Effects of Bacterial Cellulose Nanocrystals on the Mechanical Properties of Resin-Modified Glass Ionomer Cements

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Abstract

Objectives The purpose of this study was to evaluate the effect of bacterial cellulose nanocrystals (BCNCs) on the mechanical properties of resin-modified glass ionomer cements (RMGICs) including compressive strength (CS), diametral tensile strength (DTS), and modulus of elasticity (E).

Materials and Methods BCNCs were incorporated into RMGIC at various concentrations (0.3, 0.5, and 1 wt%). Unmodified RMGIC was used as the control group. The specimens were stored in distilled water at 37°C for 24 hours. CS and DTS, as well as modulus of elasticity, were evaluated using a universal testing machine. The nanostucture of BCNCs was observed via field emission scanning electron microscopy.

Statistical Analysis One-way analysis of variance and post-hoc Tukey tests were used for data analysis. Level of significance was at p < 0.05.

Results The addition of BCNCs to RMGIC led to an increase in all of the tested mechanical properties compared with the control group, with a significant increase observed for 1 wt% BCNC. CS and DTS improved up to 23%, and modulus of elasticity increased by 44%.

Conclusions The addition of BCNCs to the RMGIC improved the mechanical properties, including CS, elastic modulus, and DTS. Thus, the newly developed RMGICs with BCNCs might represent an ideal and promising novel dental material in restorative dentistry.

Introduction

Glass ionomer cements (GICs), which were introduced by Wilson and Kent in the early 1970s, are considered as the treatment choice in various clinical situations since they offer great benefits. These advantages include chemical adhesion to tooth structure, anticariogenic activity due to fluoride release, biocompatibility, and low coefficient of thermal expansion, which makes it similar to that of tooth structure.1 However, some drawbacks such as low mechanical strength, brittleness, and low wear resistance have made GICs far from being used as direct restorative materials in the load-bearing area.2

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To fortify the benefits and overcome the shortcomings of the conventional GIC, resin-modified glass ionomer cements (RMGICs) were introduced. Yet, the addition of 2-hydroxyethyl methacrylate (HEMA) in RMGICs led to greater water uptake and swelling of the resin matrix due to the hydrophilic composition of HEMA. Water leads to hydrolysis and plasticization of the resin-polyacrylate matrix, which might in turn deteriorate the mechanical properties of the cement. Therefore, RMGICs possess some desired properties over their conventional counterparts. However, drawbacks of RMGICs still need to be overcome. These deficits principally originate from their weak mechanical strength, including wear resistance, compressive strength (CS), and diametral tensile strength (DTS).

Nowadays, novel GIC designs have been introduced and various methods have been described by several researchers to ameliorate the mechanical and physical properties of glass ionomer restoratives. For instance, the addition of amagmat into the GIC powder was performed with the aim of refining the mechanical properties of these restorative materials. Yet, this combination has led to a diminished esthetics and lower bond strengths to the enamel. The mechanical properties of GICs have also been improved by incorporating with short silicon carbide fibers. However, it has been reported that these tiny fibers can accumulate in vital organs and produce toxicity similar to those of asbestos fibers. Moreover, numerous efforts have been conducted to enhance the GIC fillers by means of adding filler particles including hydroxyapatite, zirconia, ytterbium fluoride, barium sulfate, and silver into GICs. Yet, none of these methods have led to considerable reinforcements in wear resistance and mechanical strengths of GICs.

The great advancement in nanotechnology along with an urge for viable developments have led to the popularity of cellulose application because of its sustainable properties, harmlessness, abundance, low density, and little thermal expansion. The cellulose nanocrystals (CNCs) are extremely crystalline cellulose-derived structures that demonstrate outstanding mechanical strength. The CNCs, which have an average of 100 to 250 nm length and 5 to 15 nm diameter, are formed by acid hydrolysis of any natural source of cellulose. Nowadays, cellulose nanoparticles are among the most widely used materials in the improvement of mechanical properties in dental practice. CNCs form a solid scaffold in different directions within the material and greatly improve the mechanical properties by establishing hydrogen bonds. It is noteworthy to mention that the cellulose produced by some bacteria has a nanometer-sized width, even before going through processing. Bacterial cellulose is unique due to its high crystallinity, high water retention capability, and excellent mechanical and thermal properties, which is why many researchers prefer bacterial nanocelluloses for medical applications.

