Sol–gel dip coating of yttria-stabilized tetragonal zirconia dental ceramic by aluminosilicate nanocomposite as a novel technique to improve the bonding of veneering porcelain

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Abstract: The aim of this in vitro study was to evaluate the effect of silica and aluminosilicate nanocomposite coating of zirconia-based dental ceramic by a sol–gel dip-coating technique on the bond strength of veneering porcelain to the yttria-stabilized tetragonal zirconia polycrystal (Y-TZP) in vitro. Thirty Y-TZP blocks (10 mm × 10 mm × 3 mm) were prepared and were assigned to four experimental groups (n=10/group): C, without any further surface treatment as the control group; S, sandblasted using 110 µm alumina powder; Si, silica sol dip coating + calcination; and Si/Al, aluminosilicate sol dip coating + calcination. After preparing Y-TZP samples, a 3 mm thick layer of the recommended porcelain was fired on the coated Y-TZP surface. Fourier transform infrared spectroscopy (FT-IR), X-ray diffraction (XRD), scanning electron microscopy (SEM), and energy-dispersive X-ray analysis were used to characterize the coating and the nature of the bonding between the coating and zirconia. To examine the zirconia–porcelain bond strength, a microtensile bond strength (µTBS) approach was chosen. FT-IR study showed the formation of silica and aluminosilicate materials. XRD pattern showed the formation of new phases consisting of Si, Al, and Zr in coated samples. SEM showed the formation of a uniform coating on Y-TZP samples. Maximum µTBS values were obtained in aluminosilicate samples, which were significantly increased compared to control and sandblasted groups (P=0.013 and P<0.001, respectively). This study showed that aluminosilicate sol–gel dip coating can be considered as a convenient, less expensive reliable method for improving the bond strength between dental Y-TZP ceramics and veneering porcelain.

Keywords: dip coating, zirconia, porcelain, bond strength

Introduction
Development of high-strength dental ceramics such as yttria-stabilized tetragonal zirconia polycrystal (Y-TZP) has made it possible to fabricate all-ceramic restorations with superior aesthetic properties. The strengthening mechanism known as “transformation toughening”1 has made it possible to use Y-TZP as crown and bridge frameworks instead of metals. Moving from “ceramic fused to metal” toward “all-ceramic” restorations, the techniques for fabricating dental restorations must be revised and mechanisms involving porcelain to zirconia bonding must be well understood.

In dental practice, restorations having zirconia frameworks are fabricated using the computer-aided design and computer-aided manufacturing systems, which may be veneered by a tooth-colored porcelain to improve the esthetics. However, chipping and debonding of the feldspathic porcelain veneer have been one of the most common
concerns in porcelain fused to zirconia restorations. Several strategies have been developed to improve bonding between zirconia and porcelain. Despite several investigations, the bonding mechanism between zirconia and veneering ceramic remains ununderstood. The routine approach usually relies on the roughening of zirconia surface, using sandblasting technique. However, other methods such as conventional silica coating, linear application, and laser surface treatment have been introduced for porcelain to zirconia bonding.

Modification of zirconia surface using silica glasses was introduced for improving the zirconia-to-resin bonding strength. Moreover, this method would make it possible to improve the bonding between zirconia and veneering porcelain, since the feldspathic porcelain may fuse to the silica layer during firing. However, conventional methods for silica coating have been criticized for their adverse effects on mechanical properties (ie, producing surface flaws and phase transformations), as well as their expense. Therefore, other methods, such as sol–gel process, have been used for silica coating of zirconia ceramics.

A sol–gel method provides the powderless processing of glasses and thin films directly from the solution. During this process, an oxide network forms through hydrolysis and condensation of metal alkoxide precursors. Recently, sol–gel process has been used for silica coating of zirconia to improve the resin–zirconia bonding. This procedure is more conservative, less expensive, and requires less space compared to conventional sandblasting and tribochemical processes. Sol–gel dip coating, described well in 1991 by Brinker et al, consists of the withdrawal of a substrate from a sol by gravitational draining, solvent evaporation, and finally performing further condensation reactions, which makes a solid thin film. Compared to other thin film forming techniques, this method is more convenient. In this study, we have used this technique for surface modification of dental zirconia, through formation of a silica or aluminosilicate coating, in order to improve the bonding of porcelain to zirconia. The formation and characterization of the created coating was studied using Fourier transform infrared spectroscopy (FT-IR), X-ray diffraction (XRD), scanning electron microscopy (SEM), and energy-dispersive X-ray analysis (EDS). Microtensile bond strength (μTBS) test was used to quantify the bond between the Y-TZP and the veneering porcelain. In μTBS approach, test specimens are prepared as small beams, having a cross-sectional area of 1×1 mm², while the bonded interface is placed at the middle (Figure 1). The smaller cross-sectional area may decrease the defect concentration in the interface, which in turn lowers the probability of the presence of a defect with a critical size in a crack-opening orientation relative to the applied load.

