Original Article

Flexural strengths and porosities of coated or uncoated, high powder-liquid and resin-modified glass ionomer cements

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Flexural strength; Glass ionomer cement; Micro-computed tomography; Porosity; Resin coating

Abstract Background/purpose: No study has previously investigated and compared whether resin coating could prevent the effect of dehydration on flexural strengths and porosities of high powder-liquid and resin-modified glass ionomer cements (HPL-GIC and RM-GIC). The purpose of this study is to investigate the effect of resin coating on flexural strengths and porosities of HPL-GIC and RM-GIC under a dry condition.

Materials and methods: HPL-GIC (Equia Forte Fil) or RM-GIC (Fuji II LC) was mixed and loaded into a mold to create a bar-shaped specimen, n = 12 of each. The specimens were randomly divided into two groups, coated and uncoated, n = 6 of each. In the coated group, a resin coating agent (Equia Forte Coat) was applied and light cured for 20 s. After 72 h, each specimen was dried and scanned to detect porosities (% volume) using micro-computed tomography. After scanning, flexural strength (MPa) of the specimen was tested using a three-point bending method.

Results: Porosities of HPL-GIC were significantly higher than RM-GIC, either coated or uncoated group (p < .05). Flexural strengths of coated and uncoated HPL-GIC were 41.47 ± 0.89 and 15.32 ± 1.15 MPa that were significantly lower than those of RM-GIC at 104.77 ± 3.97 and 52.90 ± 2.17 MPa (p < .05). Flexural strengths of coated GICs were significantly higher than uncoated GICs (p < .05).

Conclusion: Resin coating increased flexural strengths of GICs under dry condition. HPL-GIC had higher porosities and lower flexural strength than RM-GIC.

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Introduction

Glass ionomer cement (GIC) is a water-based restorative material, composing of basic glass powder and polyacid liquid and setting by an acid–base reaction.¹ Currently, GIC is classified into two main categories: high powder-liquid (HPL) and resin-modified (RM) GIC.² HPL or highly viscous GIC is modified from the conventional material by increasing the powder-liquid mixing ratio to accelerate setting reaction and also improve material’s properties.² RM-GIC is modified by adding polymerizable monomers into the liquid and/or powder component, which the polymerization reaction forms the polymer networks that enhance the properties.³

GIC is sensitive to water imbalance during initial setting.¹ To prevent water gain or loss, thin-layer coating on GIC with a protective agent is recommended to improve strength⁴ and reduce clinical wear.⁵ Resin coating for GIC has been recently introduced to replace the varnish or petroleum jelly. At long term, GIC is still sensitive to water loss even 6–12 months after placement.⁶ Rubber dam isolation is commonly placed during adhesive and restorative procedure. GIC in the isolation area becomes dehydrated, which induces micro-crack within the material and disintegrates bond to dentin.⁶⁷ Coating GIC before rubber dam isolation might protect the material from this drying effect.⁸

Effect of surface coating on the properties of GIC tends to be different between HPL-GIC and RM-GIC.⁹ Formation of polymerized resin networks protects RM-GIC from water gain and loss, which makes the material less sensitive to water sorption or dehydration than HPL-GIC.³⁴ For this reason, surface coating may be not absolutely mandatory for RM-GIC, and the benefit is still controversial.¹⁰

GIC is a two-component material that requires mixing before use. Porosities are detected in the mixed GIC by the air is trapped inside the material during mixing. The higher viscosity of GIC, the more internal porosities are detected.¹¹ The strength of restorative GIC has an inverse relationship to the amount of porosities.¹²,¹³ HPL-GIC, either hand- or machine-mixed, contains similar amount of porosities.¹³ In contrast, hand-mixed RM-GIC contains higher porosities than the machine-mixed material.¹² Nevertheless, no previous study directly compared between the two categories of GIC in term of porosities. In addition, the porosities of dried GIC may be altered due to water loss. All of previous studies investigated the porosities of GIC in the humid condition.¹¹–¹³

Micro-computed tomography (micro-CT) is a non-destructive tool that comprehensively investigate three dimensional (3-D) details of the scanned objects.¹⁴ Recent micro-CT tool is able to detect micro gaps and voids with the smallest effective voxel size of approximately 6 microns.¹⁵ From our pilot study, we found that micro-CT could create a dry condition in the chamber during 3-D scanning. In addition, the dry condition was standardized if the scanning time was controlled.

