Effects of different surface finishing protocols for zirconia on surface roughness and bacterial biofilm formation

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PURPOSE. Surface finishing of a zirconia restoration is essential after clinical adjustment. Herein, we investigated the effects of a surface finishing protocol for monolithic zirconia on final roughness and bacterial adherence. MATERIALS AND METHODS. Forty-eight disk-shaped monolithic zirconia specimens were fabricated and divided into four groups (n = 12) based on initial surface treatment, finishing, and polishing protocols: diamond bur+polishing bur (DP group), diamond bur+stone grinding bur+polishing bur (DSP group), no diamond bur+polishing bur (NP group), and no diamond bur+stone grinding bur+polishing bur (NSP group). Initial and final surface roughness was measured with a profilometer, and shown using scanning electron microscope. Bacterial adhesion was evaluated by quantifying Streptococcus mutans in the biofilm. Kruskal–Wallis and Mann-Whitney U tests were used to compare results among groups, and two-way analysis of variance was used to evaluate the effects of grinding burs on final roughness (α=0.05). RESULTS. The DP group had the highest final Ra value, followed by the DSP, NP, and NSP groups. Use of the stone grinding bur as a coarse-finishing step significantly decreased final Ra values when a diamond bur was used (P<0.001). Omission of the stone grinding bur increased biofilm formation on specimen surfaces. Combining a stone grinding bur with silicone polishing burs produced the smallest final biofilm values, regardless of the use of a diamond bur in initial surface treatment. CONCLUSION. Coarse finishing of monolithic zirconia with a stone grinding bur significantly decreased final Ra values and bacterial biofilm formation when surfaces had been roughened by a diamond bur. [J Adv Prosthodont 2019;11:41-7]

KEYWORDS: Zirconia; Dental finishing; Dental polishing; Biofilm; Bacterial adhesion

INTRODUCTION

Monolithic zirconia restorations are alternatives to layered zirconia restorations because of their decreased potential for chipping of the porcelain veneer.¹ In a monolithic zirconia restoration, zirconia is directly exposed to the oral environment and must be polished following occlusal and axial adjustments.² Surface smoothing of dental restorations is essential for reducing plaque accumulation, improving patient comfort, preventing wear of the antagonist enamel, and improving the aesthetics of restorations.³,⁶ The effectiveness of a finishing and polishing system on dental material is evaluated by measuring the achieved surface roughness.² Restorations with surface roughness values > 0.2 µm
(Ra) could trigger bacterial plaque accumulation, which is the primary cause of gingival inflammation and secondary caries. Bacteria adhere to restorations and natural teeth through a four-phase process. The initial phase begins with transport of bacteria toward the surface. In the second phase, bacteria adhere to the surface through a combination of Van der Waals forces and electrostatic repulsive forces. The intermediate attachment phase involves the adherence of bacteria to complementary sites on the substrate surface. In the final phase, bacteria colonize on the surface and form a biofilm via proliferation of already-colonized bacteria and/or by continuous adhesion of additional salivary bacteria. Initial adhesion of bacteria to the tooth surface is a critical step in bacterial plaque formation. The irregular geometry of a rough surface shelters bacteria from shear forces, giving them more time to interact with the surface and establish strong adhesion.

Monolithic zirconia restoration fabricated from a solid block of zirconia has better mechanical properties and superior fracture strength than bi-layered restorations due to the uniform structure. The high strength and hardness of zirconia require specific surface treatment methods. Coarse-grit and fine-grit diamond rotary burs are often used at high speed for polishing monolithic zirconia because diamond burs are effective for extensive grinding of zirconia. Ho et al. stated that the use of diamond bur increased the flexural strength of zirconia, whereas Kosmac et al. concluded that use of the coarse diamond bur reduced the strength of the material due to surface flaws and microcracks. Therefore, specific zirconia finishing and polishing systems that incorporate instruments with improved diamond particle coatings have been developed. Use of zirconia-specific grinding burs are associated with higher grinding efficiency and lower heat generation than conventional stone burs.

There have been reports in the literature on the importance of using specific stone burs on zirconia. However, the effects of stone burs in relation to use of different initial surface treatments on final roughness have not been fully clarified. The purpose of the present study was to examine whether use of diamond and stone burs on zirconia affects its final roughness and bacterial adhesion properties. The null hypothesis was that the surface grinding protocol for zirconia does not influence the final roughness or bacterial biofilm formation.

**MATERIALS AND METHODS**

Figure 1 presents the study design. Forty-eight disk-shaped monolithic zirconia specimens were designed using computer software (CATIA V5R19, Dassault Systemes, Velizy-Villacoublay, France) and were manufactured with a 5-axis milling machine (Ceramill Motion 2, Amann Girrbach, Koblach, Austria) using zirconium dioxide blocks (Prettau, Zirkonzahn, Bruneck, Italy). Afterwards, sintering process of specimens was conducted using a dedicated sintering furnace (Zirkonofen 600/V2, Zirkonzahn). For the initial surface treatment, 24 specimens (15 mm in diameter, 8 mm in thickness) were randomly selected and ground for 10 s under water-cooling conditions with a coarse-grit diamond bur.
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The resulting surface roughness was measured with a contact surface profilometer (Surftest SY-400, Mitutoyo, Kawasaki, Japan) under a constant load of 4 mN at a speed of 0.25 mm/s and a measurement range of 0.8 mm. The Ra value of each specimen was determined by averaging nine measurements: three repetitions of three traces.