The aim of this in vitro study was to evaluate the effect of silica and aluminosilicate nanocomposite coating of zirconia-based dental ceramic by sol–gel dip-coating technique on the bond strength of veneering porcelain to Y-TZP.

**Materials and methods**

Frothy Y-TZP blocks (10×10×3 mm³) were prepared (DD Bio ZW iso Zirkonoxid; Dental Direkt GmbH, Spenge, Germany) and sintered as instructed by the manufacturer. The blocks were assigned to four experimental groups (n=10/group): C, without any further surface treatment as the control group; S, sandblasted using 110 μm alumina...
powder under 2.5 bar and the tip distance of 10 mm; Si, silica sol dip coating + calcination; Si/Al, aluminosilicate sol dip coating + calcination.

Sol preparation
Silica sol was prepared as described previously. Briefly, tetraethyl orthosilicate (TEOS) was added to a solution containing nitric acid (HNO₃) (~65%, Sigma-Aldrich Co., St Louis, MO, USA) and ethanol (EtOH) 99.8% (Merck & Co., Inc., Whitehouse Station, NJ, USA) under stirring. After 10 minutes, deionized distilled water (DDW) was added to the solution and the obtained sol was left under stirring for 1 hour in the ambient temperature. The molar ratio of reagents was TEOS:HNO₃:EtOH:H₂O = 1:1.7:6:2.

Aluminosilicate sol was prepared as previously mentioned by Cao et al. Partially hydrolyzed silica sol was first prepared by dissolving TEOS (67 mL) in 100 mL isopropanol and a solution containing water and HNO₃ was added with stirring at room temperature. The stirring was continued for 1 hour; thereafter, 7.36 g Al(NO₃)₃·9H₂O dissolved in 50 mL DDW was added to this sol, with vigorous stirring.

Coating of Y-TZP blocks
To provide a clean surface before coating, the blocks were vigorously washed under running distilled water, placed in an acetone/ethanol ultrasonic bath for 20 minutes, and then air-dried. The Y-TZP blocks were coated either by the silica or aluminosilicate sol using a dip-coating device (80 mm/s, dipping time: 60 seconds) and calcinated at 400°C (5°C/min) for 2 hours.

Porcelain veneering
A thermally compatible veneering porcelain (DD Nature Zr; Dental Direkt GmbH) with matching coefficient of thermal expansion (CTE) was fired on the coated Y-TZP surface under the instructed conditions in a dental porcelain firing furnace (EP3000; Ivoclar-Vivadent, Lichtenstein, Germany). The coated Y-TZP blocks were washed under running distilled water to remove any possible remnants from the coating procedure. A layer of thin porcelain dough (porcelain wash), prepared by mixing the porcelain powder and the supplied liquid, was painted on the Y-TZP surface using a fine brush and fired. In the next step, a polymethyl methacrylate split mold was used to condense a 3 mm thick layer of the porcelain (main body), while the excess water was removed using an absorbent tissue. Finally, the split mold was removed gently, and the porcelain body was fired in recommended condition as presented in Table 1.

Characterization of coatings
FT-IR, XRD, SEM, and EDS were used to characterize the coating and the nature of the bonding between the coating and zirconia. To examine the zirconia–porcelain bond strength, a microtensile approach was chosen.

The FT-IR transmittance spectra of the calcinated prepared gel were obtained in the spectral area of 4,000–600 cm⁻¹, using a Bruker FT-IR extended spectrometer (Bruker Optics Inc., Billerica, MA, USA), with a resolution of 4 cm⁻¹.