Although several studies have been performed to investigate various mechanical properties of RMGICs, no research has been conducted to assess the effect of bacterial cellulose nanocrystals (BCNCs) on mechanical properties of RMGICs. Therefore, the purpose of this study was to evaluate the effect of BCNCs in three volumetric masses of 0.3, 0.5, and 1 wt% on the mechanical properties of RMGICs. The null hypothesis of our study was that there would be no difference between the mechanical properties of RMGIC and BCNC-containing RMGIC.

Materials and Methods

A total of 80 specimens were prepared in this study. The specimens were divided into four main groups (n = 20): Group I (RMGIC powder, control), Group II (RMGIC powder with 0.3 wt% BCNC), Group III (RMGIC powder with 0.5 wt% BCNC), and Group IV (RMGIC powder with 1 wt% BCNC). Then the specimens of each group were divided into two subgroups for the CS and DTS tests (n = 10).

Sample Preparation

In this study, BCNC powder (Nano Novin Polymer Co.; Gorgan, Golestan, Iran) was used. Bacterial cellulose was extracted from Gluconacetobacter genus.

BCNC powders were weighed carefully to an accuracy of 0.001 g by means of a digital scale (GR-3000, A & D CL Toshiba, Tokyo, Japan) and were added to the previously weighed RMGIC powder (Fuji II, GC, Tokyo, Japan) containing 95% fluoroaluminosilicate glass (amorphous) and 5% polyacrylic acid using the correct concentration for each group (0.3, 0.5, or 1 wt%). To obtain a uniform powder in the specimens, initially RMGIC and BCNC powders were hand mixed and then the obtained powder was placed in amalgam capsules in an amalgamator (Ultramat 2, SDL Australia) for 20 seconds.

Then the resultant powder was mixed with RMGIC liquid (Fuji II) containing 20 to 30% distilled water, 20 to 30% polyacrylic acid, and 30 to 35% HEMA in accordance with manufacturer’s instructions (3/2 g: 1 g).

To determine CS and DTS, specimens were prepared in a cylindrical stainless-steel split mold (4 mm diameter and 8 mm height) in the same procedure described by ISO 9917-1:2017.

The mold was placed on the top of Mylar strip rested on a glass plate and the mold was filled with the material. Then a second piece of Mylar strip was placed on the material in the mold and pressed by another glass plate under hand pressure to remove excess material. The light cured specimens were eradicaded for 20 seconds trough Mylar strip using light-emitting diode at light intensity of 1,500 mW/cm^2 and a wavelength range of 440 to 480 nm (Coltulux II, Coltene, Ohio, United States). The glass plate and celluloid Mylar strip were then carefully removed. After removing the specimens, the specimens were cured from the other side to ensure that the samples were completely cured.

The specimens were placed in distilled water (37°C) for 24 hours prior to the experiment. For each test (CS and DTS), a total of 40 samples (n = 10) were placed in the universal testing machine (Instron, Z020. Zwick Roell, Germany).

Compressive Strength Test

For the CS measurement, the specimens were located lengthwise between the platens of the machine and compressed at a crosshead speed of 1 mm/min. The maximum load required to fracture each specimen was documented.
and the CS in mega Pascal (MPa) was measured using this formula: CS: \(4F/dl^2\)

Where \(d\) is sample diameter and \(F\) is the force loaded at the moment of fracture.

**Diametral Tensile Strength Test**

For the DTS, the samples were placed between the platens of the machine along their diameter. The tensile strength was measured according to the following formula:

\[ \text{Diametral traction tension: } 2F/dl \]

Where \(d\) is sample diameter, \(F\) is the force loaded at the moment of fracture, and \(l\) is the primary sample length.

