Analysis of the Porosity and Microhardness of Glass Ionomer Cements

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glass ionomer cements (GICs) have good qualities such as biocompatibility, bioactivity, adhesion to dental structures, a coefficient of thermal expansion similar to the dental structure, and fluoride release, and is the dental material with the best biomimetic properties [1, 2].

However, GICs have worse mechanical properties than other restorative materials. To improve these properties, materials such as resin monomers, zinc, fibres, silica, nanocrystals, etc. have been incorporated into the powder and liquid phases of these cements [2 – 4]. Due to the improvements obtained through this process, the clinical use of GICs may be advised in both permanent and primary teeth and in restorations in the anterior and posterior sectors [2, 5]. In fact, GICs are currently the first-choice materials in minimally invasive dentistry for cavity management and are especially used in paediatric dentistry [6].

Resin-modified glass ionomer cements (RMGICs) have better mechanical properties and aesthetics than conventional GICs [7]. In addition, they have higher initial resistance and greater control of working time due to their twin setting reaction (chemical and light) [8].

GICs are commercially available in capsules for mechanical vibrating in an automix syringe and for manual mixing with a paper block spatula. All types of mixing incorporate air, resulting in the formation of pores in the cement, which worsens its mechanical properties, in particular, a decrease in the final resistance of the filling [9 – 11], due to the appearance of microcracks which, when propagated, cause marginal gaps in the restorations [12]. The appearance of pores in GICs may also facilitate greater adhesion of microorganisms on the surface of restorations due to increased roughness [8].

Glass ionomers have differing viscosities. Some studies have associated the failure of restorations using high viscosity glass ionomer cements (HVGICs), mainly used in atraumatic restoration treatments (ART), with a larger number of pores in these ionomers [13, 14].

Currently, there is no effective method of satisfactorily reducing the porosity of GICs. To avoid the formation of pores in GICs, a better understanding of their origins and adverse effects on the mechanical properties of GICs is required. This is done using observational techniques such as Scanning electron microscopy (SEM) and Micro-Computed Tomography (Micro-CT), which permit the number, size, and distribution of pores in the material to be distinguished [11, 15]. The physical-mechanical properties of GICs are characterized by the compressive strength, flexural strength and surface microhardness [16, 17]. Microhardness, defined as the resistance of a material to indentation or penetration [18], may be affected by the porosity.

The objectives of this study were to evaluate the influence of viscosity (high and low) and type of mixing (manual, mechanical and automix syringe) on the porosity...
of GICs and study the relationship between the porosity and microhardness.

The null hypotheses were: (a) the viscosity and type of mixing did not influence the appearance of pores in GICs; (b) the porosity does not influence the micro-hardening of GICs.

2. EXPERIMENTAL DETAILS

2.1. Sample preparation

We used the following GICs and prepared five samples for each study group, according to the manufacturers’ instructions: five HVGICs with mechanical mixing:

