cervical class V cavities of approximately 2 mm in depth were prepared in the teeth using diamond stones (ISO # SB2, GC) at high speed under water spray coolant. The cavosurface margin of the cavities was always in enamel.

The cavities were etched with phosphoric acid gel (Scotchbond etchant, Lot # 4HP, 3M ESPE) for 15 seconds, followed by vigorous water spray for at least five seconds and airing for one second, keeping the cavity floor moist. Two coats of adhesive (Adper™ Single Bond, Lot # 4KE, 3M ESPE) were applied and air-thinned by a gentle stream of air for five seconds, followed by photoirradiation for 10 seconds. Dual-cure resin cement (RelyX™ ARC, Lot # ELEU, 3M ESPE) was then handmixed and applied, and hybrid composite inlays (Estenia, Kuraray), which were prepared beforehand, were bonded to the cavities.

A total of 50 cavities were divided into two groups of 25 cavities each corresponding to the different curing strategies for the resin cement. In one group, light curing was carried out using a quartz-tungsten-halogen curing unit (Curing light XL3000, 3M ESPE) for 40 seconds, with a light output not less than 550 mW/cm² (light-cure group). In the other group, the resin cement was self-cured for six minutes without photoirradiation (self-cure group). After setting for 30 minutes, resin cement flash was removed with a high-speed superfine diamond bur (ISO # V16ff, GC).

Histological evaluation

Experimental periods for the pulpal test were set at seven, 28, and 70 days to follow the ISO guidelines. Curing methods and experimental periods were
randomly assigned to each monkey’s teeth, so that each experimental period included eight to nine teeth for each curing method. After the prescribed periods, the monkeys were sacrificed by an intravenous injection of 250 mg/kg of thiopental sodium (Ravonal, Tanabe Pharmaceutical Co., Osaka, Japan), and the teeth were removed from the jaws. Following fixation in phosphate-buffered solution of 4% formalin and 1% glutaraldehyde for one day, the teeth were decalcified with EDTA solution (Decalcifying Solution B, Wako Pure Chemical Industries Ltd., Osaka, Japan) at 4°C for four days, and then washed with running water for six hours. After removing the restorations, the teeth were dehydrated and finally embedded in paraffin.

Serial histological sections of 5 μm thickness were prepared and mounted on charged slides. Hematoxylin and Eosin, and Brown & Brenn gram stains were used. For all sections, four histological features (disorder of odontoblasts, inflammatory cell infiltration, reparative dentin formation, and bacterial staining) were evaluated. The evaluated features were then graded under a light microscope as None (0), Slight (1), Moderate (2), or Severe (3)18,19. Cavities that showed apparent pulpal exposure, namely two cavities in the self-curing group and three cavities in the light-cured group, were excluded from this study. The criteria are shown as follows:

**Disorder of odontoblasts**

None (0): Characterized by an absence of remarkable changes in the odontoblasts beneath cut dentinal tubules;  
Slight (1): Characterized by a slight disarrangement of odontoblasts beneath cut dentinal tubules;  
Moderate (2): Characterized by a disarrangement of most of the odontoblasts beneath cut dentinal tubules;  
Severe (3): Characterized by a severe disarrangement or disappearance of odontoblasts beneath cut dentinal tubules.

**Inflammatory cell infiltration**

None (0): Characterized by an absence of inflammatory cells;  
Slight (1): Characterized by the scattering of a small number of inflammatory cells;  
Moderate (2): Characterized by a distinct increase in inflammatory cells;  
Severe (3): Characterized by abscess formation in the pulp or pulpal necrosis.

**Reparative dentin formation**

None (0): No additional or abnormal increased thickness in circumpulpal dentin beneath cut dentinal tubules of cavity preparation;  
Slight (1): A small thin rim of reparative dentin beneath cut dentinal tubules of cavity floor (thickness of the reparative dentin was less than twice the thickness of predentin);  
Moderate (2): A moderate bulk of new reparative dentin beneath cut dentinal tubules of cavity floor (thickness of the reparative dentin was more than twice the thickness of predentin, but less than half that of pulpal thickness);  
Severe (3): A large bulk of new reparative dentin beneath cut dentinal tubules of cavity floor (thickness of the reparative dentin was more than half that of pulpal thickness).

**Bacterial staining**

None (0): Absence of bacterial staining profiles throughout the section;  
Slight (1): Positive bacterial staining profiles along the coronal wall or axial floor of cavity;  
Moderate (2): Positive bacterial staining profiles within the cut dentinal tubules of axial floor;  
Severe (3): Positive bacterial staining profiles within dental pulp.