Most of recent studies investigated the effect of coating agent on the properties of GIC in the storage conditions without dehydration.³,⁴,¹⁶ No study has previously investigated and compared whether coating could prevent the negative effect of dehydration on the strength and porosities of HPL-GIC and RM-GIC. Therefore, the objective of this study was to investigate flexural strengths and porosities of resin-coated or uncoated, HPL-GIC and RM-GIC under the simulated dry condition.

Materials and methods

HPL-GIC (Equia Forte Fil, GC, Tokyo, Japan) and RM-GIC (Fuji II LC, GC, Tokyo, Japan) were tested in this study, which the details are present in Table 1. The encapsulated material was mixed using a triturator for 10 s according to the manufacturer’s instruction and loaded into a stainless steel mold size 25 × 2x2 mm covering with two glass slides to create a bar-shaped specimen, n = 12 of each material. The specimen of HPL-GIC was left for 5 min to ensure the initial set from the acid–base reaction. The specimen of RM-GIC was light cured from the top site using an LED light-curing unit (Demi Plus, Kavo-Kerr, Brea, CA, USA) with the 8-mm light guide for 20 s at each area. The set material was kept in a water bath at 37 ± 1°C for 15 min and then removed from the mold. The specimens were measured and polished with 400-grit silicon carbide paper to obtain the length 25 ± 0.2 mm and the width/thickness 2 ± 0.1 mm. The twelve prepared specimens of each material were randomly divided into two groups-coated and uncoated, n = 6 of each. In the coated group, a resin-based coating agent (Equia Forte Coat, GC, Tokyo, Japan) was applied on the surfaces of specimen using a micro-brush and light

| Table 1 | Manufacturer, main compositions and lot numbers of high powder-liquid and resin-modified glass ionomer cements, and a resin coating agent. |
|---------|----------------------------------------------------------------------------------------------------------------------------------|
| Materials | Manufacturer | Main compositions | Lot numbers |
|----------|---------------|-------------------|-------------|
| Equia Forte Fil (high powder-liquid) | GC corp., Tokyo, Japan | Powder: strontium fluoroalumino-silicate glass, polyacrylic acid powder Liquid: polyacrylic acid, polycarboxylic acid, tartaric acid | 1611153 and 1701241 |
| Fuji II LC (resin-modified) | | Powder: fluoroalumino-silicate glass, polyacrylic acid powder Liquid: polyacrylic acid, water, HEMA Methyl methacrylate, camphorquinone | 1704052 and 1510031 |
| Equia Forte Coat (resin coating) | | | 1502061 |
cured for 20 s at each area. In the uncoated group, no coating was applied, but the specimen of RM-GIC was also light cured for 20 s at each area to standardize the light-curing time. All specimens were kept in an incubator at 37 ± 1 °C and 100% humidity for 72 h to allow maturation of GIC before testing.

Each specimen was air dried and scanned to observe any internal porosities of the material using micro-computed tomography (micro-CT, SkyScan 1173, Bruker, Kontich, Belgium) set at 80 kV, 100 μA. The scanning concurrently created the dry condition to GIC for an approximate period of 20 min. The temperature in the operating room was set at 25 ± 2 °C with a humidity-controlled air conditioner. The 3-D images were reconstructed and analyzed using CT analyzer software (CTAn 1.16; Bruker). Volumes of the materials and porosities were identified using the CT volume software (CTVol 2.3.2.0; Bruker). Porosities were calculated and reported as percentages of the total material volume.

After micro-CT scanning, flexural strength testing using a three-point bending method was performed according to the ISO 4049-2009. A static load was applied at the center using a 2-mm diameter cylindrical tip at a rate of 1 mm/s until the specimen was fractured, and the force (N) was recorded. Flexural strength (MPa) was calculated according to the formula: \[ \text{Flexural strength (MPa)} = \left( \frac{F \times L}{2 \times W \times h^2} \right) \]
where \( F \) is the force until fracture, \( L \) is the length, \( W \) is the width, and \( h \) is the height of specimen. For statistical analysis, unpaired t-test was used to compare the flexural strengths and porosities between the uncoated and coated groups, as well as between HPL-GIC and RM-GIC.

### Results

Flexural strengths of uncoated and coated HPL-GIC were 15.32 ± 1.15 and 41.47 ± 0.89 MPa while those of RM-GIC were 52.90 ± 2.17 and 104.77 ± 3.97 MPa (Table 2). Flexural strengths of coated HPL-GIC or RM-GIC were significantly higher than those of the uncoated materials (\( p < .05 \)). Flexural strength of RM-GIC was significantly higher than HPL-GIC, either uncoated or coated (\( p < .05 \)).

