Influence of indirect reinforcements on the flexural strength of a thermally activated acrylic resin used for complete dentures

Objective: This research aimed to verify the performance of thermally activated acrylic resin (TAAR) combined with a mix of glass and aramid fibers and/or composite resin of indirect use by a tree point bending test. Material and Methods: Ten samples, with 65 x 10 x 2.5 mm, were prepared for each group (n = 10): CO, control of only TAAR; CR, in which an 60 mm indirect composite resin was polymerized together with the acrylic resin during the thermo-polymerization cycle; SS, in which ceramic glass mixed with aramid fibers cut 60 mm in length were incorporated into the samples; and SC, in which the same fibers were incorporated with addiction of the indirect composite resin. A three-point flexural strength test was performed with a universal testing machine with a load of 50Kgf at a speed of 5 mm/min in the center of the samples supported by a suitable device. The reinforced face was placed to the tensile side. The statistical one-way ANOVA and Tukey tests were made with a significance level of 95%. Results: The mean value for the CO was 60,27 ± 24,18 MPa, for CR it was 38,39±12,75 MPa, for SS it was 79,97±12,75 Mpa and for CS it was 32,40±9,05 MPa. Conclusion: The use of glass and aramid fibers at the base of a TAAR increased the three-point flexural strength, however when indirect composite resin was incorporated, a significant drop of this mechanical property was observed.

Keywords: Acrylic resin; Complete denture; Composite resin.

RESUMO

Objetivo: Esta pesquisa teve como objetivo verificar o desempenho de uma resina acrílica ativada térmicamente (RAAT) combinada com uma mistura de fibras de vidro e aramida e / ou resina composta de uso indireto por um teste de flexão de três pontos. Material e Métodos: Foram preparadas dez amostras, com 65 x 10 x 2,5 mm, para cada grupo (n = 10): CO, controle com apenas RAAT; CR, em que uma resina composta indireta de 60 mm foi polimerizada em conjunto com a resina acrílica durante o ciclo de termopolimerização; SS, em que o vidro cerâmico misturado com fibras de aramida cortadas com 60 mm de comprimento foi incorporado nas amostras; e SC, nas quais as mesmas fibras foram incorporadas em conjunto com a resina composta indireta. O teste de resistência à flexão de três pontos foi realizado em uma máquina de teste universal com uma carga de 50 KgF a uma velocidade de 5 mm/min no centro das amostras suportadas por um dispositivo adequado. A face reforçada foi colocado no lado da tração. Os testes estatísticos one-way ANOVA e Tukey foram feitos com um nível de significância de 95%. Resultados: O valor para o CO foi de 60,27 ± 24,18 MPa, para CR foi de 38,39 ± 12,75 MPa, para SS foi de 79,97 ± 12,75 Mpa e para CS foi de 32,40 ± 9,05 MPA. Conclusão: O uso de fibras de vidro e aramida na base de uma RAAT aumentou a resistência à flexão de três pontos, no entanto, quando a resina composta indireta foi incorporada, observou-se uma queda significativa dessa propriedade mecânica.

PALAVRAS-CHAVE

Resina acrílica; Prótese total; Resina composta.
INTRODUCTION

Thermally activated acrylic resin (TAAR), composed by polymethyl methacrylate, is the main material used in edentulous patients rehabilitation. Its use is justified not only for the simplicity of the technique, but also good resistance, durability, biocompatibility, cost-benefit ratio, dimensional and chemical stability. However, it still does not fulfill all the necessary requirements of a rehabilitation material [1-5]. Complete dentures execution with TAAR are done at the cost of clinical and laboratory procedures [1-3].

However, in cases of implant installation to retain and/or give support for complete dentures [6], TAAR present precarious mechanical properties and can frequently fracture [1-3,5,7-10]. This can occur as a result of occlusal disharmony, overloading, fatigue and impact caused by accidents [11-12]. Specially during the osseointegration phase of a protocol type prosthesis, which usually presents cantilever [13], provisional prosthesis made entirely of TAAR may be used over the implants [6]. Thus, the contraindication for the patient is the reduced longevity of such prosthesis.

Therefore, for mucus or implant supported complete dentures, there is the need for TAAR reinforcement, which could result in better mechanical performance, mainly in terms of fracture resistance, allowing greater longevity for these rehabilitations [1,4,8-9,12].

