**In vitro** effects of erosive challenge on the surface properties of sealants

Isabella Cavalcante Medeiros¹, Bruna Palmeira Costa¹, Brenna Louise Cavalcanti Gondim¹, Hugo Lemes Carlo¹, Rogério Lacerda dos Santos¹, Fabiola Galbiatti de Carvalho¹

¹Universidade Federal da Paraíba - UFPB, Health Science Center, Department of Clinical and Social Dentistry, João Pessoa, PB, Brazil

**Abstract**

**Aim:** To assess in vitro the surface roughness (Ra), Vickers hardness (VHN) and surface morphology of resin and glass ionomer materials used for sealants after dynamic erosive challenge.

**Methods:** Twenty specimens of each material were prepared and divided into experimental (erosive challenge) and control groups (n=10): Protect Riva (SDI), Opallis Flow (3M ESPE), Fluroshield (Dentsply), Filtek Z350 XT Flow (3M ESPE). The erosive challenge was performed 4 times per day (90 s) in cola drink and for 2 h in artificial saliva for 7 days. The control specimens were maintained in artificial saliva. Ra and VHN readings were performed before and after erosion. The percentage of hardness loss (%VHN) was obtained after erosion. The surface morphology was evaluated by scanning electron microscopy (SEM). The data were analyzed by ANOVA, Tukey and paired t tests (α=0.05).

**Results:** After erosion and saliva immersion, there was an increase in Ra values for all groups and Riva group showed the highest Ra values. After erosive challenge, Riva and Filtek groups showed significant decrease in VHN values, but Filtek group showed the greatest %VHN. For all groups there was inorganic particle protrusion and matrix degradation after erosion visualized by SEM images.

**Conclusions:** Erosive challenge affected the surface properties of all materials used as sealants, particularly in the Riva and Filtek groups.

**Keywords:** tooth erosion; composite resins; glass ionomer cements; hardness tests.

**Introduction**

Dental erosion is defined as the loss of tooth substance by the chemical process of acid exposure and dissolution, involving no bacterial plaque acid. The acids in food and drink are considered the major etiological factors responsible for erosive lesions in enamel. There is evidence that the prevalence of erosion is increasing because of the high consumption of soft drinks. The most important aspects in the treatment of patients with dental erosion are diagnosis and prevention of lesion progression to limit additional destruction of tooth tissue. Elimination of the acid source and preventive approaches using fluoride compounds are indicated to control the tooth erosion or the dental tissue softening caused by acidic solutions.

The effects of dentin-bonding agents and restorative materials used to seal dental structures have also been advocated to control or prevent the development of erosion lesions. It was shown that resin-based bonding agents could protect against erosion and abrasion in situ. Wegehaupt et al. (2012) demonstrated that resin-based materials were also able to reduce the erosive demineralization of bovine enamel after immersion in hydrochloric and citric acids over consecutive days. However, to simulate what occurs in the oral cavity, in vitro studies used...
Specimen preparation

Material and methods

Specimen preparation

Four materials used as sealants were investigated in this study: three resin-based sealants – Filtek Z350 XT Flow (3M/ESPE, St. Paul, MN, USA), Fluroshield (Dentsply, Rio de Janeiro, RJ, Brazil) and Opallis Flow (FGM, Joinville, SC, Brazil); and one glass ionomer sealant – Riva Protect (SDI, Bayswater, Victoria, Australia). The compositions of the evaluated materials are in Table 1.

Twenty specimens of each material were fabricated using silicone molds (4 mm diameter x 2 mm high), according to the manufacturers’ instructions. The capsules of Riva Protect were triturated for 10 s in an amalgamator (Ultramat 2, SDI, Bayswater, Victoria, Australia). Then, the material was inserted into the matrix with the Riva applicator. The specimens were covered with acetate strips (Probem Ltda, Catanduva, São Paulo, Brazil) and were pressed flat with a glass slide for 5 min to obtain a smooth surface. The resin-based materials were inserted into the matrix and were covered with acetate strips and polymerized for 20 s with a LED curing light (1200 mW/cm² – Radii Cal; SDI). The specimens were maintained in relative humidity for 24 h, before the baseline roughness and microhardness measurements were obtained as described above. Specimens of each material were divided in two groups (n = 10): erosion (erosive challenge) and control (artificial saliva immersion) groups.