SEM imaging (20 kV) and EDS were conducted using a SEM unit (S360; Oxford Co., Cambridge, UK) equipped with an energy-dispersive spectroscopy detector (INCA Energy 300; Oxford Instruments, Oxford, UK), after gold sputtering (Polaron E5200; Watford, Hertfordshire, UK). The phase present in the coatings was analyzed using X’Pert PRO PW 3040/60 XRD (Phillips, Almelo, the Netherlands) with a Cu-Kα monochromatic radiation (λ=1.54178 Å) source.

Microtensile bond strength test
To evaluate the efficacy of the bond between Y-TZP and overlaid porcelain, porcelain-bonded blocks were embedded in transparent polymethyl methacrylate resin and sectioned using a low-speed computerized numerical control cutting machine (Nemo, Mashhad, Iran) under water cooling to fabricate bar-shaped specimens with a 1 mm × 1 mm zirconia–porcelain bond area (Figure 1). The bars were screened under a stereomicroscope using ×40 magnification (Zoom; Blue Light Industry USA Inc., La Habra, CA, USA) for any probable cracks or chipped edges, and 12 sound bar specimens were selected randomly for each group. To simulate the thermal conditions of oral cavity, the bar specimens were subjected to a thermocycling regime (2,000 cycles, 5°C–55°C, dwell time: 20 seconds). After thermocycling, the specimens were mounted on a split holder using cyanoacrylate glue and subjected to μTBS testing using a universal testing machine.

Table 1 Firing conditions of veneering porcelain as instructed by the manufacturer

| Firing step          | Start temperature (°C) | Dry time (minutes) | Heat rate (°C/min) | Final temperature (°C) | Holding time (minutes) | Vacuum | Cooling time (minutes) |
|----------------------|------------------------|--------------------|--------------------|------------------------|------------------------|--------|------------------------|
| Wash firing          | 440                    | 2                  | 45                 | 860                    | 1.5                    | Yes    | 3                      |
| Main body firing     | 440                    | 4                  | 45                 | 830                    | 1.5                    | Yes    | 3                      |
(STM20; Santam, Tehran, Iran) equipped with a 6 kgf load cell under the load rate of 0.5 mm/min (Figure 1).

**Statistical analysis**

The effect of different surface treatments on the zirconia–porcelain µTBS was examined using one-way analysis of variance (ANOVA) procedure, which was followed by Tukey’s post hoc pairwise comparison ($\alpha=0.05$) using SPSS V11.5 software.

**Results**

**Fourier transform infrared spectroscopy**

Figure 2 shows the FT-IR spectra of calcined silica and aluminosilicate. Silica spectrum shows the IR band near 800 cm$^{-1}$, which can be assigned to Si–O–Si symmetric stretching vibrations and a peak at 464 cm$^{-1}$, which is due to O–Si–O bending vibrations. The strong and broad band at 1,082 cm$^{-1}$ with a shoulder at ~1,200 cm$^{-1}$ is in the region that is usually assigned to the transverse and longitudinal optical modes of the Si–O–Si asymmetric stretching vibrations. Moreover, the position and shape of the main Si–O vibrational band at 1,082 cm$^{-1}$ show a stoichiometric silicon dioxide structure.

The wide IR band in the range of 3,000–3,800 cm$^{-1}$ is due to the stretching vibration of H$_2$O molecules, and the band at 1,644 cm$^{-1}$ is related to the corresponding bending.

FT-IR spectrum obtained from aluminosilicate composite indicated that the peak at 1,082 cm$^{-1}$ was shifted to 1,073 cm$^{-1}$, due to the incorporation of aluminum ions, confirming the formation of Al–O–Si bonds. In the presence of Al$^{3+}$ ions as the second neighbor of the Si–O, the Al$^{3+}$ ions will attract the oxygen molecule stronger than Si. Hence, the Si–O–Al asymmetric stretching would be located at a lower wavenumber than Si–O–Si stretching.

