Adhesive Strength of Paint-on Resins to Crown and Bridge Composites

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This paper examined the adhesive strength of paint-on resin to crown and bridge composites after soaking in water and thermal-cycling. Three shades of paint-on resin were coated on three kinds of crown and bridge composite under four surface treatment conditions (a combination of sandblaster and pretreatment liquid). These specimens were soaked in water at 37°C for 1 day, 1 month, and 1 year, and at 4°C and 60°C alternatively for 1-minute periods for 10,000 cycles by thermal-cycling machine. The adhesive strengths were obtained by shear test. There were no significant differences among the adhesive strengths of three shades of paint-on resin to three composites after storage (p>0.05). The adhesive strengths to composites with sandblasting showed higher values than those without it (p<0.01).

Key words: Adhesive strength, Crown and bridge composite, Paint-on resin

INTRODUCTION

Crown and bridge (crown & bridge) composites are used as esthetic restorative materials for jacket crowns and facing crowns. Therefore, an excellent color match between the crown & bridge composite and the tooth is vital to esthetic success. However, color tone of the crown & bridge composite can be changed by various factors. It can be changed by conditions in the oral cavity, due to consumption of foods and drinks, tobacco, etc. It can also be changed by the finished surface conditions of the crown & bridge composite. In particular, the thin resin layer makes it difficult to reproduce a matching and stable color tone for the crown & bridge composite. This is because the thin resin layer reflects the background color. Recently, a light-curing paint-on resin — which has the selling points of abrasion resistance and color tone stability — has been on the market to solve these problems. This material is first applied to the surface of the crown & bridge composite or artificial plastic tooth. Then, the following are carried out: last shade adjustment, correction, characterization. To find out more about this resin, we have been studying its resistance to tooth-brush abrasion and its optical properties.

The aim of this investigation was to evaluate the adhesive strength and durability of paint-on resin. So the adhesive strengths between three kinds of crown & bridge composite and three shades of paint-on resin were measured under various surface treatment conditions and storing conditions of test specimens.

MATERIALS AND METHODS

Materials

Three kinds of commercial crown & bridge composite and three shades of paint-on resin were used in this investigation, as shown in Table 1. A pretreatment agent (C & B liquid, Heraeus Kulzer Japan, Osaka, Japan) was used to process the test specimens according to the surface treatment conditions given in Table 2.

Test specimens preparation

The dentin or body paste of the crown & bridge composites shown in Table 1 was filled into the hole of a stainless steel holder (20 × 30 × 1 mm with ø10 mm hole). Subsequently, the enamel or incisal paste
Table 2 Four surface treatment conditions of cured enamel or incisal crown and bridge composites before painting the paint-on resin

<table>
<thead>
<tr>
<th>Surface treatment</th>
<th>Code</th>
</tr>
</thead>
<tbody>
<tr>
<td>-</td>
<td>NT</td>
</tr>
<tr>
<td>Sandblaster treatment</td>
<td>SB</td>
</tr>
<tr>
<td>Pretreatment agent (c &amp; b liquid)</td>
<td>PA</td>
</tr>
<tr>
<td>Sandblaster treatment + Pretreatment agent (c &amp; b liquid)</td>
<td>SP</td>
</tr>
</tbody>
</table>

Table 3 Three conditions of preparing the test specimens for adhesive test

<table>
<thead>
<tr>
<th>Condition</th>
<th>Composite</th>
<th>Paint-on resin</th>
<th>Surface treatment</th>
<th>Conservation</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>DS</td>
<td>MA, MO, MT</td>
<td>SP</td>
<td>In water at 37°C for 1 month (1M)</td>
</tr>
<tr>
<td>2</td>
<td>DS</td>
<td>MA</td>
<td>NT, SB, PA, SP</td>
<td>In water at 37°C for 1 month (1M), and in water at 37°C for 3 months before surface treatment and for 1M after all curing processes (3/1M)</td>
</tr>
<tr>
<td>3</td>
<td>DS, SO, EP</td>
<td>MA</td>
<td>SP</td>
<td>In water at 37°C for 1 month (1M), 1 year (1Y), and thermal-cycled in water between 4°C and 60°C for 10,000 cycles (TC)</td>
</tr>
</tbody>
</table>

Adhesive strength measurement

After storage or thermal-cycling, the shear bond strengths (adhesive strengths) of test specimens were measured with a universal testing machine (TCM-1kNNB, Minebea, Nagano, Japan) at a cross-head speed of 0.5 mm/min. A stainless steel wire with 0.43-mm square was connected to a load cell and the stainless steel holder attached to the jig (Fig. 2).

