

Research article

Characterization and enhancement of physico-mechanical properties of glass ionomer cement by incorporating a novel nano zirconia silica hydroxyapatite composite synthesized via sol-gel

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Abstract: In restorative dentistry, there has been a growing shift towards using nanoparticles dispersed in the polymer matrix to improve properties of dental restorative materials. A new nano zirconia–silica–hydroxyapatite (nanoZrO₂–SiO₂–HA) was synthesized by one-pot synthesis and characterized using transmission electron microscopy (TEM), scanning electron microscopy (SEM), Fourier transform infrared spectroscopy (FTIR), energy dispersive X-ray (EDX) and dot mapping. The effect of addition of nanoZrO₂–SiO₂–HA to the conventional GIC (cGIC) on the compressive strength, flexural strength and surface roughness was also evaluated. The characterization studies confirmed that all particles were in the nanoscale range with spherical zirconia and silica particles embedded in the voids between rod-shaped HA crystallites. The nano particles were evenly and homogeneously dispersed throughout the sample with high density patterns visible for zirconia, calcium and phosphorus. The incorporation of 5% nanoZrO₂–SiO₂–HA has resulted in considerable improvement in the compressive and flexural strengths of cGIC. The GIC 5% nanoZrO₂–SiO₂–HA exhibited an increase in compressive (144.12 ± 13.88 MPa) and flexural strength (18.12 ± 2.33 MPa).

over cGIC which was statistically significant ($p \leq 0.05$). It also demonstrated surface roughness profile ($0.15 \pm 0.029 \mu\text{m}$) similar to that of cGIC ($0.15 \pm 0.019 \mu\text{m}$). Therefore, the nanoZrO₂-SiO₂-HA can be a promising filler for GIC to be used as restorative dental material in high stress bearing areas.

Keywords: glass ionomer cement; nanomaterials; mechanical properties; characterization; zirconia

1. Introduction

Many restorative materials have been used in dentistry over the decades. There is now a trend towards the use of tooth-colored restorative materials such as conventional glass ionomer cement (cGIC) and composite resins. Composite resins are, however, often considered unsuitable for special needs and pediatric patients due to the difficulty of isolation and moisture contamination during placement of the restoration [1]. On the other hand, glass ionomer cements do not require such stringent moisture control and isolation protocols [2,3]. They were first introduced in dentistry by Wilson et al. at the Laboratory of the Government Chemist in 1969 [4,5].

GIC is the generic name for a group of classifying materials originally produced from aluminosilicate polyacrylates [6,7]. The powder of the ionomer glass is manufactured by fusion of its main components (SiO₂, Al₂O₃ and CaF₂) at a temperature between 1100 and 1500 °C to form a homogeneous fluoro-aluminosilicate glass [8,9]. Silica and alumina are responsible for the mechanical strength of the material, while calcium fluoride contributes to the F ions release at the surrounding media [10,11].

As aqueous polyelectrolyte systems, GICs are known for their relative ease of use, chemical bonding to tooth substrate, fluoride ion release and recharge, low coefficient of thermal expansion and acceptable aesthetic quality [12]. GIC is also found to be biocompatible with pulp tissue and with primary cultures of bone cells. In the clinic, however, the use of GIC is limited due to its relatively inferior mechanical properties and sensitivity to initial desiccation and moisture [13–15].

To expand the usage of cGIC, especially for restoration of adult teeth and for anterior teeth restorations, the physical, mechanical and aesthetic properties must be improved. The polymeric substrate of these ionomeric cements act as a vehicle or carrier for different organic and inorganic fillers or additives. Glass ionomers have already distinguished themselves as unique restorative materials possessing the potential for remineralisation and antimicrobial action owing to their fluoride uptake and elusion ability. As a result of this unique ability, GICs have potentially been employed as templates for releasing other active antimicrobial components, thereby, further expanding their use as a restorative material for deciduous teeth and in atraumatic restorative therapy (ART) [16].

The earliest attempt has been reported by Simmons in 1983 via simple addition of silver amalgam alloy powder to GIC powder and subsequently, McLean and Gasser fused and sintered amalgam powders to basic glass particle [17,18]. The resulting cermet particles exhibited strong bonding between the metallic and glass particles. However, these modifications were still not enough

for GICs to be used as a posterior restorative material [19,20]. Metal powders [21–25], reactive glass fibres [26,27] and other non-reactive fillers have been evaluated as potential reinforcement phases in order to improve the mechanical properties of GICs without compromising their adhesive and fluoride release properties. In one such study, researchers added micron size hydroxyapatite (HA: $\text{Ca}_{10}(\text{PO}_4)_6\text{OH}_2$) to cGIC resulting in improved biological properties of GIC [28]. Later, Yap et al. reported that the addition of HA enhanced the hardness of GIC [29]. Hydroxyapatite is an integral part of enamel and dentine with an ability to remineralize initial incipient caries and improve the bond strength to dentine [2,30].