Initially the use of metallic alloys as reinforcement was made in the form of molten metal or stainless-steel wires [14-15]. Although these metallic reinforcements showed a good performance regarding the mechanical aspect [16-18], they presented big interfacial stress and aesthetic disadvantages [3,11]. Alternatively, various reinforcing fibers, such as glass [13,19-20], aramid, carbon and polyethylene are commercially available [18-20]. Carbon fibers were useful in strengthening the polymers process but they are not used due to the easy lateral fibers spreading during the processing phase and their low aesthetic quality [3,21], which is also the main disadvantage of aramid fibers [3]. Contrary, silanized glass fibers show better adhesion to the polymeric matrix and superior aesthetics when correctly inserted in the base of complete dentures [3,21].

Polymers are cured by the polymerization reaction and one of the consequences of this reaction is the contraction that occurs at different intensities regarding different types of resins. Considering a possible improvement in the compatibility of silanized glass fibers with the composite resin and considering the difference in the contraction degree of the composite resin compared to the acrylic one, it was supposed the creation of tensions that could improve the final resistance of this aggregation [22].

Considering this, this research aimed to verify the flexural strength of TAAR reinforced with different fibers and composite resin. The null hypothesis tested was that there would be no difference in flexural strength between the control group, without reinforcement, and the different experimental groups, reinforced either with composite resin and/or aramid and glass fibers.

MATERIAL & METHODS

Preparation of the samples:

Three stainless steel patterns with dimensions of 67 x 12.6 x 2.55 mm were used as standard to make a silicone matrix in flasks. Initially, the flasks were filled with type III gypsum (Polident, Agudos, SP, Brazil) in the proportion specified by the manufacturer's instructions (Figure 1).

After the separating agent dried, a laboratory silicone (Rohhorsil, Bluestar Silicons, France) was used around the three standards with a 20 ml syringe, leaving only one side of the patterns without covering. After 30 minutes (silicone polymerization time), the counter part of the flask was positioned and
filled with type III gypsum in small portions, using a vibrator. After the flasks were properly prepared, they were pressed for 1 hour. After the flasks re-opening, the stainless steel standards were removed and a thin layer of the gypsum surface separation agent was re-applied and the same process was performed with smaller additional metal dies to standardize the space for the reinforcement.

Figure 1 - A) insertion of the patterns on the gypsum with a 10mm space between them; B) wall construction in condensation silicone (Vipisil; VIP, Pirassununga, São Paulo, Brazil); C) flasks prior to removal of the metallic pattern; D) flasks after removal of the metallic pattern; E) additional metal spacer pattern to standardize the reinforcement space; F) final matrix obtained for the samples.
After that, ten samples were prepared for each group (n = 10): CO, control TAAR (Lucitone 550, Dentsply Sirona, São Paulo, SP, Brazil); CR, in which an indirect composite resin (SR Chromasit, Ivoclar Vivadent, Schaan, Liechtenstein) was prepared with 60 mm in a Centrix syringe and polymerized together with the acrylic resin during the thermopolymerization cycle (Figure 2); SS, in which ceramic glass mixed with aramid fibers (Superfiber, Superdont, Rio de Janeiro, RJ, Brazil) cut 60 mm in length were incorporated into the samples; and SC, in which the same fibers were incorporated to the samples with addition of CR (Table 1).

Table 1- Groups, reinforcements types and materials used to make the samples

<table>
<thead>
<tr>
<th>GROUPS</th>
<th>REINFORCEMENTS TYPES</th>
<th>MATERIALS</th>
</tr>
</thead>
<tbody>
<tr>
<td>CO</td>
<td>None</td>
<td>TAAR</td>
</tr>
<tr>
<td>CR</td>
<td>Indirect composite resin</td>
<td>TAAR + CR</td>
</tr>
<tr>
<td>SS</td>
<td>Aramid and glass fibers</td>
<td>TAAR + fibers</td>
</tr>
<tr>
<td>SC</td>
<td>Indirect resin, aramid and glass fibers</td>
<td>TAAR + CR + fibers</td>
</tr>
</tbody>
</table>

TAAR was manipulated as recommended by the manufacturer. The flasks were slowly and gradually pressed in a hydraulic press until establishing 1000 Kgf, and then maintained for 30 minutes. After the pressing time, the flasks were transferred to an individual press and brought to the polymerization process. The polymerization process was undertaken using a short cycle, as recommended by the manufacturer; the flasks were positioned in cool water and brought to 100°C, maintaining for 45 minutes at that temperature. Once the process finished, a two-hour period was waited for cooling the flasks at room temperature.

The final dimensions of the samples were 65 mm x 10 mm x 2.5 mm [23], using two stainless steel devices that served as guides to match the length, width and thickness of the specimens. Specimens were wet grounded to the final dimensions using sandpaper in respective granulation of 180, 320 and 600. After finishing, specimens were stored in a recipient with distilled water inside a kiln at 37°C for 48 ± 2 hours.