Results pertaining to disorder of odontoblasts, inflammatory cell infiltration, reparative dentin formation, and bacterial staining were statistically analyzed using the Kruskal-Wallis test, with statistical significance defined as p<0.05.

The remaining dentin thickness between the cavity floor and pulp (RDT) was also recorded, and variances in thickness were analyzed using one-way ANOVA (p<0.05).

**RESULTS**

Table 1 shows a summary of the findings in the histological sections and RDT. No statistically significant differences in RDT were found among the groups (one-way ANOVA, F=1.060, p=0.3973).

**Disorder of odontoblasts**

Moderate to severe disorders were detected with both curing methods for all experimental periods. In the self-cured group, one moderate and three slight disorders of eight specimens were seen at seven days; one severe, one moderate, and one slight disorder of six specimens were seen at 28 days; and one moderate and five slight reactions of nine specimens were seen at 70 days (Fig. 3).

The light-cured group showed one moderate and three slight disorders of seven specimens at seven days; two moderate and one slight disorder of seven specimens at 28 days; and one moderate and one slight reaction of eight specimens at 70 days (Fig. 4).

Statistical analysis showed no significant differences in odontoblastic disorder between the two curing methods in any of the experimental periods.

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*pulpal reaction to a dual-cure resin cement*
Inflammatory cell infiltration

Inflammatory cell infiltration was seen only at 28 days. Two moderate and one slight reaction of six specimens were detected in the self-cured group, and one moderate reaction was seen in the light-cured group (Fig. 5).

Statistical analysis showed no significant differences in inflammatory cell infiltration between the two curing methods in any of the experimental periods (Kruskal-Wallis test, p=0.5994).

Reparative dentin formation

At seven days in both test groups, there were no indications of any reparative dentin beneath the cut tubules of the remaining dentin.

The self-cured group showed one case of moderate and one case of slight dentin formation of six specimens at 28 days, and three cases of moderate and one case of slight formation of nine specimens at 70 days (Figs. 5 and 6). In the light-cured group, one case of moderate formation of seven specimens was seen at 28 days, and one case of moderate and one case of slight formation of eight specimens were seen at 70 days (Fig. 7).

Statistical analysis showed no significant differences in reparative dentin formation between the two curing methods in any of the experimental periods (Kruskal-Wallis test, p=0.5805).

Bacterial staining

Bacterial penetration along the cavity walls could not be detected in any of the experimental groups.
DISCUSSION

This *in vivo* study evaluated the effects of two curing strategies of a dual-cure resin cement on pulpal irritation level toward resin inlay restorations using monkey pulp. Owing to the limitations and difficulties of animal treatment, resin inlays were fabricated in advance and set immediately after cavity preparation. Impression or provisional restoration could be omitted in this process due to the employment of standardized cavity preparation and standard-sized resin inlays. Although a few points of discrepancy exist from the clinical steps for indirect restorations, a previous *in vivo* study indicated slight pulpal irritation resulting from impression and temporization. This is because a provisional restoration helps to heal pulpal damage resulting from cavity preparation, and therefore irritation levels following the clinical steps may be lower.

Results of this study indicated that both light-curing and self-curing methods of resin cement induced slight to moderate levels of odontoblastic disorders in the pulp after a seven-day test interval. At seven days, since similar irritation levels of odontoblastic disorders without any inflammatory cell infiltration were observed in both curing methods, these reactions seemed to arise from the deep cavity preparation or bonding process for the setting of resin inlay. It is widely recognized that pulpal irritation levels of cavity preparation increase if the remaining thickness of cavity floor dentin is thin. In the present study, cavity depth was controlled to approximately 2 mm. Employing this standardized cavity preparation, the remaining dentin thickness (RDT) of three-day specimens ranged from 0.20 to 0.85 mm. While a few specimens with extremely deep cavity preparation showed moderate levels of disarrangement, the correlation between RDT and odontoblastic disorders could not be detected due to the controlled cavity depth.

In this study, phosphoric acid etching gel attached to the resin cement was applied to the cavity wall to remove the smear layer. Several studies have shown that total-etched dentin by phosphoric acid sometimes enhanced the reduction of odontoblasts more than nonetched cavities at the initial stage, but these differences did not persist over the long term. After etching and rinsing, we kept the etched dentin moist prior to application of the adhesive, following the instructions for this material. Since the observed odontoblastic changes with the two curing methods were only mild to slight with lower degree of odontoblastic reduction, the wet bond method could indeed reduce the irritation level of the etching process. Meanwhile, the effects of air-drying on odontoblastic changes have been discussed, especially...
After total etching, and this mild level of reaction appeared to be due to the less air-drying.