**X-ray diffraction**

Figure 3 shows the XRD pattern of silica and aluminosilicate-coated Y-TZP samples after calcination. Analysis of the diffraction pattern of silica-coated sample showed the presence of silicon dioxide and tetragonal structure of zirconium silicate (ZrSiO$_4$). XRD pattern of aluminosilicate sample showed the formation of aluminum silicates (Al$_2$SiO$_5$), predominantly as sillimanite, kyanite, and tetragonal ZrSiO$_4$. Moreover, hexagonal zirconium aluminum oxide with the chemical formula of Zr$_5$Al$_3$O$_{10.5}$ was also detected. Overall, the

![Figure 2 FT-IR spectra of silica (red line) and aluminosilicate (purple line) coating materials. Abbreviation: FT-IR, Fourier transform infrared spectroscopy.](image-url)
Scanning electron microscopy and energy-dispersive X-ray analysis

Figures 4A to D show the SEM images obtained from the surface of specimens after different surface treatments. SEM examination showed the formation of a uniform silica or aluminosilicate layer on the Y-TZP surface (Figure 4C and D). The cross-sectional SEM examination (figures 5A to D) indicated that interfacial silica or aluminosilicate layers were ~10 μm thick (Figure 5C and D).

EDS pattern demonstrated the presence of silicon and oxygen (silica) on the silica-coated samples and a combination of silicon, aluminum, and oxygen on the aluminosilicate-coated surfaces (Figure 4).

Microtensile bond strength

The mean values for μTBS were in order of Si/Al>Si>C>S among the groups in this study. Minimum bonding strength was observed in the sandblasted group (16.74±2.00 MPa), which was significantly lower than in the other groups (Figure 6). The maximum μTBS mean value was obtained in the Al/Si group (24.08±3.44 MPa), followed by the Si group (22.82±2.44 MPa), having no statistically significant difference (P=0.67). The descriptive and analytic statistical data for μTBS are presented in detail in Figure 6. All specimens were fractured in a mixed fashion including the interfacial area and the porcelain part.

Discussion

In this study, we have used a sol–gel dip-coating approach to improve the bonding between the dental (Y-TZP) and the veneering porcelain. Sol–gel approach for Y-TZP coating is a noninvasive method for surface modification to improve the Y-TZP bonding to silicate-based veneering ceramics. Tribochemical methods also can be used for surface modification of zirconia; however, similar to sandblasting in the present study, they may have adverse effect on the bond strength through phase transformation, induced by the collision of silica-coated alumina particles. Sol–gel technique requires less expensive equipment and chemical reagents, compared to the sandblasting unit and silica-coated alumina sand particles.
Figure 4 SEM and EDS (insert) examination of different surface treatments/coatings.  
**Notes:** (A) No treatment (c), (B) sandblasted zirconia surface (z), (C) silica coating (Si), and (D) aluminosilicate coating (Si/Al). Magnification ×600.  
**Abbreviations:** SEM, scanning electron microscopy; EDS, energy-dispersive X-ray analysis.

Figure 5 Cross-sectional SEM imaging of porcelain bonded to zirconia.  
**Notes:** (A) Control group, (B) sandblasted zirconia, (C) Al/Si sol-coated, and (D) Si sol-coated. White arrows indicate the cross-sectional view of aluminosilicate (Al/Si) and silica (Si) coatings. Magnification ×600.  
**Abbreviations:** SEM, scanning electron microscopy; Y-TZP, yttria-stabilized tetragonal zirconia polycrystal; P, porcelain; Si, silica coating; Al/Si, aluminosilicate coating.
A recent study\(^2\) has shown that the veneering porcelain would not chemically interact with the Y-TZP during the firing. However, during sol–gel dip-coating process used in this study, a silica or an aluminosilicate network forms in two steps through a sol–gel process, consisting of a hydrolysis and the consequent polycondensation reactions to form a silica or an aluminosilicate network. Both XRD and FT-IR analysis in this study confirmed the formation of silica and aluminosilicate after heat treatment. During the dip-coating process, a silica or an aluminosilicate layer forms on the Y-TZP surface by the interaction of hydroxyl groups of silica or aluminosilicate particles with the hydroxyl groups of Y-TZP on the surface.\(^3\) This is followed by rapid concentration due to gravitational draining, concomitant with evaporation and condensation reaction.\(^4\) By heat treatment, the diffusion of metallic ions provokes the formation of new intermediate phases, as indicated in this study.

In the present study, a microtensile design was chosen to evaluate the bond strength between porcelain and zirconia. Fischer and Stawarczyk\(^2\) have concluded that shear bond strength design cannot evaluate the interfacial bond strength between Y-TZP and porcelain, since the shear force may develop tensile stresses in porcelain adjacent to the interface, which will result in cohesive fracture in the porcelain substrate.