<table>
<thead>
<tr>
<th>Glass ionomer cement</th>
<th>Composition</th>
<th>Mode of application</th>
</tr>
</thead>
<tbody>
<tr>
<td>EQUIA® Forte Fil (GC Corp., Tokyo, Japan)</td>
<td>Powder: 95 % strontium fluoro-alumino-silicate glass; 5 % polyacrylic acid Liquid: 40 % aqueous polyacrylic acid</td>
<td>– Apply cavity conditioner (10 sec) or dentin conditioner (20 sec). – Mix the capsule for 10 sec. – Load in GC capsule applicator and apply. – A protective finished varnish (EQUIA Forte Coat) should be applied.</td>
</tr>
<tr>
<td>Ionostar® Plus (Voco GmbH, Cuxhaven, Germany)</td>
<td>Powder: fluoro-alumino-silicate glass, polyacrylic acid, tartaric acid. Liquid: polyacrylic acid solution</td>
<td>– Activate the capsule. Mix the capsule for 10 sec, insert it in an application device, and apply the material directly into the cavity. – A protective finished varnish should be applied.</td>
</tr>
<tr>
<td>3M™Ketac™ Universal Aplicap™ (3M Deutschland GmbH, Neuss, Germany)</td>
<td>Powder: oxide glass chemicals Liquid: water, copolymer of acrylic acid-maleic acid, tartaric acid</td>
<td>– Activate the capsule. – Mix the capsule (10 sec) and insert the capsule into the Aplicap™Applicator. – Apply the glass ionomer cement.</td>
</tr>
<tr>
<td>Riva Light Cure HV® (SDI Limited. Bayswater, Victoria, Australia, Germany)</td>
<td>Liquid: acrylic acid homopolymer; 2-hydroxyethyl methacrylate; dimethacrylate cross-linker; acid monomer; tartaric acid. Powder: glass powder.</td>
<td>– Apply Riva Conditioner for 10 sec and rinse with water. – Activate the capsule and immediately mix in an amalgamator. – Immediately place into the capsule applicator and light cure, each increment 20 sec.</td>
</tr>
<tr>
<td>Riva Silver® (SDI Limited. Bayswater, Victoria, Australia, Germany)</td>
<td>Liquid: acrylic acid homopolymer; tartaric acid. Powder: glass powder; acrylic acid homopolymer; alloy powder.</td>
<td>– Apply Riva Conditioner for 10 seconds and rinse with water. – Remove excess water. Keep moist. – Mix the capsule in an amalgamator OR mix the powder/liquid combination on a mixing pad. Apply Riva Protect to tooth surface. Apply Riva Coat and light cure (20 sec).</td>
</tr>
<tr>
<td>Riva Protect® (SDI Limited. Bayswater, Victoria, Australia, Germany)</td>
<td>Liquid: acrylic acid homopolymer; tartaric acid. Powder: glass powder; acrylic acid homopolymer, calcium amorphous phosphate [Ca3(PO4)2].</td>
<td>– Apply Riva Conditioner for 10 sec and rinse with water. – Mix the capsule in an amalgamator OR mix the powder/liquid combination on a mixing pad. Apply Riva Protect. – When the material has lost its surface gloss, apply a thin film of Riva Coat. – Light cure for 20 sec.</td>
</tr>
<tr>
<td>Vitremer™ (3M Deutschland GmbH, Neuss, Germany)</td>
<td>Powder: silane treated glass, potassium persulfate. Liquid: copolymer of acrylic and itaconic acids, water, HEMA, ethyl acetate, diphenyl iodonium hexafluorophosphatate</td>
<td>– Mix the powder into the liquid and place it into a 3M ESPE dispenser. Place the material in a dry field. – Light cure the glass ionomer for 40 sec.</td>
</tr>
<tr>
<td>Vitrebond™ Plus (3M Deutschland GmbH, Neuss, Germany)</td>
<td>Liquid: copolymer of acrylic and itaconic acids, water, hydroxethyl methacrylate (HEMA), ethyl acetate, diphenyl iodonium hexafluorophosphatate, tetrahydrofuran. Paste: silane treated glass, HEMA, water, bisphenol A diglycidyl ether dimethacrylate (Bis-GMA), silane-treated silica.</td>
<td>– Apply dentin conditioner. – Mix paste/liquid components (10 – 15 sec) and apply. – Light activation (20 sec).</td>
</tr>
<tr>
<td>ACTIVA BioACTIVE Restorative™ (Pulpdent®)</td>
<td>Paste/paste. Blend of diurethane and other methacrylates with modified polyacrylic acid (44.6 %), amorphous silica (6.7 %), and sodium fluoride (0.75 %).</td>
<td>– Etch 10 sec, rinse and dry. – Apply a bonding agent and place mix tip against the floor of the cavity. – Place a 1 – 2 mm insulating layer of ACTIVA and light cure before bulk filling.</td>
</tr>
</tbody>
</table>
For groups with mechanical mixing, the capsules were mixed in a rotary vibrator (3M™ RotoMix™ Rotating Capsule Seefeld, Germany) and the material was introduced into standardized polypropylene tubes with an internal diameter of 4 mm and a height of 6 mm using an applicator gun. All samples, except for the self-curing cements, were photopolymerized for 20 sec using a SmartLite LED lamp (Dentsply®, USA) at 1250 W/cm². The samples were photopolymerized for 20 sec using a SmartLite LED lamp (Dentsply®, USA) at 1250 W/cm². The samples were removed from the tube and cross-sectioned into 1 mm sections with a precision cutting machine (IsoMet 1000, BUEHLER, Illinois, USA), providing four sections per cylinder. Twelve sections were used for SEM and image analysis and five sections were used for microhardening study. The samples used for SEM and image analysis were etched with 37 % orthophosphoric acid for 5 sec, washed, and dried. They were then introduced in an ultrasonic machine (Calypso®- MESTRA) for 15 min immersed in 100 % ethyl alcohol and left at room temperature for dehydration.

