INTRODUCTION

The advance of adhesive techniques in Dentistry has increased the use of composite resins. Currently, this material is widely used in direct and indirect restorative procedures. Composite resin can be used to make cores on prefabricated posts (1), as a cavity filling material (2), as a dentin sealing material in the resin-coating technique, (3) and also as a luting agent (4).

It is known that during light curing of resin materials, the presence of oxygen does not allow the complete polymerization of the surface, which favors the bonding of new increments of chemically similar materials (5).

In adhesive cementation, there is an interaction between the luting cement and the non-polymerized surface layer of the adhesive applied to the dental substrate (6). In addition, in situations when the tooth is rebuilt with a composite core, an interaction between this material and the luting cement should occur. Although both materials are chemically similar, the suitable bond achieved by the non-polymerized surface layer is harmed because in the composite resin core, the surface is completely polymerized and contaminated by saliva and temporary cement residues (7).

Several techniques have been suggested for removing temporary cement (8-10). Schwartz et al. (10) reported that cleaning with pumice is effective, while, Paul et al. (11) found that pumice is inefficient for surface cleaning.

The bond between resin materials has been extensively investigated (7,12,13). Nevertheless, there is no consensus in the literature about which is the best surface treatment to achieve a good adhesion between contaminated polymerized resins and resin-based luting systems. Therefore, the aim of this study was to assess the influence of surface treatment on the shear bond strength of a composite resin, previously submitted to the application of a temporary cement (TC), to an adhesive luting cement.
application of a temporary cement, and an adhesive luting cement. The null hypothesis was that there is no significant difference in shear bond strength between treatment protocols.

MATERIAL AND METHODS

Eighty-four cylindrical composite specimens (Esthet-X; Dentsply, Petrópolis, RJ, Brazil) were made using a silicon mold with inner diameter of 5 mm and 2 mm high. The resin was inserted in a single increment and a polyester strip was placed over the material. Light curing was done onto the polyester strip for 40 s, using a halogen lamp (Curing Light XL 3000; 3M ESPE, St. Paul, MN, USA) with light intensity of 600 mW/cm².

The cylinders were divided into 6 groups (n=12). G1 (control) did not receive temporary cement or any surface treatment. The composite resin specimens in Groups 2 to 6 received a coat of Temp Bond NE (Kerr Corp., Orange, CA, USA) temporary cement onto its entire surface. The cement was mixed according to the manufacturer's instructions. After waiting 6 min for cement setting, the cylinders were stored at 37°C in relative humidity for 24 h. After this period, the temporary cement was removed with a dentin excavator, followed by rinsing with an air-water spray and air drying. After temporary cement removal, the composite resin cylinders received the following surface treatments: G2 - Cleaning with a cotton pellet containing ethanol for 10 s and drying with jets of compressed air (3); G3 - Rotary brush with pumice applied for 15 s, followed by washing with air-water spray for 30 s and drying with jets of compressed air (8); G4 - Air-abrasion with 50 µm grain alumina (Microetcher, San Ramon, CA, USA) for 10 s (9) followed by washing with air-water spray for 30 s and drying with compressed air streams; G5 - Air-abrasion with 50 µm grain alumina for 10 s (9), followed by rinsing with air-water spray for 30 s, drying with compressed air streams, coating with a layer of adhesive system (ScotchBond Multi Purpose; 3M ESPE) using a disposable microbrush tip (Microbrush Corp.; Grafton, WI, USA), and photoactivation for 20 s (Curing Light XL 3000; 3M ESPE); G6 - Air-abrasion with 50 µm grain alumina for 10 s, followed by rinsing with air-water spray for 30 s, drying with compressed air streams, etching with 37% phosphoric acid (10), and coating with a layer of adhesive system applied and photoactivated as described above.

All blocks were then adapted to a polytetrafluoroethylene matrix (3 mm diameter and 2 mm high), and the composite resin surfaces received a 2 mm increment of Panavia F adhesive luting cement (Kuraray, Kurashiki City, Japan), which was photoactivated for 40 s, according to the manufacturer's recommendations.