Fillers and monomers analyses

Each paint-on resin was put into a porcelain crucible and weighed ($W_a$) with an analytical balance. Then it was fired in electric furnace at 700°C for 6 hours, and the inorganic filler was weighed ($W_e$) after it had cooled. The inorganic filler content ($W_e/W_a$) was calculated and averaged for three samples.
After weighing, the inorganic filler was fixed in a specimen holder and subjected to X-ray diffraction (XRD, Rint 2000, Rigaku, Tokyo, Japan). The diffraction conditions were CuKα, 40 kV-30 mA, at a scanning rate of 2°/min.

The paint-on resin paste was uniformly applied between two KBr plates and measured with Fourier transform infrared spectrometer (FTIR, FT/IR-460, JASCO, Tokyo, Japan).

**Statistical analysis**

Seven test specimens were used for the adhesive strength measurement. Two-way analysis of variance (ANOVA) was conducted with storing conditions and materials as the independent variables and adhesive strengths as dependent variables. The test specimens stored in water at 37°C for 1 day (1D) was used as control, and Tukey’s test was carried out to analyze the difference.

**RESULTS**

The adhesive strengths measured under Condition 1 are shown in Fig. 3. The test specimens used three shades of paint-on resin (MA, MO, and MT), and their adhesive strengths ranged from 8.2 to 9.1 MPa. There were no statistical differences among the values of the paint-on resins, and between the values of 1D- and 1M-specimens (p>0.05).

The adhesive strengths measured under Condition 2 are shown in Fig. 4. The adhesive strengths of 1D- and 1M-specimens under four surface treatment conditions (NT, SB, PA, and SP) ranged from 8.0 to 9.1 MPa, and no statistical differences were observed among the values of four surface treatments (p>0.05). The adhesive strengths of 3/1M-specimens in SB and SP (8.3 and 8.4 MPa) were higher than those in NT and PA (4.9 and 6.1 MPa) (p<0.01).

The adhesive strengths measured under Condition 3 are shown in Fig. 5. The 1D-, 1M-, and 1Y-specimens used three kinds of light-curing crown & bridge composite (DS, SO, and EP), and their adhesive strengths ranged from 9.0 to 9.4, 8.9 to 9.6, and 6.0 to 7.1 MPa respectively. No statistical differences were observed among the values of storing conditions (p>0.05). However, the adhesive strengths of EP specimens decreased statistically compared with those of DS and SO specimens (p<0.05). No statistically significant differences were found between 1D-, 1M-, and 1Y-specimens and thermal-cycled specimens (TC) (p>0.05).
The inorganic filler contents of MA, MO, and MT were 37.9%, 39.0%, and 35.9% respectively. The XRD patterns of the inorganic fillers are shown in Fig. 6. The mark (♦) in MA and MO corresponds to ZrO₂ peak, and broad patterns corresponding to amorphous material were observed in all three.

FTIR spectra are shown in Fig. 7. MA, MO, and MT spectra were similar to each other. Since the characteristic absorption of 1533 cm⁻¹ corresponds to δ N-H, and those of 1608 and 1509 cm⁻¹ correspond to benzene, paint-on resins should be composed of UDMA- and Bis-GMA- or Bis-MEPP-type monomers.