Dynamic erosive pH-cycling challenge

The specimens were immersed in cola drink (Coca-Cola®, SP, Brazil - pH 2.3) at room temperature in individual containers (10 mL/specimen) for 90 s four times/day. Subsequently, the specimens were rinsed thoroughly with deionized water and were immersed in artificial saliva with a pH of 7.0 (10 mL/block). This erosive challenge was repeated for 7 days. The cola drink and artificial saliva were changed after every cycle. During the acidic cycles, the samples were kept in hermetically sealed containers to prevent the loss of carbonation from the cola drink. The specimens in the control group were immersed in artificial saliva for 7 days. The artificial saliva was changed every day.

Surface roughness measurements

At the end of the erosive challenge, the specimens were ultrasonically washed for 10 min and dried with absorbent paper. They were then fitted to a surface roughness-measuring instrument (TR200, Digimess, São Paulo, SP, Brazil). In each specimen, three successive measurements in the central area

<table>
<thead>
<tr>
<th>Sealant/Batch number</th>
<th>Type</th>
<th>Compositions</th>
</tr>
</thead>
<tbody>
<tr>
<td>Riva Protect (SDI, Bayswater, Victoria, Australia, F1111031)</td>
<td>Glass ionomer</td>
<td>Capsule: Compartment 1 (powder): Fluoro aluminosilicate glass (90% load size not specified) and polyacrylic acid (10%); Compartment 2 (liquid): polyacrylic acid (25%), tetraric acid (10%) and balanced ingredient (non-hazardous) (65%)</td>
</tr>
<tr>
<td>Fluroshield (Dentsply, Rio de Janeiro, RJ, Brazil, 9462229G)</td>
<td>Resin</td>
<td>UED-Bis-GMA (&lt;40%), barium aluminoborosilicate glass (30%), polymerizable dimethacrylate resin (&lt;10%), Bis-GMA (&lt;5%), sodium fluoride (&lt;5%), dipentaerythritol hexa-acrylate phosphate (&lt;5%), titanium dioxide (&lt;3%), amorphous silica (&lt;2% load size not specified)</td>
</tr>
<tr>
<td>Filtek Z350 XT (3MESPE, St. Paul, MN, USA, N085003)</td>
<td>Resin</td>
<td>Silane-treated ceramic (52-60%), Bis-GMA (10-15%), TEGDMA (10-15%), Bis-EMA (1-5%), silane-treated silica (3-11%), silane-treated zirconium oxide (3-11% mean load size 0.01-6 µm), functionalized dimethacrylate polymer (1-5%)</td>
</tr>
<tr>
<td>Opallis Flow (FGM, Joinville, SC, Brazil, 040612)</td>
<td>Resin</td>
<td>UDMA (5-10%), TEGDMA (5-10%), Bis-EMA (5-10%), silanized inorganic filler, Ba-Al-Si micro-particles and SiO₂ in nanoparticles (0.05 and 5.0 µm) (~72%)</td>
</tr>
</tbody>
</table>

*Bis-GMA: bisphenol-A-glycidyl methacrylate; TEGDMA: triethylene glycol dimethacrylate; UED-Bis-GMA: urethane-modified Bis-GMA dimethacrylate; Bis-EMA: etoxylated bisphenol-A-diglycidyl methacrylate
in different directions were performed by the same examiner, and the mean surface roughness values (Ra) were obtained and expressed in micrometers. The roughness test was performed at baseline and 24 h after erosive challenge.

Surface Vickers microhardness

The microhardness measurements were obtained with a hardness tester (HMV II; Shimadzu Corporation, Kyoto, Japan), using a Vickers indenter (VHN) and a 200 g load, with a 15 s dwell time. Five indentations were made in each specimen, at least 50 µm apart, and the mean VHN value was obtained. In addition, the percentage of microhardness loss (%VHN) was calculated using the following formula:

\[ \%VHN = \frac{100 \left( VHN_{I} - VHN_{F} \right)}{VHN_{I}} \]

where VHN(I) is the average of the initial (baseline) microhardness measurements, and VHN(F) is the average of the final (after erosive challenge) microhardness values.

Scanning electron microscopy (SEM)

Three representative specimens from each group were mounted on aluminum stubs and sputter-coated with gold in vacuum (Balzers-SCD 050 Sputter Coater, Balzers, Liechtenstein). Three extra specimens were prepared for baseline (without saliva or coca immersion) evaluation. A LEO 1430 scanning electron microscope (Zeiss Inc., Thornwood, NY, USA) was used for SEM analyses. Analyses were performed at 1500 X and 2000 X magnification before and after the erosive challenge and immersion in artificial saliva.