The test specimens were stored in distilled water at 37°C for 24 h, and then to a universal test machine (DL-1000; Equipamentos e Sistemas de Ensaio Ltda., São José dos Pinhais, PR, Brazil). A shear force was applied with a load cell of 100 kgf in the composite resin/cement interface with a chisel-shaped rod attached to the universal testing machine at a crosshead speed of 1 mm/min until bonding failure. The chisel had a 3-mm-diameter saddle-shaped end (Fig. 1). The load at fracture was divided by the crosssectional area of the bonding surface (7.07 mm²) to calculate the nominal stress across the specimen at fracture in MPa. All fractured surfaces were examined with a stereomicroscope (Stemi 2000-C; ZEISS, Petrópolis, RJ, Brazil), at ≈25 magnification to determine the failure mode (adhesive or cohesive).

Data obtained were analyzed by ANOVA and Tukey’s multiple-comparison test at a 5% significance level using Minitab statistical software, version 14.12, 2004 (Minitab Inc., State College, PA, USA).
RESULTS

There were significant differences among the groups with different surface treatments ($f=9.71$; $p=0.000$). G5 and G6 presented the highest shear bond strength values and did not differ significantly from G1. G2 did not differ significantly from G1 either (Table 1).

Table 2 shows the failure mode distribution in the groups. G3 presented 100% of adhesive failures while G5 and G6 presented 100% of cohesive failures. All cohesive failures were in the composite resin Esthet-X.

DISCUSSION

Under clinical conditions, contamination by temporary cement can influence the shear bond strength between resinous materials. Thus, effective surface cleaning is necessary. This study assessed the influence of surface treatment of a composite resin, previously submitted to application of a temporary cement, on the shear bond strength of an adhesive luting cement.

In this study, the methodology used to assess bond strength was the shear test as previously used (7,14). It is an acknowledged methodology and is easy to perform. Different devices can be used to perform the shear strength test, such as chisel systems (straight chisel, saddle-shaped chisel), stainless steel tape and piston (15). In the present study, the option was to use a saddle-shaped chisel (Fig. 1) because there was a larger area of contact between the chisel and the specimen during the test, thus with better distribution of forces. One disadvantage of the shear test is the size of the bond area.

According to Della Bona et al. (16), the larger the bond area, the greater the possibility of formation of bubbles and porosities, which will affect the adhesion. In this study, the bond area was 7.07 mm$^2$, whereas microtensile studies have areas close to 1 mm$^2$. The results of this study showed that the lowest shear bond strength values between the composite resin and luting cement were observed in G3 (rotary brush + pumice) and G4 (air-abrasion with 50 µm grain alumina), being significantly lower than the control group (G1), which was not contaminated by temporary cement. Thus, these two treatments were not considered efficient for surface cleaning. With regard to treatment with pumice, Paul and Scherer (11) reported its inefficiency in dentin cleaning after contamination with temporary cement. However, Schwartz et al. (10) found that the rotary brush associated with pumice was effective.

Although air-abrasion with 50 µm grain alumina

<table>
<thead>
<tr>
<th>Composite surface treatment (Group)</th>
<th>Mean (SD)</th>
<th>Homogeneous Sets*</th>
</tr>
</thead>
<tbody>
<tr>
<td>No treatment (G1)</td>
<td>13.343 (3.928)</td>
<td>A C</td>
</tr>
<tr>
<td>Ethanol (G2)</td>
<td>10.188 (3.471)</td>
<td>A B</td>
</tr>
<tr>
<td>Rotary brush + pumice (G3)</td>
<td>8.530 (3.962)</td>
<td>B</td>
</tr>
<tr>
<td>Air-abrasion (G4)</td>
<td>8.628 (3.752)</td>
<td>B</td>
</tr>
<tr>
<td>Air-abrasion + adhesive (G5)</td>
<td>14.785 (3.622)</td>
<td>C</td>
</tr>
<tr>
<td>Air-abrasion + acid etching + adhesive (G6)</td>
<td>15.856 (2.308)</td>
<td>C</td>
</tr>
</tbody>
</table>

* Sets accompanied by the same letters presented no significant differences ($p > 0.05$).