**Three-point bending test**

A three-point flexural strength test was performed using a universal testing machine (EMIC, São José dos Pinhais, Paraná, Brazil) with a load of 50 kgf at a cross-head speed of 5 mm/min at the center of the samples supported by a suitable device. The reinforced face was placed to the tensile side (Figure 3).

The fracture load was recorded in Newtons (N) and the flexural strength (FS) was calculated in MPa with the following equation $FS = \frac{3FD}{2LH^2}$, where $F$ is the maximum load supported by the specimen before its fail, which was determined in Newton (N); $D$ is the span distance (50mm); $L$ is the length of the specimens (10mm); $H$ is the thickness of the specimens (2.5mm). The dimensions were checked with the aid of a digital pachymeter (Mitutoyo, Suzano, São Paulo, Brazil).

Data were tabulated and submitted to the normality test, which indicated the possibility of performing a one-way ANOVA and Tukey tests with a significance level of 95%.
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Miranda JS et al.

Braz Dent Sci 2018 Apr/Jun;21(2)

RESULTS

Through the ANOVA statistical test, it was possible to verify that there was a significant statistical difference between the groups (p = 0.01). Tukey test indicated that the application of the indirect composite resin reduced the flexural strength of the samples, while the use of glass and aramid fibers was beneficial to the resistance. Therefore, the group that presented the greatest flexural strength was SS, followed by CO, CH and CS, respectively (Table 2).

Table 2 - Three-point flexural strength data of the control and experimental groups

<table>
<thead>
<tr>
<th>GROUPS</th>
<th>MEAN ± STANDARD DEVIATION (MPa)</th>
</tr>
</thead>
<tbody>
<tr>
<td>CO</td>
<td>60.27±24.18 B</td>
</tr>
<tr>
<td>CR</td>
<td>38.39±12.75 A</td>
</tr>
<tr>
<td>SS</td>
<td>79.97±12.75 C</td>
</tr>
<tr>
<td>CS</td>
<td>32.40±9.05 A</td>
</tr>
</tbody>
</table>

DISCUSSION

In this research, glass and aramid fibers and/or indirect composite resin were added as an attempt to reinforce TAAR. To evaluate these treatments, a three-point bending test was performed. This test is the most used to evaluate acrylic resin resistance [1-3,7,10-12,14,21,22], because the three-point test simulates a bending failure caused by deformation of the base of the complete prosthesis [3,12,14], being preferable to torsion, compression or shear tests [7].

Trying to simulate the conditions of a complete prosthesis, the thickness of the experimental bars was 2.5 mm, which is the dimension also used in others researches [7,11-12] and it is close to the one clinically used in a prosthesis base.

Insertion sites of the reinforcement material in the TAAR can be in the upper face of the prosthesis base [1], in the medium [1,10] or in the lower region [1,9]. Agha, Flinton and Vaidyanathan [1] evaluated the strength influence of the glass fiber reinforcement on TAAR bars and reported that the placement of the fibers closer to the lower region was more favorable to this material resistance, fact that guided the preparation of the specimens in our research. The beneficial effects of fiber reinforcement on the tensile side [9], face subjected to increased stress, can be attributed to the ability of the fiber to absorb more energy by stretching of the fibers, before the occurrence of the fracture. As a result, the stress required for the initiation of crack growth and its subsequent propagation is postponed until a greater stress is required or the fracture to be properly initiated [1].

Analyzing the obtained results, the null hypothesis was rejected, since all the experimental groups obtained different results when compared to the control group. In groups which indirect resin was inserted, a decrease in resistance was found. Although da Silva et al. [22] reported the opposite, this decrease in resistance values can be justified by the fact that the elastic modulus of the composite resin is higher than the acrylic resin [24,25], not allowing the bars with this composite to undergo so much plastic deformation compared with pure acrylics or fibers reinforced bars and fractured with a lower load.

Another factor to be considered is that the composite resin used in this research did not promote an efficient penetration into the surface of the acrylic resin, resulting in a structure with faults, in which the fractures could be originated.
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Miranda JS et al.
Braz Dent Sci 2018 Apr/Jun;21(2)

References


Conclusion

The use of glass and aramid fibers at the base of a thermally activated acrylic resin increased the three-point flexural strength, however when indirect composite resin was incorporated into the specimens a significant drop of this mechanical property was observed.
Influence of indirect reinforcements on the flexural strength of a thermally activated acrylic resin used for complete dentures

Miranda JS et al.


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