In restorative dentistry, there has also been a growing shift towards using nanoparticles to improve properties of cGICs [31]. In the last decade, nanoscale HA [32,33], combination of HA–silica (SiO_2) [34,35] have been successfully used to improve GICs. Recently, nanophases of zirconia (ZrO_2) or a combination of HA– ZrO_2 [36–38] have been incorporated in attempts to strengthen GIC with enhanced outcomes. Zirconium and its oxide, due to their good dimensional stability and toughness (on the same order as stainless-steel alloys), have been used widely for the toughening and strengthening of brittle HA bio glasses in biomedical applications [39]. Furthermore, ZrO_2 has been used in fabrication of high strength core for dental implants [40]. Another more recent study was done to evaluate the microhardness of a HA– SiO_2 – ZrO_2 incorporated GIC with some encouraging results [41].

However, there is still dearth of information on the assessment of physico-mechanical properties of nano ZrO_2 – SiO_2 –HA GIC hybrid in terms of compressive strength (CS), flexural strength (FS) and surface roughness (SR). Therefore, the aim of the present study was to evaluate the effectiveness of a novel nano ZrO_2 – SiO_2 –HA reinforcement phase on the physico-mechanical properties of cGIC. The nano powder was synthesized and characterized morphologically and qualitatively prior to the evaluation of CS, FS and SR of the GIC nano ZrO_2 – SiO_2 –HA hybrid material. The null hypothesis tested was that the addition of ZrO_2 – SiO_2 –HA to the cGIC does not improved its physico-mechanical properties.

2. Materials and methods

2.1. Materials

The alumino-silicate glasses and the polyacids used in the research study were from commercially available Fuji IX[®] (GC, Tokyo; Japan). All the other chemicals and materials were of analytical grade and used as received from the manufacturer. Tetraethyl orthosilicate (TEOS, >99%), ammonia (NH_3 , 25%) and calcium hydroxide ($\text{Ca}(\text{OH})_2$, >98%) were obtained from Merck Serono Pvt. NSW, Australia. Phosphoric acid (H_3PO_4 , >99%), and ethanol ($\text{C}_2\text{H}_5\text{OH}$, >99%) were procured from Sigma-Aldrich Co. LLC, MO, USA, and Zr nanopowder (ZrO_2 , >99%) was from US Research Nanomaterials Inc., TX, USA.

2.2. Synthesis of nanoZrO₂-SiO₂-HA

The nanoZrO₂-SiO₂-HA was synthesized using a modification of the sol-gel technique [42] described by Ab Rahman et al. [41]. Initially, Ca(OH)₂ was dissolved in distilled water using a magnetic stirrer until a suspension was formed. After this, a stoichiometric amount (i.e., Ca/P molar ratio of 1.67) of H₃PO₄ (0.075 M) held in a burette was added drop wise to the suspension and the pH of the suspension was adjusted by adding ammonia till it was stabilized in the range of 11–12. The resulting suspension was stirred continuously for 48 h. Then, a quantity of 5 mL of TEOS and 25 wt% ZrO₂ dissolved in C₂H₅OH was added drop wise. C₂H₅OH was mixed with TEOS because it is immiscible in water, but dissolves easily in C₂H₅OH. Hence, the mixing greatly affects the homogeneity of TEOS in the reaction. Finally, the sol was stirred for an additional 12 h and then centrifuged (Centrifuge 5702, Eppendorf[®], Hamburg; Germany), washed with C₂H₅OH, freeze dried (ScanVAc CoolSafe, LaboGene ApS; Lyngø) and calcined (WiseTherm[®], Witeg Lab. GmbH; Germany) at 600 °C for 1 h.

2.3. Characterization of nanoZrO₂-SiO₂-HA nanopowder

The morphology and microstructure of the samples was studied by using a fully analytical energy filtered transmission electron microscopy (EFTEM) with an acceleration voltage of 120 kV and objective lenses with resolution point of 0.34 nm (Zeiss Libra 120 plus). Dilute dispersion of 100 mg of nano powder was washed in 3 mL of ethanol and was subjected to ultrasonication for 10 min to disperse the nano particles and to obtain a homogeneous suspension. One drop of this suspension was carefully placed with the help of a pipette on a 2.3 mm carbon-coated copper grid with a thickness of 10–20 nm.

The microstructure of the nanoZrO₂-SiO₂-HA dehydrated powder was also observed under scanning electron microscopy (SEM, QuantaTMFEG, ThermoFisher[®], OR; USA). Prior to evaluation, each powder specimen was coated with gold using a sputter coater (Desk-II, Denton Vacuum Inc., NJ; USA) at a gas pressure of ~50 mTorr, and ~40 mA current. The coating time required was 180 s. Energy dispersive X-ray analyzer (EDX, Rontec; Germany) which was directly connected to SEM was used to investigate semi quantitatively the chemical composition of the nanoZrO₂-SiO₂-HA.