*S. mutans (ATCC 25175) was used to assess biofilm formation. S. mutans was maintained in a brain heart infusion (BHI) medium and grown under aerobic conditions. An in vitro biofilm formation assay was performed in accordance with published protocol. The specimens were cleaned with 70% alcohol and sterilized by UV light for 1 hour before the biofilm formation test. Briefly, S. mutans colonies were inoculated into BHI-1% sucrose broth and incubated overnight. The culture broth was inoculated into 2 mL of the same liquid medium to reach 1% volume/volume in a 12-well polystyrene plate containing each specimen. After incubation for 16 hours, the streptococcal broth was removed, and the specimens were washed three times with sterile phosphate buffered saline solution to remove loosely attached biomass. For biofilm quantification, 200 μL of crystal violet solution (0.2 % w/v in 10% ethanol) was added to each specimen, followed by incubation for 1 hour. After washing with phosphate-buffered saline solution three times, the specimen was air-dried. Crystal violet retained by the streptococcal biofilm was redissolved in 200 μL acidic solvent (10% acetic acid in distilled water); the absorbance was determined with a microplate reader (Molecular Devices, San Jose, CA, USA) at 595 nm.

For scanning electron microscope (SEM) analysis, the specimens were attached to an aluminum metal platform with a double sided adhesive tape. Then the edges of the specimens were painted with carbon paint (Pelco Colloidal Graphite, Ted Pella, Redding, CA, USA). After coating the specimen with spotted platinum with ion sputter (E-1030, Hitachi, Tokyo, Japan), they were analyzed with SEM (SU8230, Hitachi) at ×200 magnification.

All statistical analyses were conducted using the IBM SPSS Statistics v22.0 statistical software package (IBM Corp., Armonk, NY, USA). Surface roughness for each group was expressed as mean ± standard deviation. Kruskal-Wallis and Mann-Whitney U tests were used to compare the effects of stone grinding bur use among all subgroups. Two-way analysis of variance (ANOVA) was performed to investigate relationships between surface preparation and use of a stone grinding bur. Results were visualized using interaction plots. Statistical significance was set at P < .05.

Table 1. Instrument codes and revolutions per minute used for finishing and polishing

<table>
<thead>
<tr>
<th>Category</th>
<th>Diamond bur</th>
<th>Stone grinding bur</th>
<th>Silicone polishing bur</th>
<th>Fine silicone polishing bur</th>
</tr>
</thead>
<tbody>
<tr>
<td>Instrument codes*</td>
<td>V847KR</td>
<td>8001.050.HP</td>
<td>3041.HP</td>
<td>30041.HP</td>
</tr>
<tr>
<td>Revolutions per minute</td>
<td>200,000</td>
<td>12,500</td>
<td>20,000</td>
<td>10,000</td>
</tr>
</tbody>
</table>

*As provided by manufacturers

Fig. 2. Microscopic views of burs at ×300 magnification. (A) Diamond bur, (B) Stone grinding bur, (C) Silicone polishing bur, (D) Fine silicone polishing bur.
RESULTS

The initial surface roughness values were $Ra = 1.07 \ \mu m$ for the diamond bur-treated group and $Ra = 0.42 \ \mu m$ for the non-treated group. Table 2 presents the roughness of zirconia specimens by surface treatment. In the initial measurements, the use of the diamond bur at high speed significantly increased Ra values. After different finishing and polishing protocols, the DP group had the highest final Ra value, followed by the DSP, NP, and NSP groups. Addition of a finishing step using a stone grinding bur significantly decreased the final Ra values for diamond bur-treated specimens; however, the effect was not statistically significant when the specimens had not been treated with the diamond bur ($P = .70$). Two-way ANOVA revealed a strong interaction between the use of the diamond bur in the initial surface treatment and the use of the stone grinding bur in the finishing step on the final Ra ($P < .001$) (Table 3).

In vitro streptococcal biofilm masses on the surfaces of the specimens, as determined by crystal violet staining, were compared (Fig. 3). In general, biofilm formation of S. mutans significantly increased on the surfaces of specimens treated with the diamond bur, especially when use of a stone grinding bur was omitted ($P < .001$). The combination of a stone grinding bur and a silicone polishing bur produced small final biofilm mass.

![Figure 3](image)

**Fig. 3.** Biofilm formation on zirconia specimen from each group after finishing and polishing. Black lines indicate initial biofilm formation for diamond bur-treated and nontreated specimens.