2.2. SEM and image analysis

Sample surfaces were coated with a thin layer of gold (BIO-RAD Polaron SEM Coating System) and placed in a vacuum chamber at a voltage of 2.5 kV and an intensity of 20 mA. The coating time of each plate was 4 min. The coating time of each plate was 4 min. The samples were subsequently placed for analysis and dried. They were then introduced in an ultrasonic machine (Calypso®- MESTRA) for 15 min immersed in 100 % ethyl alcohol and left at room temperature for dehydration.

2.3. Microhardness measurements

Five samples in each group were used to study surface microhardness (SMH). SMH measurements were made using a Vickers diamond under a load of 100 g using a Microhardness Tester FM-310 (FUTURE TECH CORP., Kawasaki, Japan).

The force was applied for 15 sec. Each specimen was indented three times and the mean Vickers hardness value was determined for each sample.

2.4. Statistical analysis

Independent variables were pore diameter, number of pores/areas, and microhardness, and the dependent variables were the total area and percentage of area per section. The sample size (n = 12 per group) was calculated using pore diameter values, and accepting an alpha risk of 0.05, a beta risk of 0.20 (power 0.8) in a two-sided test to recognize a minimum difference of 4 units between any pair of groups, assuming that there were 9 groups. Common deviation was assumed to be 2.4. A drop-out rate of 0 % was anticipated.

Data are expressed as means ± standard deviation (SD). The Shapiro-Wilk normality test was applied. As there was no normality, the Kruskal Wallis test was used to detect between-group differences and Dunn’s test was used for two-by-two comparisons. To determine the interactions between the type of mixing and viscosity, we applied a robust ANOVA test. To establish possible correlations between variables, we used Pearson’s correlation test. Statistical significance was established as p < 0.05.

3. RESULTS

3.1. Number of pores per area

Ionostar® Plus (Figure 1A), EQUIA® Forte Fil (Fig. 1B) and Riva Light Cure HV® (Fig. 1 c) showed a significantly higher number of pores per section than other materials (Table 2).

3.2. Pore size

Riva Protect® had the largest pore-size (32.213 ± 8.519) (Fig. 1 d). ACTIVA BioACTIVE Restorative® (3.990 ± 3.034) (Fig. 1 e) and Vitrebond® Plus (7.194 ± 4.530) had the smallest pore sizes (Fig. 1 f) (Table 2).

<table>
<thead>
<tr>
<th>Material</th>
<th>Number of pores/area (82944 μm²)</th>
<th>Mean pore diameter</th>
<th>Total area occupied by pores</th>
<th>% of area occupied by pores</th>
</tr>
</thead>
<tbody>
<tr>
<td>EQUIA® Forte Fil</td>
<td>11.081 ± 5.955</td>
<td>15.890 ± 4.085</td>
<td>2749.746 ± 1779.295</td>
<td>3.315 ± 2.145</td>
</tr>
<tr>
<td>Ionostar® Plus</td>
<td>15.300 ± 7.498</td>
<td>10.379 ± 4.252</td>
<td>1804.628 ± 1386.520</td>
<td>2.20 ± 1.67</td>
</tr>
<tr>
<td>Riva Silver®</td>
<td>6.467 ± 1.833</td>
<td>22.313 ± 9.150</td>
<td>3506.631 ± 2696.618</td>
<td>4.228 ± 3.251</td>
</tr>
<tr>
<td>Riva Protect®</td>
<td>6.100 ± 1.853</td>
<td>32.213 ± 8.519</td>
<td>6147.767 ± 2828.360</td>
<td>7.412 ± 3.410</td>
</tr>
<tr>
<td>Vitremer®</td>
<td>7.867 ± 7.376</td>
<td>14.828 ± 6.895</td>
<td>2357.998 ± 1260.369</td>
<td>2.843 ± 1.520</td>
</tr>
<tr>
<td>Vitrebond® Plus</td>
<td>6.000 ± 6.641</td>
<td>7.194 ± 4.530</td>
<td>441.055 ± 682.527</td>
<td>0.532 ± 0.823</td>
</tr>
<tr>
<td>ACTIVA BioACTIVE Restorative®</td>
<td>7.400 ± 4.262</td>
<td>3.990 ± 3.034</td>
<td>395.719 ± 1309.223</td>
<td>0.477 ± 1.578</td>
</tr>
</tbody>
</table>

Statistical significance: a vs. Ionostar®Plus; b vs. VitrebondTM Plus; c, vs. Activa BioActive RestorativeTM; d vs. Riva Protect®
3.3. Total area and percentage of area per section

ACTIVA BioACTIVE Restorative™ (395.719 ± 1309.223) (Fig. 1 e) and Vitrebond™ Plus (441.055 ± 682.527) (Fig. 1 f) had the smallest area occupied by pores. Riva Protect® (6147.767 ± 2828.360) (Fig. 1 d) followed by Riva Light Cure HV® (4221.295 ± 3450.718) (Fig. 1 c) had the largest area occupied by pores (Table 2).