Statistical analysis

The data were analyzed using GraphPad Instat software, version 2.0 (GraphPad Software, La Jolla, CA, USA) at a significance level \( \alpha = 0.05 \). The sample size was calculated considering the minimum difference between the average of treatments (mean ± standard deviation) of 0.5 ± 0.05 µm for roughness testing and 13.0 ± 3.0 VHN for microhardness testing. With a significance level of 0.05 and a power of 95%, a minimum of four specimens per group was required. All tested variables satisfied the assumptions of normal distribution, therefore two-way ANOVA and Tukey’s test were performed for statistical comparisons of Ra and VHN measurements among the sealants. Student’s paired t-test was used compare Ra and VHN measurements before and after erosive challenge for the same sealant. Unpaired t-test was used to compare Ra and VHN values after erosive challenge and artificial saliva immersion for each sealant.

Results

The results of the roughness and microhardness tests are in Tables 2 and 3, respectively. When the Ra values between baseline and post-treatment (after erosion or after saliva) were compared, there was a statistically significant increase in Ra values in all the groups (\( p=0.001 \)), except for the Fluroshield and Filtek groups, which did not show significant differences in Ra values after artificial saliva immersion (\( p=0.06 \) and \( p=0.08 \), respectively) (Table 2). When the comparison was made among sealants after erosion
and after saliva immersion, the Riva group showed the highest Ra values (p=0.001) and there were no significant differences among other groups after both treatments (p=0.07) (Table 2).

After erosive challenge, the Riva and Filtek groups showed significant decrease in VHN values (p=0.03 and p=0.001, respectively), but after saliva immersion, there were no significant VHN alterations in any group (Table 3). After erosion, when the %VHN values were compared, the Filtek group had the greatest value (p=0.001) and Riva, Fluroshield and Opallis groups did not show significant differences among them (p=0.08). However, after artificial saliva immersion, there were no significant differences in %VHN among the groups (p=0.10) (Table 3).

Tables 4 and 5 show the comparison of Ra and VHN values of each material between “after erosion” and “after saliva immersion” treatments, respectively. For all the materials, the erosion treatment showed significantly higher Ra values when compared to saliva immersion treatment, except for the Opallis group (p=0.08) (Table 4). For VHN measurements, Riva and Filtek groups showed higher VHN values after saliva immersion compared to after erosion treatment (p=0.03 and p=0.001, respectively) (Table 5).

Figures 1, 2, 3 and 4 show the SEM images at baseline, after erosive challenge and after saliva immersion. For all the groups, after the erosive challenge, a protrusion of the inorganic particles and degradation of the ionomer or resin matrix (Figures 1B, 2B, 3B and 4B) was observed. The Riva group had greater surface degradation than the other groups, showing a porous surface and matrix deterioration (Figure 1B). The cracks visualized on ionomer sealant images (Figure 1) were artifacts caused by vacuum during sample preparation. After artificial saliva immersion, slight surface degradation was observed in the Riva and Fluroshield groups (Figures 1C and 2C), but the other groups did not show morphological differences between baseline and the treatments. Furthermore, the Opallis group showed the smallest surface morphological changes among baseline, erosive challenge and artificial saliva immersion (Figure 4).

### Discussion

Resin and ionomeric materials can be used in clinical practice to seal tooth structures and to prevent dental erosion. It is known that, during consumption, beverages contact only with the tooth surface and restorative materials for a short time before they are washed away by saliva. In previous studies, the sealant surfaces have usually contacted acidic beverages for prolonged periods of time, or the studies did not include the saliva in their methodologies. The

**Table 4** - Surface roughness (Ra) measurements of sealant materials after erosive challenge and artificial saliva immersion. Values are expressed as mean ± standard deviation.

<table>
<thead>
<tr>
<th>Sealants</th>
<th>Roughness values (Ra) (µm)</th>
<th>After erosion</th>
<th>After saliva</th>
</tr>
</thead>
<tbody>
<tr>
<td>Riva Protect</td>
<td>0.45 ± 0.06</td>
<td>0.35 ± 0.02</td>
<td></td>
</tr>
<tr>
<td>Opallis Flow</td>
<td>0.07 ± 0.01</td>
<td>0.06 ± 0.01</td>
<td></td>
</tr>
<tr>
<td>Filtek Z350 XT</td>
<td>0.10 ± 0.02</td>
<td>0.06 ± 0.01</td>
<td></td>
</tr>
<tr>
<td>Fluroshield</td>
<td>0.13 ± 0.02</td>
<td>0.08 ± 0.01</td>
<td></td>
</tr>
</tbody>
</table>

* The same uppercase letters indicate that there was no significant difference in Ra values between the erosion and saliva treatments of each sealant material (unpaired t test, p > 0.05).