<table>
<thead>
<tr>
<th>Composite surface treatment</th>
<th>Adhesive (%)</th>
<th>Cohesive (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>No treatment (G1)</td>
<td>75</td>
<td>25</td>
</tr>
<tr>
<td>Ethanol (G2)</td>
<td>58.3</td>
<td>41.7</td>
</tr>
<tr>
<td>Rotary brush + pumice (G3)</td>
<td>100</td>
<td>--</td>
</tr>
<tr>
<td>Air-abrasion (G4)</td>
<td>83.3</td>
<td>16.7</td>
</tr>
<tr>
<td>Air-abrasion + adhesive (G5)</td>
<td>--</td>
<td>100</td>
</tr>
<tr>
<td>Air-abrasion + acid etching + adhesive (G6)</td>
<td>--</td>
<td>100</td>
</tr>
</tbody>
</table>
(G4) produced low shear bond strength values, its association with the adhesive system (G5 and G6) increased significantly the bond strength to the adhesive luting cement. According to Padipatvuthikul and Mair (14), there are two explanations for the action of adhesive system in these cases. Firstly, the airborne particle abraded surface contains micro-defects that can be penetrated by the adhesive fluid, resulting in micromechanical retention. Secondly, the solvents present in the adhesive systems may cause an alteration in the surface layer, allowing the monomer of the resinous present in the adhesive systems may cause an alteration in the surface layer, allowing the monomer of the resinous cement to react with the non converted vinyl groups (-C=C) at the subsurface.

As regards the effectiveness of phosphoric acid application before adhesive application, although there was higher shear bond strength in this group (G6), it did not differ significantly from the group without acid application (G5). The action of phosphoric acid at the surface of the composite resin is a cleansing action. With the previous use of air abrasion, this action may not have caused significant effects on the bond strength.

Surface cleaning with ethanol (G3) was shown to be a simple and effective technique, as it did not differ significantly from the control group (G1), although it had lower values than groups G5 and G6. This technique is recommended by some authors (17,18) in the resin coating technique, in which dentin sealing is done soon after preparation, before the stages of molding and making temporary restorations. Ethanol is a solvent of organic material and promote surface cleaning, which facilitates the bond between composite resin and resinous cement (18).

With regard to the bond failures, it was observed that in the groups that presented higher shear bond strength (G5 and G6), all tested specimens had cohesive failures in the composite resin. Della Bona et al. (19) and Della Bona and Van Noort (16) discussed this problem and concluded that shear tests measure the resistance of the adjacent composite rather than the resistance to bonding. However, the adjacent resin is only going to fracture if the shear bond strength at the interface is high. In this study, the fractures occurred at the bond interface in the groups in which the shear bond strength presented lower values (Table 2). In the groups with the highest shear bond strength values (G5 and G6), only cohesive failures occurred, demonstrating that the interface had a good resistance, fracturing the base composite.

Within the limitations of this study, it was concluded that ethanol, air-abrasion associated with adhesive system and air-abrasion associated with the phosphoric acid were effective for cleaning the surface of composite resin from temporary cement residues, promoting an effective bond of composite to an adhesive luting cement.

RESUMO

A influência do tratamento de superfície sobre a resistência adesiva entre uma resina composta (RC), previamente submetida à aplicação de um cimento temporário (CT), e um cimento resinoso foi avaliada. Oitenta e quatro cilindros de RC (5 mm de diâmetro e 3 mm de altura) foram confeccionados e incluídos em resina acrílica. Os conjuntos foram divididos em 6 grupos (G1 a G6) (n=12). Os grupos de 2 a 6 receberam uma camada do CT. Após 24 h, o CT foi removido e as superfícies de RC receberam os seguintes tratamentos: G2: limpeza com etanol; G3: limpeza com escova rotatória e pedra pomes; G4: jateamento; G5: jateamento e aplicação de adesivo; G6: jateamento, condicionamento ácido e adesivo. O G1 (controle) não recebeu CT ou nenhum tratamento de superfície. Os conjuntos foram adaptados à matriz e receberam incremento de cimento resinoso. Os corpos de prova foram submetidos ao ensaio de cisalhamento. Os testes ANOVA e Tukey (p=0,001), demonstraram que os grupos G3 (8,53 MPa) e G4 (8,63 MPa) diferiram estatisticamente do grupo G1 (13,34 MPa). As maiores valores médios de resistência ao cisalhamento foram encontradas nos grupos G5 (14,78 MPa) e G6 (15,86 MPa). O jateamento da superfície da resina composta associado a um sistema adesivo, independente do pré-tratamento com o ácido fosfórico, proporcionou uma efetiva união ao cimento resinoso.

REFERÊNCIAS


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