At 28 days, the light-cured group showed a moderate reaction of inflammatory cell infiltration. At the same time, one slight and two moderate reactions were observed in the self-cured group. Although no significant differences in inflammatory cell infiltration were detected between the two curing methods, the self-cured specimen showed a slightly higher number of inflammatory reaction cases. It is well known that the conversion of monomers to polymers is never complete, regardless of the type of activation. Unreacted resinous materials like Bis-GMA, HEMA, UDMA, TEGDMA, initiators, and solvents leach out from the material and have been shown to cause cytotoxic reactions in in vitro studies.

Since light irradiation can achieve a relatively higher conversion ratio of resin cements over the self-cure method, pulpal irritation due to unconverted resinous materials might be lower. However, these cases of inflammatory cell infiltration observed at 28 days disappeared at 70 days for both curing methods. Several reports of in vitro studies have indicated that the release of unconverted components into an adjacent aqueous phase was at its highest during the initial immersion, with levels decreasing thereafter. Accordingly then, this result seemed to indicate that the irritation levels due to unconverted products from the cement were low, even when it was self-cured.

In this study, prior to the application of RelyX™ ARC cement, total-etch adhesive was applied to the cavity wall and light-cured, creating a resin-tooth hybridized zone with an approximate depth of 5 to 10 mm. Since the sealing effect of this hybridized zone has been shown in a series of studies, it could be said that this zone blocked the diffusion of unreacted resinous materials to the pulp, resulting in lower levels of pulpal reaction with both self-cure and light-cure methods.

It is noteworthy that odontoblastic disorders remained in both self-cured and light-cured specimens even after 28- or 70-day test intervals. Since traumatic effects due to cavity preparation should have largely disappeared by the intermediate ISO-stipulated time interval, these remaining odontoblastic disorders could be an indication of some stimulant effect from the restoration itself. Although the reason for this remaining disorder was still unclear, unconverted resinous materials within the cement might in part induce this reaction. It is well documented that, after dental injury and irreversible odontoblastic damage, dental pulp is able to produce a reparative dentin matrix - which is secreted by a secondary generation of odontoblast-like cells. On this note, our results were in agreement with the previous report: whereby slight to moderate levels of reparative dentin formation were found at the 70-day test interval. Consequently, it was highly probable that this remaining disorder indicated a remodeling of odontoblasts and that the healing process still continued at 28 and 70 days. Indeed, the relatively lower level of disorders observed in the 70-day light-cured specimens compared with the 28-day specimens seemed to support this speculation.

Despite rapid progress in the advancement of adhesive materials coupled with high bond strength rendered by recently developed resinous materials, microleakage along restorative sites remains a significant challenge in bonded inlays. However, in this in vivo study, no bacterial penetration was detected the class V cavity wall for up to 70 days, when the resin inlays were bonded with RelyX™ ARC. In light of this result, we could conclude that the bonding performance of this resin cement was excellent for both self-cure and light-cure methods. It should also be mentioned that the adhesive employed in this resin cement seemed to contribute to this ideal result regarding sealing capacity.

In view of previous adhesion results obtained from bonded resin inlays to dentin using self-cured and light-cured resin cements, it was of interest for us to determine the bacterial leakage behavior of this material. Nonetheless, it is common knowledge that adhesively luted resin inlays significantly reduce the side effects of polymerization stress of resin composites, and that the bonding capacity of self-cured resin cements is generally inferior to that of light-cured cements. Within the limitations of this in vivo study, self-cured RelyX™ ARC could bond the resin inlay up to 70 days without bacterial penetration. At this juncture, it should be put into perspective that the curing process of resin cements is a complicated one. During the transition from liquid to viscoelastic phase, the resin cement can be used without inducing damage to the internal structures of both the cement and adhesive. Since this soft and flowable stage continues longer in the self-cure method, movement of resin cement during shrinkage might compensate for the shrinkage, thus resulting in an unexpected but excellent marginal seal.

The results of this present study were limited to biocompatibility and in vivo microleakage in monkey teeth only. Therefore, it was deemed both expedient and necessary to carry out further bonding durability tests under occlusal stress as well as more detailed clinical evaluations of the resin cement, RelyX™ ARC.

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REFERENCES