**Modulus of Elasticity**

Young’s modulus for each specimen was determined according to the slope of the plotted graphs for CS produced by the universal testing machine.

\[ E: \sigma = (F/A)/(\Delta L/L) \]

Where \(A\) is the diagonal cross-section, \(\Delta L\) is the length change during compression test, \(L\) is the primary sample length, and \(F\) is the force loaded at the moment of fracture.

**Field Emission Scanning Electron Microscopy**

For field emission scanning electron microscopy (FESEM) analysis, a thin layer of gold-palladium was used to cover the representative specimen in BCNC experimental group. Using 10–15 kV electron beam, the specimens were witnessed by FESEM (CS-3500, Shimadzu, Kyoto, Japan) at magnifications of 1,000× and 75,000× (►Fig. 1).

**Statistical Analysis**

For data analysis, the mean and standard deviation (SD) values of all groups were obtained using Statistical Package for the Social Sciences version 15.0 (Microsoft, Illinois, United States). The average and SD were used for data description. To evaluate the homogeneity of data, the Kolmogorov-Smirnov test was used. One-way analysis of variance (ANOVA) and post-hoc Tukey tests were used for data analysis. Significance level was \(p < 0.05\).

**Results**

The mean ± SD values of CS, DTS, and modulus of elasticity (E) of various trial groups are shown in ►Table 1. The results of one-way ANOVA revealed that there was a significant difference between CS values of different tested groups \((p = 0.010)\). As shown in ►Table 1, the highest CS was observed in group IV (1% wt), which was significantly greater than the CS of the control group \((p = 0.007)\). No significant difference was found between the CS of the 1% wt group and that of 0.5% wt \((p = 0.804)\) and 0.3% w/w \((p = 0.565)\). Furthermore, a significant difference was observed between modulus of elasticity \((E)\) of the CS values of different experimental groups \((p < 0.001)\).

As shown in ►Table 1, the modulus of elasticity \((E)\) of the CS values of the control group was significantly lower than 0.3% w/w, 0.5% wt, and 1 wt% groups \((p < 0.001)\). Moreover, there was a significant difference between DTS values of different experimental groups \((p = 0.002)\).

As revealed in ►Table 1, the highest DTS was observed in group IV (1% wt), which was significantly greater than the DTS of the control group \((p = 0.001)\). No significant difference was found between the DTS of the 1 wt group and that of 0.5% wt \((p = 0.249)\) and 0.3% wt \((p = 0.055)\).

►Figure 1 shows representative FESEM images of 1 wt% BCNC in cementitious mass of RMGIC with a net-like structure and reveals the size of BCNC.

**Discussion**

The current study evaluated the effect of adding BCNCs on the mechanical properties of RMGICs. The null hypothesis was rejected. The result of this study showed that the RMGICs containing 1 wt% BCNCs represented significantly higher CS and DTS compared with the control group. Thus, adding BCNC to the RMGIC can increase the strength of the newly developed GICs without any difficulty with regard to clinical applications.

Nowadays, attention has been given to modern technologies in dentistry for the introduction of new restorative materials. In this regard, the introduction of cellulose fibers, especially CNCs, aimed to meet such demands and were in line with the objectives for the production of renewable biomaterials and green development. 26

The cellulose examined in the present study was derived from bacteria as this type of cellulose can be readily processed into nanocrystals. These nanocrystals are applied as strengthening constituents in producing polymers with high-performance applications. Bacterial cellulose presents a wider range of application compared with its herbal counterparts due to its exclusive thermo-mechanical characteristics and biocompatibility. 27,28 Therefore, the reinforced mechanical properties of bacterial cellulose-modified GICs might offer more durable and long-term restorations prepared with this restorative cement.

Previously, low concentrations of CNCs have been found considerably sufficient to enhance the mechanical strength of composites. In fact, according to Silva et al., 29 when the

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**Fig. 1** Field emission scanning electron microscopy images of 1 wt% bacterial cellulose nanocrystal BCNC-containing resin-modified glass ionomer cement: (A) 1,000× magnification showing the net-like structure of BCNC within cementitious mass; (B) 75,000× magnification showing the size of BCNC.
concentrations of CNCs were more than 1 wt%, the nanoparticles accumulated, leading to the failure of composites’ mechanical properties. Thus, in the present study, concentrations of 0.3, 0.5, and 1 wt% of BCNCs were used for the investigation of mechanical properties.