This study showed that sandblasting might significantly weaken the porcelain–Y-TZP interface. There is a general agreement among studies that sandblasting of Y-TZP has negative effect on the bond strength of veneering porcelain.\(^3,24,26,27\) This destructive effect has been related to the tetragonal to monoclinic solid phase transformation, induced by the collision of alumina particles during sandblasting.\(^10,24,26,27\) Significantly lower CTE in monoclinic form may induce tensile (rather than compressive) stresses at the surface layer, which causes failure in lower shear stresses.\(^25\) Wang et al also have mentioned that the toughness of zirconia/veneer interface without any treatment is significantly higher than that of interfaces subjected to airborne-particle abrasion.\(^9\) Using fracture mechanics, they have suggested another reason for the diminishing effect of sandblasting. Their study showed that sandblasting using 110 \(\mu\)m alumina particles may act as a critical factor in developing flaws in the surface, decreasing the interfacial fracture toughness.\(^9\) Other previous studies also have shown the weaken interface between Y-TZP and layering ceramics after sandblasting surface treatment.\(^28-30\) So, it can be said that although sandblasting increases the surface roughness and bond area, the effect of resulted mechanical retention may not outweigh the influence of the mechanical and thermal residual stresses.\(^8\)

Maximum bond strength values were obtained in aluminosilicate-coated followed by silica-coated samples. The difference between these two groups was not statistically significant; however, in comparison with sandblasted and control groups, the mean values in Al/Si group were significantly increased. The evidence of formation of ZrSiO\(_4\) and zirconium aluminum oxide and intermetallic compounds (ie, a chemical bond) in the present study can

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Figure 6 Mean values and standard deviation for \(\mu\)TBS experiment.

**Note:** Arrows indicate the Tukey’s analysis results. Solid arrows indicate statistically significant difference, dotted arrows indicate no statistically significant difference.

**Abbreviation:** \(\mu\)TBS, microtensile bond strength.
explain the relative higher bond strength in silica- and aluminosilicate-coated samples. Furthermore, the cell volume of ZrSiO$_4$ is four times larger than that of zirconia; hence, a compressive stress would be created in the coating, which can enhance the bonding. In addition, it is possible that after porcelain firing, such new phases appear as the consequence of silica content of the veneering porcelain in the control group. This can be evident in the relatively equal bond strength of silica-coated and control specimens. However, as the bond strength in aluminosilicate-coated samples was significantly increased, it can be postulated that the formation of new phases such as Al$_2$SiO$_5$ and zirconium aluminum oxide may improve the thermal and mechanical compatibility (elastic modulus [EM] and CTE of two adherents) at the interface, which must be studied in more detail in future studies.

The mode of failure must be considered when discussing about an interfacial bond strength. In this study, all specimens were fractured in a mixed fashion including the interfacial area and the porcelain part. The failure pattern is related to factors that cause increased stress. These factors are the difference and mismatch of the EM and CTE of porcelain and zirconia, as well as the test methodology.$^{31-33}$ The difference in EM and CTE of porcelain and zirconia is recognized as the determinant factor for inducing residual stresses in the adherent.

Studies about the mechanics of veneered Y-TZP debonding are under progress yet, although Nishigori et al.$^8$ reported that the fracture initiates from the veneer ceramic. In this study, the veneering porcelain may integrate to the silica and aluminosilicate coating, due to the sintering of silica content of porcelain and coating, while the newly formed interface between coating and Y-TZP remains the weakest zone of porcelain–Y-TZP junction. Similar failure pattern with different mean bond strength levels between coated and uncoated groups may imply that in the present study, the fracture had initiated in the interfacial zone, but not in the porcelain, which must be studied in more detail in the future.

**Conclusion**

Considering the limitations of this in vitro study, the results showed that aluminosilicate sol–gel dip coating can be considered as a convenient, less expensive, reliable method for improving the bond strength between Y-TZP ceramics and the veneering porcelain. Further studies are needed to evaluate the effect of different variables such as the calcination and dip-coating processes on the quality of porcelain–Y-TZP bond.

In addition, the fracture mechanics of bond failure must be studied in more detail.

**Acknowledgment**

This manuscript is a part of thesis for postgraduate degree in prosthodontics. The research was funded by the Vice Chancellor for Research and Technology, Mashhad University of Medical Sciences, Mashhad, Iran.

**Disclosure**

The authors report no conflicts of interest in this work.

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