In addition, Fourier transform infrared spectra (Spectrum TwoTM FTIR, PerkinElmer, MA; USA) of the GIC nanoZrO₂-SiO₂-HA powder was measured in the spectral range of 400–4000 cm⁻¹. The standard spectral resolution of the instrument was between 0.16 and 0.5 cm⁻¹ suitable for most applications.

2.4. Fabrication of GIC nanoZrO₂-SiO₂-HA hybrid samples

Conventional GIC, Fuji IX[®] was used as control and base material. The nanoZrO₂-SiO₂-HA powder was added at varying percentage of 3%, 5%, 7% and 9% by weight into the cGIC to prepare a total of four experimental groups. The mixture was then ground manually using a pestle and mortar in a control grinding process for about 10 min. All the samples (cGIC and GIC nanoZrO₂-SiO₂-HA) were prepared by mixing with Fuji IX-liquid at the recommended powder/liquid ratio (P/L) of 3.6.

The physico-mechanical properties evaluated were compressive strength, flexural strength and surface roughness after 24 h of setting.

2.4.1. Compressive strength testing

Cylindrical samples measuring 4 mm × 6 mm were prepared in stainless-steel split mould following the procedures outlined in ISO 9917-1: 2007. The moulds were coated with a thin layer of cocoa butter provided by the manufacturer of Fuji IX prior to filling the moulds to facilitate the easy retrieval of the set samples. They were then filled with the material and covered with polyester strips and glass slides and gently pressed by hand to remove air bubbles from uncured cement paste. The split mould was carefully disassembled after 30 min and the samples were gently retrieved by hand from the moulds and conditioned in distilled water at 37 °C for 24 h. Then, each sample was placed with the flat ends between the platens of a universal testing machine (AG-X plus 20 KN, Shimadzu Corp.; Japan) and a compressive load was applied along the long axis of the sample at a crosshead speed of 1 mm/min until the sample fractured. The compressive strength (CS) was calculated in mega pascals (MPa) using the following equation: $CS = 4P/\pi d^2$, where P is the load at fracture (N), and d is the diameter (mm) of the cylindrical sample and π is Pi which has a numerical constant of 3.14.

2.4.2. Flexural strength testing

Beam-shaped samples measuring 3 mm × 3 mm × 25 mm were prepared in a stainless-steel split mould following the procedures outlined in ISO 9917-2: 2007 and as aforementioned. Samples were kept in 100% humidity at 37 °C for 50 min before being immersed in distilled water and stored for 23 h at 37 °C [43]. Then, they were subjected to a three-point bending test in a universal testing machine (AG-X plus 20 KN, Shimadzu; Japan) at a crosshead speed of 0.5 mm/min with a 20 mm span or at a rate of loading of 50 ± 16 N/min and the flexural strength (FS) was recorded.

2.4.3. Surface roughness evaluation

A polytetrafluorethylene mould (5 mm diameter × 2 mm deep) was used to prepare how many samples according to the procedure outlined before. The prepared samples were then stored at 100% relative humidity for 24 h. Only a polyester strip and a glass slide were used with the intention of obtaining a smooth surface. Any form of additional polishing can lead to an increase in surface roughness (SR) and hence no polishing of the samples was carried out [44].

Each sample was blot dried and subjected to SR testing using a profilometer (Surfcom Flex, Accretch, Tokyo Seimitsu Co. Ltd.; Japan) and the value (Ra) obtained was in μm . To measure the roughness profile value, the diamond stylus (5 μm tip radius) was moved across the surface under a constant load of 4 mN. Five measurements passing through the center of the sample were performed and the average Ra taken. The cut-off value for surface roughness will be 0.8 mm based on mathematical cut-off filters in-built in the processor unit.

2.4.4. Statistical analysis

The data was analysed using SPSS version 22.0 (SPSS Inc., Chicago, IL; USA). Numerical data was tabulated and presented as mean \pm SD based on their normality distribution. A $p \leq 0.05$ was considered as statistically significant. One-way analysis of variance (ANOVA) was performed to determine whether p -value was statistically significant, and a post-hoc Tukey's HSD multiple range test was used to determine which experimental groups were statistically different from one another in the evaluation of *CS*, *FS* and *SR*.

3. Results and discussion

3.1. Characterization interpretations

3.1.1. FTIR spectra analysis of GIC nanoZrO₂-SiO₂-HA hybrid

FTIR absorption spectra (Spectrum Two™ FTIR, PerkinElmer, MA; USA) of the GIC nanoZrO₂-SiO₂-HA hybrid containing 3 to 9% of nanoZrO₂-SiO₂-HA filler was measured in the spectral range of 400–4000 cm⁻¹. The standard spectral resolution of the instrument was between 0.16 and 0.5 cm⁻¹ suitable for most applications.