<table>
<thead>
<tr>
<th>Source</th>
<th>Sum of squares</th>
<th>d.f.</th>
<th>Mean square</th>
<th>F</th>
<th>P</th>
</tr>
</thead>
<tbody>
<tr>
<td>Diamond bur</td>
<td>2.417</td>
<td>1</td>
<td>2.417</td>
<td>331.675</td>
<td>&lt; .001</td>
</tr>
<tr>
<td>Stone bur</td>
<td>.189</td>
<td>1</td>
<td>.189</td>
<td>25.907</td>
<td>&lt; .001</td>
</tr>
<tr>
<td>Diamond bur × Stone bur</td>
<td>.121</td>
<td>1</td>
<td>.121</td>
<td>16.608</td>
<td>&lt; .001</td>
</tr>
<tr>
<td>Error</td>
<td>.321</td>
<td>44</td>
<td>.007</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Total</td>
<td>16.583</td>
<td>48</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Corrected total</td>
<td>3.047</td>
<td>47</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

**Table 2.** Means ± SDs for surface roughness (Ra) of specimens by surface treatment protocol

**Table 3.** Two-way ANOVA for surface roughness
For SEM images, the specimen treated with a diamond bur alone showed sharp groove patterns (Fig. 4A), as opposed to the other specimens (Fig. 4B, 4C). The specimen of DSP group showed more evenness of the surface compared to rounded edges of the large grooves of DP group (Fig. 4B, 4C).

DISCUSSION

This study was conducted to evaluate the effects of surface finishing of monolithic zirconia with a coarse stone grinding bur on final surface roughness and bacterial adhesion. Based on our results, the null hypothesis was rejected because the final surface roughness and biofilm formation were significantly different, depending on the coarse-finishing protocols used.

A zirconia-specific stone grinding bur was used in this study in a coarse-finishing step for specimens with different initial surface roughness values. When initial surface treatment is performed using a high-speed diamond bur on monolithic zirconia specimens, a very rough surface is produced. The finishing and polishing procedure is used to achieve a smooth and regular surface for the restoration. However, deep scratches created by the diamond bur are difficult to remove with silicone polishing burs because of the extreme strength and hardness of zirconia material. Finishing and polishing instruments comprise of three major components: primary abrasive, supplemental abrasive, and abrasive-fixing substance. Among these, the primary abrasive is the main factor determining the grinding efficiency. Since zirconia has a value of 9 on the Mohs hardness scale, diamond grinding burs for occlusal adjustment have been manufactured using a coarse diamond grain size and serve as the primary abrasive. Silicone polishing burs, on the other hand, contain very fine diamond grains; thus, these burs exhibit significantly lower grinding efficiency. Therefore, use of a stone grinding bur may be considered an important intermediate step in filling the gap between the extremely rough surface of the diamond bur and the fine surfaces of silicone polishing burs. Previous studies have reported that use of zirconia stone grinding burs for zirconia materials plays an important role in removing deep grooves and achieving a finer surface texture for subsequent polishing steps. Therefore, addition of a coarse-finishing step using a stone grinding bur is essential and should not be omitted in clinical practice, particularly after extensive reduction of the zirconia restoration is accomplished with a coarse-grit diamond bur.

Surface roughness is considered the most important factor in oral biofilm formation. A correlation between surface roughness and bacterial adhesion was also observed in this study. Biofilm formation was substantially reduced after stone grinding, independent of whether specimens had been roughed by diamond burs. Interestingly, when use of the stone grinding bur was omitted, use of silicone polishing burs alone increased the biofilm thickness in specimens treated with a diamond bur. This phenomenon may be explained by the surface topography of treated zirconia (Fig. 4, Fig. 5). The initial surface treatment process that used a coarse-grit diamond bur generated deep, sharp grooves on the surface. The silicone bur is not suitable for grinding irregular macrostructures because of the low hardness and fine grit of the diamond particles in the instrument. Moreover, silicone burs cannot reach the inner sides of the narrow deep grooves, but they can be used to round the sharp outer edges of these grooves. Although this partial surface alteration generally decreases the Ra value of polished zirconia, this surface structure may provide more favorable conditions for bacterial adhesion. Han et al. also reported the similar bactericidal effects in the surface with sharp grooves and pits. These results confirm the clinical importance of sequential finishing and polishing regimens.

There are various clinical and technical factors that determine the final roughness of zirconia. The roles of various polishing systems and sintering processes in the final roughness of zirconia need to be evaluated in future studies. Moreover, the effects of grinding direction, time and pressure, and use of water coolants are also important clinical factors that require further investigation.

Fig. 4. Scanning electron microscope images of diamond bur-treated specimens. (A) Surface after use of diamond bur alone, (B) Surface after use of diamond and polishing burs, (C) Surface after use of diamond bur, stone grinding bur, and polishing bur.
CONCLUSION

Within the limitations of this study, a coarse-finishing step using a stone grinding bur significantly decreased the final Ra values of monolithic zirconia when the surface had been roughed by a diamond bur. Biofilm formation of S. mutans increased on the surfaces of specimens treated with a diamond bur, especially when use of a stone grinding bur was omitted.

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Fig. 5. Schematic image of bacterial adhesion to zirconia; heavy bacterial inhabitation is observed in the partially altered surface treated by diamond and polishing burs. (A) Surface after use of diamond bur alone, (B) Surface after use of diamond and polishing burs, (C) Surface after use of diamond bur, stone grinding bur, and polishing bur.