**Table 5** - Microhardness (VHN) measurements of sealant materials after erosive challenge and artificial saliva immersion. Values are expressed as mean ± standard deviation.

<table>
<thead>
<tr>
<th>Sealants</th>
<th>Microhardness values (VHN)</th>
<th>After erosion</th>
<th>After saliva</th>
</tr>
</thead>
<tbody>
<tr>
<td>Riva Protect</td>
<td>59.4 ± 3.3</td>
<td>64.0 ± 2.4</td>
<td></td>
</tr>
<tr>
<td>Opallis Flow</td>
<td>40.2 ± 1.9</td>
<td>39.6 ± 2.1</td>
<td></td>
</tr>
<tr>
<td>Filtek Z350 XT</td>
<td>24.1 ± 0.9</td>
<td>43.1 ± 1.2</td>
<td></td>
</tr>
<tr>
<td>Fluroshield</td>
<td>20.4 ± 2.4</td>
<td>19.9 ± 1.9</td>
<td></td>
</tr>
</tbody>
</table>

* The same uppercase letters indicate that there was no significant difference in VHN values between the erosion and saliva treatments of each sealant material (unpaired t test, p > 0.05).
present study was designed to overcome these limitations of *in vitro* erosion studies, using a dynamic erosive pH-cycling model. This dynamic erosive model simulated the typical consumption of individuals considered to be at risk for dental erosion44, using a beverage (Coca-Cola, pH 2.3) that is widely consumed by the population and that has high erosive potential due to its low pH and fluoride/calcium concentrations5.

The results showed an increase in Ra values for all of the sealants after the erosive challenge (Table 2). Francisconi et al.12 (2008) found wear of approximately 0.3 µm for resin and glass ionomer restorative materials after an erosive challenge similar to that used in the present study. According to these authors, roughness values of materials after erosive challenge of approximately 0.1-0.4 µm could be considered low, like the Ra values in the present study. For resin-based sealant materials, the phosphoric acid found in cola beverage could induce softening of the bisphenol-A-glycidyl methacrylate (Bis-GMA) polymers in resin sealants, which could result from the leaching of diluent agents, such as triethylene glycol (Bis-GMA) polymers in resin sealants, which could result in the organic matrix degradation of the tested materials. In glass ionomer cements, there is microcrack formation in a ionic-crosslinked polyalkenoate matrix, leading to subsequent loss of particle adherence, which could also cause increases in roughness and wear17. Similar to resin-base sealants, acidic beverages could dissolve the siliceous hydrogel layer and the matrix peripheral to the glass particles in ionomeric sealant, causing a rougher surface.

The ionomeric sealant (Riva group) showed the highest Ra values before and after erosion and artificial saliva immersion (Table 2). SEM images also showed greater surface degradation of in the Riva group than in other groups, with a porous surface and higher matrix deterioration (Figure 1B). Because the size of the glass filler particles in ionomeric cements is larger than in resin materials17 (Figure 1), there is less homogeneity between the filler and matrix, thus increasing its surface roughness in both immersion media. Furthermore, in glass ionomer cements, there is microcrack formation in a ionic-crosslinked polyalkenoate matrix, leading to subsequent loss of particle adherence, which could also cause increases in roughness and wear17. Similar to resin-base sealants, acidic beverages could dissolve the siliceous hydrogel layer and the matrix peripheral to the glass particles in ionomeric sealant, causing a rougher surface.

The dissolution of siliceous hydrogel matrix could also explain the significant decrease in VHN values in the Riva group compared to other groups after the erosive challenge (Table 3). The study by Francisconi et al.12 (2008) also showed that glass ionomer cements had greater microhardness losses compared to resin composites, because the acid attack on the resin matrix occurred in a lesser extent than in the siliceous hydrogel matrix. After artificial saliva immersion, there were no significant differences among the sealants in %VHN loss; it seems that the acidic pH of beverages could interfere to a greater extent in the organic matrix degradation of the tested materials. In the present study, the Filtek group showed the greatest %VHN loss, compared to other resin-based sealants after erosive challenge (Table 3).