FTIR analysis shows some variations in adsorption peaks of the GIC nanoZrO₂-SiO₂-HA hybrid material when compared to the FTIR of nanoZrO₂-SiO₂-HA nanopowder [41] (Figure 1). The band at 3436.60- is due to ν_2 bending mode of absorbed water OH stretching vibration confirming the presence of alcohols and phenols [34,41,45]. The band at 1709.51 is due to stretching vibration of carboxyl group of ketones or quinone. This confirms ν_3 stretching vibration of PO₄³⁻. The peaks at 964.41, 1041.56 and 1093.64 observed in the FTIR spectra of nanoZrO₂-SiO₂-HA [41] (representing the bending mode of PO₄³⁻ and the stretching vibration of Si-O, Zr-O) are replaced by a prominent peak at 1034.28 which becomes a bit broad in the GIC nanoZrO₂-SiO₂-HA hybrid but is otherwise stable in position. It arises due to symmetric and asymmetric stretching of Si-O-Si, i.e., with the addition of Zr it increases the distance of Si-O [34,41,46]. It specifies the Zr-O-Si bond formation and is because of the large ionic radius of Zr [38,47].

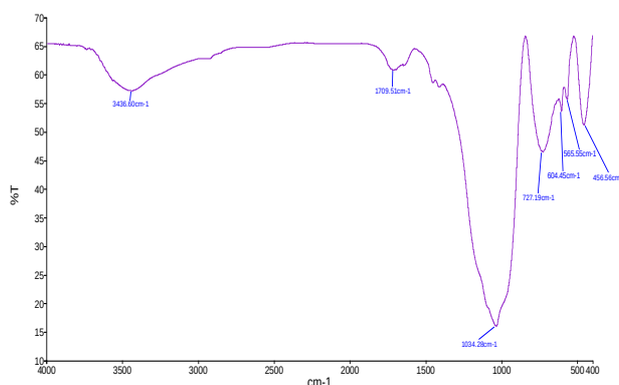


Figure 1. FTIR spectra of GIC nanoZrO₂-SiO₂-HA hybrid material.

The small peak at 731.02 seen in the FTIR spectra of nanoZrO₂-SiO₂-HA [41] is replaced by a prominent peak at 727.19 that arises due to CaF₂; this peak might also be due the presence of θ -Al₂O₃. A new peak 456.56 appears due to the bending vibration of Si-O-Si, AlPO₄ and α -Al₂O₃. These results indicate some molecular interaction presented between the nanoZrO₂-SiO₂-HA and GIC in the composite.

3.1.2. TEM evaluation

Comparative morphologies of nanoZrO₂, nanoSiO₂ and HA prepared by the modified sol-gel technique [34,41,43], as observed under EFTEM in a 200 nm micrograph are shown in Figure 2. The image reveals a narrow size and shape distribution in the morphology of the particles with roughly low agglomeration and aggregation.



Figure 2. TEM micrograph of nanoZrO₂-SiO₂-HA at low magnification (40000 \times)

The 100 nm micrograph confirm that the asymmetrically elongated rod like structures are indeed HA crystallites (Figure 3a), which has been documented in previous studies [35,36] and apart from the elongated rod shaped HA structures, smaller spherical structures embedded between and along the elongated HA are the silica particles [34,56,58]. These are also accompanied with oval to spherical slightly larger and darker structures which are most likely the nanoZrO₂ particles (Figure 3a,b) and these observations are in agreement with findings of a previous study [38]. These micrographs confirmed our assumption that all the powder particles were in or close to the nano scale range with the elongated HA measured to a mean length of 114 nm with SiO₂ having a mean size of approximately 18 nm (Figure 3a, Inset). These observations were in agreement with previously published literature [34,41,42,56]. The nanoZrO₂ had a mean size of 39 nm which confirms that the particles are indeed ZrO₂ as the commercially supplied ZrO₂ used in the one-pot synthesis was of the dimension of 40 nm and these dimensions are similar to many published studies [37,38,41]. The overall smaller particle sizes were attributed to the modified one-pot synthesis technique and the prolonged freeze drying. Slowing the rate at which ammonia is fed resulted in smaller ZrO₂ and SiO₂ nanoparticles and freeze drying produced mesoporous crystallites with high surface area.

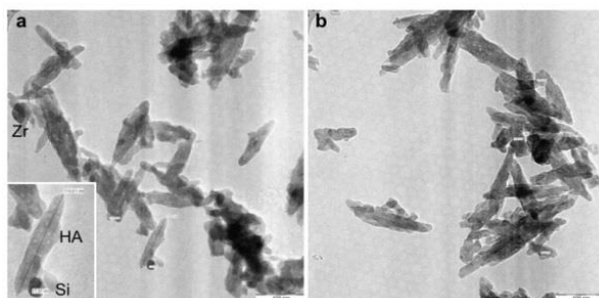


Figure 3. TEM micrograph of nanoZrO₂-SiO₂-HA at high magnification (60000×) at two different locations (a) and (b), Figure 3(a) inset shows the close up microstructure of HA and Si.