3.4. Interaction between the type of mixing and the degree of viscosity

We found no association in the number of pores per area between the type of mixing and the degree of viscosity (p = 0.769), but there was an association between the mean pore diameter (p = 0.001) and the total area occupied by pores (p = 0.002), showing that high viscosity ionomers were little influenced by these two factors, while in low viscosity ionomers (fluids), mechanical mixing produced larger pores and a larger occupied area than manual mixing.

3.5. Microhardness

Riva Silver® (116.633 ± 14.659) and EQUIA® Forte Fil (81.105 ± 7.716) showed the highest microhardness values.

Vitrebond™ Plus (27.335 ± 6.019) followed by Vitremer™ (23.375 ± 4.648) showed the lowest (Table 3).

There was no correlation between microhardness and the number of pores per area, pore diameter, total area, and percentage of area per section (Pearson’s correlation test) (Table 4).

Table 3. Surface microhardness (Vickers hardness-VHN) values for each group. Expressed as mean ± standard deviation

<table>
<thead>
<tr>
<th>Material</th>
<th>Microhardness</th>
<th>Statistical significance</th>
</tr>
</thead>
<tbody>
<tr>
<td>EQUIA® Forte Fil</td>
<td>81.105 ± 7.716</td>
<td>a, b, c</td>
</tr>
<tr>
<td>Ionostar® Plus</td>
<td>45.237 ± 10.70</td>
<td>a, b, c</td>
</tr>
<tr>
<td>3M™ Ketac™ Universal Aplicap™</td>
<td>76.494 ± 6.364</td>
<td>a</td>
</tr>
<tr>
<td>Riva Light Cure HV®</td>
<td>54.182 ± 8.865</td>
<td>a</td>
</tr>
<tr>
<td>Riva Silver®</td>
<td>116.633 ± 14.659</td>
<td>a, b, c, d, e</td>
</tr>
<tr>
<td>Riva Protect®</td>
<td>54.777 ± 6.469</td>
<td>a</td>
</tr>
<tr>
<td>Vitremer™</td>
<td>23.375 ± 4.648</td>
<td>a, b, c, d, e</td>
</tr>
<tr>
<td>Vitrebond™ Plus</td>
<td>27.335 ± 6.019</td>
<td>a, b, c, d, e</td>
</tr>
<tr>
<td>ACTIVA BioACTIVE Restorative™</td>
<td>36.975 ± 7.3</td>
<td>a, b, c</td>
</tr>
</tbody>
</table>

Statistical significance: a, vs. Riva Silver®; b, vs. EQUIA® Forte Fil; c, 3M™ Ketac™ Universal Aplicap™; d, Riva Protect®; e, Riva Light Cure HV®

Table 4. Pearson’s correlation test results of surface microhardness and number of pores per area, pore diameter, total area and percentage of area per section

<table>
<thead>
<tr>
<th></th>
<th>Microhardness</th>
<th>% of area occupied by pores</th>
<th>Total area occupied by pores</th>
<th>Number of pores per area</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>cc: 0.492; p = 0.179</td>
<td>cc: 0.981; p &lt; 0.001</td>
<td>cc: 0.981; p &lt; 0.001</td>
<td>cc: 0.314; p = 0.410</td>
</tr>
<tr>
<td>Pore diameter</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Microhardness</td>
<td>cc: 0.429; p = 0.179</td>
<td></td>
<td></td>
<td>cc: 0.0785; p = 0.841</td>
</tr>
<tr>
<td>% of area occupied by pores</td>
<td></td>
<td></td>
<td></td>
<td>cc: -0.209; p = 0.589</td>
</tr>
<tr>
<td>Total area occupied by pores</td>
<td></td>
<td></td>
<td></td>
<td>cc: 1.000; p = 0.001</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>cc: -0.0785; p = 0.841</td>
</tr>
</tbody>
</table>
| cc: correlation coefficient; p: P value.
4. DISCUSSION

The results of this study show that two HVGICs, Ionostar® Plus and EQUIA® Forte Fil, had the highest number of pores per section, and a LVGIC (Vitrebond™ Plus), had the fewest pores per section. This direct relationship between the high viscosity of glass ionomer cement and the presence of a large proportion of pores has also been described by other authors as a result of increased incorporation of air bubbles during the mixing process in HVGICs [10, 11, 19, 20 – 22]. Other authors, however, found more pores in LVGICs [21, 22].