DISCUSSION

For the test specimens which used three shades of paint-on resin (MA, MO, and MT), there were no statistical differences among their adhesive strength values (p>0.05). This being so because there was hardly any difference in the filler content and in the filler and monomer compositions. The dental composite resin cures by radical polymerization. Therefore when oxygen exists, free radicals formed from an initiator are scavenged and the non-polymerized monomer remains on the surface. An incremental method using this non-polymerized monomer is widely used for composite resins and crown & bridge composites. Bagis et al.¹² reported that a heat treatment at 50-125°C significantly decreased the amount of non-polymerized monomer remaining in a post-light-cured composite resin. This fact shows that non-polymerized monomer — which is required for polymerization — remains primarily in the surface layer.

In SB, the non-polymerized monomer in the surface layer was removed with a sandblaster. As a result, there were no statistical differences in adhesive strength between SB and NT specimens (Fig. 4). In the same fig., the adhesive strengths of 3/1M-specimens in NT and PA significantly decreased compared with those of 1D- and 1M-specimens in NT and PA. These specimens exhibited interface fracture between paint-on resin and the crown & bridge composite (Fig. 8, left). When specimens are stored in water, little by little the monomer near the surface dissolves into the water, and the moisture penetrates into the void. It is well known that absorbed water in the resin matrix obstructs monomer infiltration, hence lowering the adhesive property. The sandblaster treatment is well used to remove the surface layer of water-absorbed resin matrix. As a result, the sandblaster treatment leads to formation of mechanical holding, extension of surface area, and generation of a fresh surface. However, Tachii et al.¹³ mentioned that the rough surface by sandblaster treatment resulted in no adhesive strength increase because the composite paste has high viscosity. In the present study, the increase in adhesive strength was probably obtained due to a sufficient wetting effect — the viscosity of paint-on resin being lower than that of composite resins. These results showed that if the elapsed time following specimen production were short, sufficient adhesive strength can be
obtained without extra labor (such as painting of the c & b liquid).

When comparing among the types of light-curing crown & bridge composite, the adhesive strength in EP was lowered significantly (p<0.05). However, cohesive fracture was sustained in the entire crown & bridge composite (Fig. 8, right). Miyairi et al. measured the adhesive strengths of composite resins and resin cements, and reported that when compared to resin cement, the composite resin became insensitive to stress concentration due to relaxation. Filler content and filler size of the composite resin would influence this factor.

The adhesive strengths of 1D- and 1M-specimens were not significantly different from those of TC-specimens (p>0.05), hence highlighting the excellent durability of the paint-on resins' adhesive property. The 10,000 times thermal-cycling test used in this investigation — which corresponds to two-and-a-half years in the oral cavity by Miyazaki et al. — is a considerably severe test condition. Generally, degradation induced by thermal-cycling is attributed to internal stress caused by the repetitions of expansion and contraction. The internal stress then generates a crack between filler and resin matrix, and soon after water penetrates into the crack. Low filler content and small filler size would delay the degradation of paint-on resin.

The XRD study revealed that MA and MO contained ZrO$_2$ and amorphous material, whereas MT contained amorphous material only (Fig. 6). ZrO$_2$ is widely used as a colorant. Since MA and MO were colored and MT was superior in transparency, ZrO$_2$ was probably added as a colorant in these pastes. Arikawa et al. reported that the optical transparency of MA and MO was lower than that of MT, and that it greatly decreased with increase in thickness. This property is most likely influenced by the diffusion of inorganic filler.

The filler content of paint-on resins ranged from 35.9 to 39.0%. The previous study also reported similar values: 37.8 wt% in MA, 38.8 wt% in MO, and 36.1 wt% in MT. In addition, the three shades of paint-on resin (MA, MO, and MT) showed similar FTIR spectra (Fig. 7) — which meant that the same monomer types were used. These facts thus indicated that no difference in the filler and monomer contents of paint-on resins would affect adhesive strength and physical properties after curing.

This study has proved that there were no significant adhesive strength differences among the three shades of paint-on resin (Fig. 3) and among the three kinds of crown & bridge composite (Fig. 5). In practical clinical service, the surface color of crown & bridge composites post-restoration may be changed by some conditions. Assuming that these conditions exist, the adhesive strengths of the crown & bridge composites stored in water for 3 months were measured before painting the paint-on resin (3/1M). It was shown through this study that by creating a new surface with sandblaster treatment, sufficient adhesive strength was obtained.

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REFERENCES