The results showed that the addition of 1 wt% BCNC to RMGIC led to a significant increase in the mechanical properties of the RMGIC. Furthermore, the addition of 0.3, 0.5, and 1 wt% of BCNCs to RMGICs resulted in a significant increase in the modulus of elasticity of the CS of these materials compared with the control group.

CS and flexural strength are regarded as load-bearing capacity indicators of a restorative dental material. GIC is a brittle material with a tensile strength distinctly lower than its CS. In fact, this material fails by dissemination of crack under tensile rather than compressive forces. Studying scanning electron microscopy confirmed that the addition of nanoparticles reduced porosity, with no differences between the materials. Therefore, it seems that the addition of BCNCs can fill spaces in the defects and impede the dissemination of previously present pores by blocking their paths, which in turn stops the pores from forming cracks and improves the DTS of RMGICs. It has been shown that an improvement in this property is hardly achievable in GICs. Therefore, this finding can be of high clinical significance since it is essential for GICs located in the load-bearing areas to resist the masticatory occlusal forces produced within their structure.

The FESEM image in this study revealed that the incorporation of BCNC to the RMGIC led to the creation of an interconnected network of BCNC, which was randomly distributed within the GIC matrix. Silva et al. have previously shown that the nanoscale size of the CNC fibers assists the dissemination of CNCs in the GIC matrix and facilitates the creation of a web-like arrangement within the matrix. As a consequence, this significant improvement in the mechanical properties of the BCNC-incorporated RMGICs can be attributed to the uniform dissemination of the interrelated nanocrystals in the cement matrix. In addition, BCNC’s ability to form a hydrogen bonding with the hydroxyl groups of the glass particles and carboxylic groups of the polyacrylic acid can be another possible explanation for the RMGIC's improved mechanical strength.

The formation of the network made by BCNCs adhering to the GIC particles may be explicated by the inherent self-association of the BCNCs. In fact, an electrostatic interaction occurs between the positive charges of the GIC and the negative charges of the BCNC that, in turn, causes cement reinforcement. This innate feature of the BCNC also helps in forming auxiliary architectures for the percolation of load into the cement matrix. Following this network formation, a substantial strengthening effect of the RMGIC was found in all the tested properties. DTS, CS, and elastic modulus increased from 18.64, 133.60, and 1,151.60 MPa in the control group to 22.84, 164.54, and 1,663.64 MPa in the 1 wt% added BCNCs, respectively.

In line with our findings, Silva et al. showed that the addition of a small amount of a nanoparticulate renewable material in the form of CNCs significantly increased the CS and DTS strengths of restorative GICs. In a recent study, CNCs were used in combination with titanium oxide (TiO₂) nanoparticles as an additive to GIC. It was reported that the physical properties of the modified GIC reinforced with 2 wt% TiO₂ nanoparticles and 1 wt% of CNC showed significant improvement.

Given the desired mechanical strength of the BCNC-incorporated RMGICs observed in this study, the RMGIC modified with 1 wt% BCNC might provide a practicable restorative material to be used under stress, which could help decrease tooth loss.

One of the limitations of this study was that the current research was conducted in-vitro. The in-vitro results obtained in the present study do not necessarily validate those achieved in vivo. For this matter, there is an urge for clinical studies to confirm in-vitro findings. Moreover, only compressive and tensile tests were performed in our study. The future of RMGICs modified with BCNCs still requires further investigations on other properties of these novel materials. Properties such as color stability and biocompatibility are needed to be studied for the comprehensive finding of the properties and characteristics of these materials.

### Conclusions

It was found that the addition of 1 wt% of BCNC to the RMGIC considerably improved all the tested properties, including CS, DTS, and elastic modulus. Thus, the newly developed RMGICs with BCNCs might represent an ideal and promising novel dental material in restorative dentistry.

### Conflict of Interest

None declared.

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