The amount of porosities of HPL-GIC were significantly higher than RM-GIC (\( p < .05 \)), either in coated (0.20 ± 0.10% vs. 0.07 ± 0.03%) or uncoated group (0.18 ± 0.02% vs. 0.05 ± 0.02%) (Table 3). The porosities were not significantly different between the coated and uncoated groups (\( p > .05 \)).

The representative micro-CT images of uncoated and coated GIC materials are present in Fig. 1. The specimens of HPL-GIC showed many porosities distributed within the material. In contrast, the specimens of RM-GIC only contained few porosities within the material. The patterns of porosities were similar between the uncoated and coated specimens.

### Discussion

Under the simulated dry condition, the flexural strength of GICs coated with a resin coating agent were significantly higher than the uncoated cements. GIC is a water-based material that is sensitive to water loss, so the properties of the cements are negatively affected in the dry condition. Coating on GIC prevents water loss and, as a result, increases the flexural strength in this study.

When comparison between HPL-GIC (Equia Forte Fil) and RM-GIC (Fuji II LC), the flexural strength of the former was much significantly lower than the latter, regardless of coating or non-coating. Current HPL-GIC is clinically set within 5 min, yet the acid-base reaction still progresses until the maturation stage. HPL-GIC at early stage is partially mature with limited strength, as reported in our study. Hence, HPL-GIC strictly requires protection from resin coating after restoration placement or whenever water imbalance is expected on the restoration, such as after rubber dam isolation.

In addition to the polyacid salt matrix formation in HPL-GIC, the polymerized resin networks in RM-GIC protect the material from water gain and loss, and also improved the strength of the material. Our results also showed the strength of RM-GIC was much higher than HPL-GIC, regardless of coating. However, coating RM-GIC is still required since the coating significantly improved the strength of RM-GIC under the dry condition, as reported in our study. The importance of coating to RM-GIC is confirmed in the other studies that investigated the effect on gap formation and fracture toughness.

In our study, the porosities of GICs were not significantly affected by coating. Coating could prevent water loss from GICs under the dry condition, but it had no effect on the formation of porosities. However, we found that the amount of porosities of HPL-GIC was significantly higher than RM-GIC. The result is in correspondence to the study that investigated gaps and voids of HPL-GIC and RM-GIC when used as a base in restoration of endodontically treated teeth. This phenomenon might be explained by the difference in viscosity between the two materials. From the specimen preparation, the author noticed that the viscosity of the HPL-GIC was relatively higher than the RM-GIC, which might cause more voids or air bubbles inside the material during mixing. Furthermore, the higher amount of porosities in HPL-GIC might also take a part in the lower flexural strength when compared to RM-GIC.

### Table 2

| Flexural strengths (MPa) | Uncoated | Coated |
|-------------------------|----------|--------|
| Equia Forte Fil         | 15.32 ± 1.15\(^a\) | 41.47 ± 0.89\(^b\) |
| Fuji II LC              | 52.90 ± 2.17\(^c\) | 104.77 ± 3.97\(^d\) |

The superscripts with different small letters indicate a significant difference in flexural strength between the groups.

### Table 3

| Porosities (% volume) of the coated and uncoated specimens of high powder-liquid and resin-modified glass ionomer cements. |
|---------------------------------------------------------------|
| **Porosities (% volume)** | **Uncoated** | **Coated** |
|---------------------------|--------------|------------|
| Equia Forte Fil           | 0.18 ± 0.02\(^a\) | 0.20 ± 0.10\(^a\) |
| Fuji II LC                | 0.05 ± 0.02\(^b\) | 0.07 ± 0.03\(^b\) |

The superscripts with different small letters indicate a significant difference in % porosities between the groups.
In conclusion, coating with a resin-based agent significantly increased the flexural strength of either HPL-GIC (Equia Forte Fil) or RM-GIC (Fuji II LC) under the dry condition. RM-GIC had a significantly higher flexural strength than HPL-GIC, regardless of coated or uncoated. The porosities of HPL-GIC were significantly higher than RM-GIC. On the contrary, the porosities of coated and uncoated GICs were not significantly different.

Declaration of Competing Interest

The authors deny any conflict of interest.

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