There was presumably a homogenous mix of the rod-shaped HA, spherical ZrO₂ and SiO₂ particles in the nano powder. Therefore, it can be inferred that the morphology of the nanoZrO₂-SiO₂-HA powder consists of a mixture of spherical ZrO₂, SiO₂ particles embedded in the voids or spaces between rod-shaped HA.

3.1.3. SEM evaluation

SEM micrographs of nanoZrO₂-SiO₂-HA powder in Figure 4a,b clearly reveal that nano hybrid powder is a mixture of elongated rod-shaped structures which is HA crystallites, smaller spherical powder shaped structures which are SiO₂ particles in close proximity of which are the spherical slightly larger and whiter nanoZrO₂ particles. These observations are in line with many of the previously reported literature [34,35,38,57]. The SEM images also confirm that nanoSiO₂ and nanoZrO₂ have occupied the spaces present between HA rods compacting the nano powder and forming a denser and stronger nanoZrO₂-SiO₂-HA powder [36,41,43,59]. The higher magnification (200000×) SEM micrograph (Figure 4b) reveals that the nanoHA rods have a rough outer texture which suggests that they may be covered by very fine SiO₂ and probably ZrO₂ residue. Similar conclusions have been recorded by authors in another study [43].

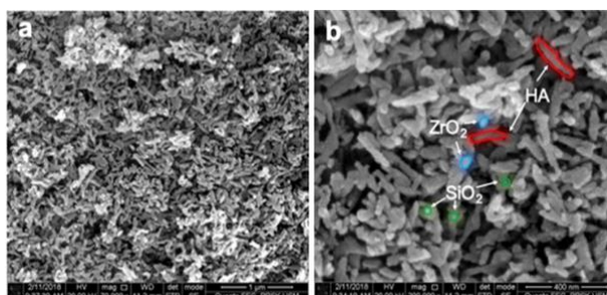


Figure 4. SEM micrograph of nanoZrO₂-SiO₂-HA at (a) low magnification (70000×) and (b) high magnification (200000×).

3.1.4. EDX and dot mapping analysis

Energy dispersive X-ray results in Figure 5 shows that the main elements present in the nanoZrO₂-SiO₂-HA powder are Ca, P, Zr, Si and O. A total of 4 iterations were done and peaks possibly omitted were at 9.703, 11.508 keV. L-alpha line of Ca was present while K-alpha lines of Si, O, P and Ca are observed. Thus, it is affirmed that a nanoZrO₂-SiO₂-HA nano composite is formed after analyzing the quantitative presence of Ca, P and Si in the EDX spectrum.

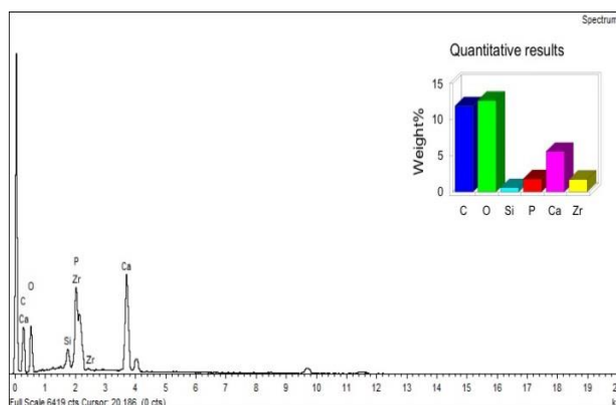


Figure 5. EDX spectrum of nanoZrO₂-SiO₂-HA.

It is possible to analyse the spatial distribution and density of Zr, Si, HA in the powder by performing a dot mapping using EDX. Figure 6 shows the SEM dot mapping characteristics for (a) Zr, (b) Ca, (c) P and (d) Si. All of these elements were evenly and homogeneously dispersed throughout the sample scan area with high density patterns visible for Zr, Ca followed by P without any evident of clumping or aggregation [41]. Si was the least dense with uniform distribution which is in accordance with the low quantity of TEOS (5 mL) used in the one-pot synthesis.

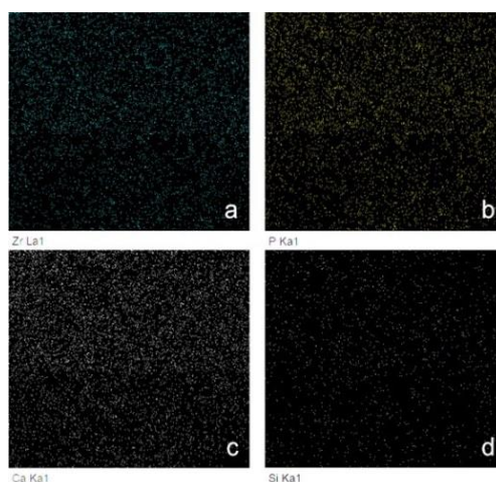


Figure 6. SEM dot mapping of (a) Zr, (b) P, (c) Ca, (d) Si of nanoZrO₂-SiO₂-HA.