It is known that inorganic filler particles reduce polymerization shrinkage at the same time as they enhance the mechanical properties of the resin material18. The higher percentage of inorganic filler (72%) and the lower organic content (30%) of Opallis sealant, compared with Filtek Z350 XT (52-60% and 3-11%, respectively) (Table 1) probably resulted in a smaller %VHN loss in the Opallis group19. The higher percentage of inorganic filler and its possible displacement due to matrix degradation could also explain the higher Ra values after saliva immersion for Opallis group. Furthermore, the percentage of ethoxylated bisphenol-A dimethacrylate (Bis-EMA) in the organic matrix of Opallis was higher than in the Filtek group. The decreased flexibility and elimination of the hydroxyl groups from the Bis-GMA monomer to Bis-EMA increased the hydrophobicity of Bis-EMA monomer20. This characteristic reduced the water uptake by the matrix and its plasticization after contact with oral liquids. Thus, the percentage of Bis-EMA monomer may be partially responsible for the biochemical stability of Opallis sealant in aqueous environments, regardless the pH of the solution.. The Opallis group was the only one that showed no significant differences in Ra or VHN values after erosive challenge and immersion in artificial saliva (Tables 4 and 5). After erosion, SEM images also showed a surface with matrix degradation and the protrusion of inorganic filler in the Fluroshield and Filtek groups (Figures 2B and 3B, respectively), whereas the Opallis group showed a homogeneous surface similar to that at baseline (Figures 4A and B). The concentration and particle size of the glass ionomer cement is greater than that of the resin sealants. Thus, it was possible to visualize more easily the matrix degradation and the protrusion of the particles after erosive challenge. A limitation of this study was not showing a higher magnification of SEM images of resin sealants, but even in 1500x magnification it was still possible to verify the same degradation characteristics (protrusion of the particles) after erosion in Figures 2B and 3B related to Fluroshield and Filtek Z350 materials. For Opallis group (Figure 4) it was not possible to verify the degradation on the surface because this material suffered less degradation, as explained before.

Hardness is a physical property possibly related to the degree of conversion and to the amount of filler particles in resin-based materials21. Thus, it is likely that the relatively high degree of conversion of the Bis-GMA/UED-Bis-GMA resin matrix for Fluroshield might have compensated for the small percentage of filler particles, resulting in a smaller %VHN loss than in the Filtek group (Table 3). It is generally accepted that crosslinked polymers are more resistant to degradation and solvent uptake in aqueous environments, whereas linear polymers present more spaces and pathways for molecules to diffuse within their structures and to degrade the material21. These facts may also be the reasons why the Fluroshield group did not show significant differences between VHN values before and after erosive challenge and saliva immersion (Tables 3 and 5). For the other groups, except for Opallis, the erosive challenge caused greater degradation of the surface properties than artificial saliva immersion (Tables 4 and 5). SEM images also showed similarities between the surface morphology after erosion and after saliva immersion for the Fluroshield and Opallis.
groups (Figures 2B, 2C and 4B, 4C).

The oral cavity is the adequate environment for predicting the behavior of dental materials, but in vitro models are very important for providing insight into the fundamental mechanisms of biodegradation\(^1\). Clinically, a thin layer of sealant is applied on tooth structure to prevent against erosion. This study used specimens 2 mm thick, but the analysis was restricted to surface properties and the thickness of specimens did not interfere in the results.

Although sealants materials may degrade with time\(^9\), it may have a role in prevention of tooth erosion and may be a less patient-dependent approach compared to fluoride application, because it does not depend on the patient compliance. In general, the present study showed that that erosive challenge with cola beverage caused changes in the surface properties of sealant materials, and the Opallis group had a better in vitro performance related to surface properties.

The hypothesis tested in the present study was accepted because there were differences among the sealants in microhardness, roughness or morphological surface characteristics after erosive challenge. Longer periods of erosive challenge should be used, and other important properties of the sealants, such as adhesion and microleakage, should be studied under erosive challenge.

Acknowledgements

The authors are grateful to Laisa Daniel Gondim for contributions to the roughness surface analysis and to Professor Severino Jackson Guedes de Lima and Isaque Jerônimo Porto of Rapid Solidification Laboratory/Federal University of Paraíba (LSR/UFPB) for use of SEM.

References