Taking into account all materials, manual mixing generated pores with a mean diameter of 8.702 ± 1.146 µm, mechanical mixing 20.006 ± 0.999 µm and the automix syringe 2.783 ± 1.83 µm. For the diameter of the pores, the type of mixing showed a significant interaction with the viscosity level of the ionomers, so that in HVGICs the type of mixing did not significantly influence the pore diameter, while it was decisive for LVGICs. Thus, manual mixing in fluid ionomers generated smaller diameter pores than mechanical mixing.

Vitrebond™ Plus, with low viscosity and manual mixing, showed pores with a diameter of 7.194 ± 4.530 µm, while Riva Protect® with low viscosity and mechanical mixing, had the largest diameter pores (32.213 ± 8.5197 µm). Al-Kadhim et al. [23] also found that glass isomer cement in capsules (Fuji I CAPSULE) had pores with a greater diameter than manually mixed cement (Fuji I).

The same interaction between the type of mixing and the degree of viscosity observed for the size of the pores was also found for the area the pores occupied. Thus, the type of mixing was decisive for LVGICs. Manual mixing in Vitrebond™ Plus generated the lowest proportion of pores and mechanical mixing in Riva Protect® the highest proportion. With the great speed of mechanical mixing, the low viscosity of the material would allow the formation of more and smaller air bubbles, similar to a "foth". The slower process of manual mixing would prevent the inclusion of air and cause some bubbles to collapse [23 – 26]. However, some reports have described conflicting results, suggesting that manual mixing can incorporate air during the process and that automatic mixing would help prevent this by producing smaller pores and fewer large pores [15, 27].

The ionomer with the smallest diameter pores and the smallest area/proportion of pores was ACTIVA BioACTIVE Restorative™, available in an automix syringe, probably because it is not subjected to spatulation or mechanical mixing, thus reducing the incorporation of air bubbles.

Further study of the advantages that the automix syringe system could provide in the porosity of GICs is required.

Riva Silver® (116.633 ± 14.659) and EQUIA® Forte Fil (81.105±7.716) had the highest micro-hardening values. Riva Silver® showed the highest microhardness. These results are similar to those obtained by Nanda and Naik [28] and Yin et al. [29], who compared a conventional GIC with others whose composition included silver particles which, in addition to improving the mechanical properties of these materials, has an antibacterial effect, and is often used in restorative dentistry.

Hershkovitz et al. [30] obtained better microhardness results at 60 min. with EQUIA Fil versus Riva Self Cure and Ketac Molar. Sidhu [31] suggested the better mechanical properties of this high viscosity cement was due to greater crosslinking of the finest glass particles.

Two GICs using manual mixing had the lowest microhardness; Vitrebond™ Plus (27.335±6.019) and Vitremer™ (23.375± 4.648). Al-Taee et al. [17] also found that microhardness, the modulus of elasticity, and compression force, among other factors, were higher in GICs and RMGICs with mechanical mixing compared with those with manual mixing. They also found less porosity of the material if it used mechanical mixing, with an improvement in mechanical properties. However, we could not correlate the microhardness with the porosity of the GICs studied. Nor could we find any correlation between the viscosity and microhardness of the GICs, as has been shown in other studies [13,17,32].

Although the microhardness and the modulus of elasticity of the dentin are similar to those of GICs [33], these properties may be influenced by the operator's handling and insertion techniques of the GICs, and therefore the lower microhardness values in GICs with manual mixing may be due to unintentional alterations in the powder/liquid proportions when mixing the cement [13]. Therefore, in order not to alter this proportion and its mechanical properties, many authors suggest using only GICs with mechanical mixing [34].

According to the results of our study, we can partially reject the first null hypothesis since, regarding porosity, the type of mixing was only decisive in low viscosity ionomers and accept the second null hypothesis, since porosity did not influence the microhardness of the GICs studied.

5. CONCLUSIONS

The main strength of this study is the large number of GICs studied and the detailed analysis.

The results show that when the viscosity of glass ionomer cement is low, the size of the pores and the area they occupy depend on the type of mixing while, when the viscosity is high, the type of mixture has less influence. The automix syringe system seems to provide better porosity results. There was no relationship between porosity and microhardness.

Among all materials studied, EQUIA® Forte Fil and 3M™Ketac™UniversalAplicap™ may be recommended for clinical use as they had a high level of microhardness and intermediate porosity.

REFERENCES


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