3.2. Evaluation of physico-mechanical properties

Mechanical properties are very important for a dental restorative material. In order to obtain maximum service, these materials should withstand the forces generated during mastication [60]. The stability of a solid material under the applied force is determined by the nature of the material and atomic bond strengths. Materials used in restorative dentistry should have sufficient strength to withstand the complex forces that repeatedly can arise from mastication during the chewing process. The average force that is observed in chewing is about 200–400 N [61].

Physical properties especially surface roughness (SR) of restorative materials in the oral environment is an important criterion to predict their clinical deterioration. Surface characteristics of GIC restorations are particularly of concern because rough surfaces are more prone to faster bacterial colonization and plaque maturation, thereby increasing the risk of caries [62] ultimately leading to staining and poorer aesthetics. In addition, an increase in SR might indicate material deterioration [63–65].

3.2.1. Compressive strength evaluation

Compressive strength is useful for comparing materials that are brittle and generally weak (such as GICs) in tension [66]. The compressive strength of a material is defined as the maximum stress that the material can undergo without fracture. Compressive strength is widely used as a method of measuring the strength of GICs. Fuji IX Universal was selected as the control GIC because it is a widely popular commercially available cGIC and is recommended for this reason by WHO for ART [33]. Also, Fuji IX contains glass predominantly composed of SiO_2 , Al_2O_3 , CaF_2 , Na_3AlF_6 , and AlPO_4 in which the Ca is substituted with Sr without affecting the setting reaction of GIC. Therefore, the absence of Ca in the glass composition of Fuji IX was considered an advantage as it would not interfere with the characterization and quantitative assessment of Ca release characteristic from HA of the nanoZrO₂–SiO₂–HA powder. The similarity between HA and the chemical and skeletal structure of bone and teeth makes it an obvious candidate to expand the applications of GIC [67]. Similarly, high strength, fracture toughness and biocompatibility of ZrO₂ make it a valuable adjunct for overcoming the brittleness of HA [36,37,68,69].

Table 1 shows the CS of GIC nanoZrO₂–SiO₂–HA experimental groups and cGIC which were tested after being immersion in distilled water bath at 37 °C for 1 d. The data demonstrate that compared with the control, the GIC nanoZrO₂–SiO₂–HA groups exhibited considerably higher compressive strength ($p < 0.05$) for GIC 5% nanoZrO₂–SiO₂–HA group (144.12 ± 13.88 MPa). This finding is in agreement with previous results which reported the highest increase in CS for 4% GIC–HA as compared to Fuji IX [29]. The GIC 5% nanoZrO₂–SiO₂–HA exhibited a ~30% increase in CS over Fuji IX. These values were higher than the increase of ~18.42% reported by Moheet et al. [43] for GIC nanoHA–SiO₂ as well as the ~14% in CS of a GIC–HA material developed by Moshaverinia et al. [70]. This was probably due to the high ZrO₂ content (25%) of the nano powder. The results also exhibited a linear correlation up to 5 wt% addition of nanoZrO₂–SiO₂–HA powder to the GIC after which the CS values decreased. This could be due to the overcrowding of the nano filler at over 5 wt%, resulting in the disruption of the poly salt bridge formations in the GIC matrix.

It is postulated that the glass component of this hybrid GIC undergoes finely controlled micronization to achieve optimum particle size and characteristics [71]. In addition, the homogeneous and dense presence of ZrO_2 particles in the nano powder further reinforces and improves the mechanical properties of GIC as verified by SEM dot mapping scans. The nano ZrO_2 - SiO_2 -HA particles occupy the empty voids between the glass particles, thereby enhancing the packing density and acting as a reinforcement phase in the GIC composition [41,59].

Table 1. Compressive strength comparison between the various experimental groups.

Group	Comparison	Mean compressive strength \pm SD (MPa)	<i>p</i> -value
Control (117.64 \pm 16.40 MPa)	GIC 3% nano ZrO_2 - SiO_2 -HA	126.78 \pm 9.369	0.215
	GIC 5% nano ZrO_2 - SiO_2 -HA	144.12 \pm 13.88*	0.000
	GIC 7% nano ZrO_2 - SiO_2 -HA	133.84 \pm 15.77*	0.003
	GIC 9% nano ZrO_2 - SiO_2 -HA	120.47 \pm 8.59	0.964
GIC 3% nano ZrO_2 - SiO_2 -HA	GIC 5% nano ZrO_2 - SiO_2 -HA	144.12 \pm 13.88*	0.001
	GIC 7% nano ZrO_2 - SiO_2 -HA	133.84 \pm 15.77	0.471
	GIC 9% nano ZrO_2 - SiO_2 -HA	120.47 \pm 8.59	0.583
GIC 5% nano ZrO_2 - SiO_2 -HA	GIC 7% nano ZrO_2 - SiO_2 -HA	133.84 \pm 15.77	0.125
	GIC 9% nano ZrO_2 - SiO_2 -HA	120.47 \pm 8.59*	0.000
GIC 7% nano ZrO_2 - SiO_2 -HA	GIC 9% nano ZrO_2 - SiO_2 -HA	120.47 \pm 8.59*	0.020

*Denotes statistically significant difference at *p*-value \leq 0.05.

3.2.2. Flexural strength evaluation

The present study compared the flexural strength of varying wt% of nano ZrO_2 - SiO_2 -HA to the control (Table 2). There was a linear correlation between the *FS* values and the wt% addition of nano powder to the GIC up to 7% following which further addition of 9% nano ZrO_2 - SiO_2 -HA to the GIC led to a decrease in the *FS*. These observations were in agreement with previous findings [32,70] and could be due to overcrowding of filler particles as well as dry cement mix owing to the high powder/liquid ratio needed to spatulate higher wt% nano ZrO_2 - SiO_2 -HA powder into the GIC. Arita and colleagues added HA granules at different weight (8–25%) percentages and concluded that HA addition to GIC at 8% had the highest *FS* values. In our study, similarly the highest *FS* values recorded were for GIC 5% nano ZrO_2 - SiO_2 -HA \sim 18.12 \pm 13.88 MPa instead of GIC 9% nano ZrO_2 - SiO_2 -HA. Similar observations were made in another study in which the authors added HA and Si nano powder to cGIC at varying w/w% (5–20%) and reported the highest *FS* values of 17.68 \pm 1.81 MPa for 10% GIC nanoHA- SiO_2 . They concluded that further addition of higher percentages (15% and 20%) of nanoHA- SiO_2 to cGIC resulted in decreased *FS* values [43].

Table 2. Flexural strength comparison between the various experimental groups.

Group	Comparison	Mean flexural strength \pm SD (MPa)	<i>p</i> -value
Control (14.38 \pm 3.01 MPa)	GIC 3% nanoZrO ₂ -SiO ₂ -HA	14.44 \pm 2.17	1.000
	GIC 5% nanoZrO ₂ -SiO ₂ -HA	18.12 \pm 2.33*	0.000
	GIC 7% nanoZrO ₂ -SiO ₂ -HA	16.26 \pm 2.93	0.148
	GIC 9% nanoZrO ₂ -SiO ₂ -HA	14.61 \pm 2.55	0.999
	GIC 5% nanoZrO ₂ -SiO ₂ -HA	18.12 \pm 2.33*	0.000
GIC 3% nanoZrO ₂ -SiO ₂ -HA	GIC 7% nanoZrO ₂ -SiO ₂ -HA	16.26 \pm 2.93	0.171
	GIC 9% nanoZrO ₂ -SiO ₂ -HA	14.61 \pm 2.55	1.000
GIC 5% nanoZrO ₂ -SiO ₂ -HA	GIC 7% nanoZrO ₂ -SiO ₂ -HA	16.26 \pm 2.93	0.155
	GIC 9% nanoZrO ₂ -SiO ₂ -HA	14.61 \pm 2.55	0.000
GIC 7% nanoZrO ₂ -SiO ₂ -HA	GIC 9% nanoZrO ₂ -SiO ₂ -HA	14.61 \pm 2.55	0.258

*Denotes statistically significant difference at *p*-value \leq 0.05.

In general, all the GIC nanoZrO₂-SiO₂-HA experimental groups had higher *FS* values than the control group. This is probably due to the toughening effect of ZrO₂ and additive effect of HA on the degree of polysalt bridge formation in the GIC matrix. Moreover, ZrO₂ does not dissolve in the distilled water upon immersion for extended periods [36].

There might be higher degrees of acid-base reaction within the structure of the setting cement due to presence of HA resulting in enhanced mechanical properties of GIC [72]. The addition of HA hastens the development of early *FS* of GIC in moist or wet conditions regardless of its form [32]. Another study, also reported a higher *FS* values when porous HA (~28%) was added to GIC as compared to control [73]. However, they reported *FS* of upto ~14 MPa which were much lower than the *FS* values recorded for GIC 5% nanoZrO₂-SiO₂-HA (18.12 \pm 13.88 MPa) in our study.

The increase in *FS* of the GIC could also be due to the fact that the spherical nano Si particles occupy the voids present between the HA rods resulting in increased packing density in the GIC matrix, thereby, reducing crack propagation [34,41,43]. The GIC 5% nanoZrO₂-SiO₂-HA in our study exhibited the highest *FS* values by up to ~26% as compared to the control group 14.33 \pm 3.01 MPa. Similarly, another study also reported the highest *FS* values at 5% addition of HA to GIC [70].

3.2.3. Surface roughness evaluation

In the present study, the average *Ra* values of SR for all tested GICs were within the range of 0.13–0.65 μ m. The lowest SR were reported for GIC 3% nanoZrO₂-SiO₂-HA (0.13 \pm 0.59 μ m) and GIC 5% nanoZrO₂-SiO₂-HA (0.15 \pm 0.29 μ m) among the experimental groups (Table 3). The GIC 5% nanoZrO₂-SiO₂-HA group and control group had nearly identical SR profiles. These values from our study are in agreement with the findings of another recent study that reported the *SR* values (0.20 \pm 0.07 μ m) for a GIC carbomer which contains HA [74]. However, the *SR* profile of our material was smoother (0.15 \pm 0.29 μ m).

Table 3. Surface roughness comparison between the various experimental groups.

Group	Comparison	<i>Ra</i> value \pm SD (μm)	<i>p</i> -value
Control ($0.151 \pm 0.019 \mu\text{m}$)	GIC 3% nanoZrO ₂ -SiO ₂ -HA	0.13 ± 0.059	0.912
	GIC 5% nanoZrO ₂ -SiO ₂ -HA	0.15 ± 0.029	0.997
	GIC 7% nanoZrO ₂ -SiO ₂ -HA	$0.33 \pm 0.062^*$	0.000
	GIC 9% nanoZrO ₂ -SiO ₂ -HA	$0.65 \pm 0.051^*$	0.000
GIC 3% nanoZrO ₂ -SiO ₂ -HA	GIC 5% nanoZrO ₂ -SiO ₂ -HA	0.15 ± 0.029	0.762
	GIC 7% nanoZrO ₂ -SiO ₂ -HA	$0.33 \pm 0.062^*$	0.000
	GIC 9% nanoZrO ₂ -SiO ₂ -HA	$0.65 \pm 0.051^*$	0.000
GIC 5% nanoZrO ₂ -SiO ₂ -HA	GIC 7% nanoZrO ₂ -SiO ₂ -HA	$0.33 \pm 0.062^*$	0.000
	GIC 9% nanoZrO ₂ -SiO ₂ -HA	$0.65 \pm 0.051^*$	0.000
GIC 7% nanoZrO ₂ -SiO ₂ -HA	GIC 9% nanoZrO ₂ -SiO ₂ -HA	$0.65 \pm 0.051^*$	0.000

*Denotes statistically significant difference at p -value ≤ 0.05 .

The highest *Ra* values were recorded for GIC 9% nanoZrO₂-SiO₂-HA ($0.65 \pm 0.05 \mu\text{m}$) followed by GIC 7% nanoZrO₂-SiO₂-HA ($0.33 \pm 0.06 \mu\text{m}$) and they were statistically different from the other groups. This suggests that the composition of the glass content may be responsible for these differences. As GIC 9% nanoZrO₂-SiO₂-HA had the highest filler content and since ZrO₂ is insoluble in the poly acid when compared to HA which dissolves easily, this probably was the reason for its increased *Ra*. Similarly, in another study, higher *Ra* values ($0.64 \mu\text{m}$) were reported for a silver-reinforced GIC, probably because silver particles are also insoluble in poly acids [75].

As mentioned previously, *SR* higher than $0.2 \mu\text{m}$ is likely to increase significantly bacterial adhesion, plaque maturation, thus increasing caries risk [76]. An increase in *SR* can result in alterations in the way light is reflected that in turn can make the material appear opaque. It has been reported that a surface may be considered reflective when its imperfections are well below $1 \mu\text{m}$ [77]. In this study, the GIC 5% nanoZrO₂-SiO₂-HA and GIC 3% nanoZrO₂-SiO₂-HA presented *SR* below $0.2 \mu\text{m}$ value without any efforts at polishing. Therefore, they can be considered to have an ideal surface profile.

4. Conclusion

The morphologic and microstructural characterization of the nanoZrO₂-SiO₂-HA synthesized by a modified one-pot synthesis followed by the evaluation of physico-mechanical properties revealed the following:

- TEM and SEM micrographs confirmed that all particles were in the nano-scale range with spherical ZrO₂ and SiO₂ particles embedded in the voids between rod-shaped HA crystallites.
- FTIR spectra confirmed the presence of functional groups corresponding to each element and the results of EDX and dot mapping revealed that these elements were evenly and homogeneously dispersed throughout the sample with high density patterns visible for ZrO₂, Ca followed by P.
- The incorporation of nanoZrO₂-SiO₂-HA resulted in considerable improvement in the compressive and flexural strengths of cGIC. The GIC 5% nanoZrO₂-SiO₂-HA exhibited a

~30% and ~26% increase in compressive and flexural strength over cGIC, respectively. It also demonstrated surface roughness profile similar to that of cGIC.

Therefore, these nanoceramics can be considered as promising fillers for GIC to be used as a restorative dental material in high stress areas.

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Conflict of interests

All authors declare no conflicts of interest in this paper.

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