Effects of Zirconia Addition on Fracture Toughness and Bending Strength of Dental Porcelains

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Zirconia dispersed composite porcelains with glass and aluminous porcelain as matrix were prepared as models of dental porcelains. The bending strength and fracture toughness of the composite porcelains were examined. The bending strength and fracture toughness of composite porcelains containing 50 wt% zirconia were 20 to 80 % greater than in glass alone. However, bending strength and fracture toughness decreased upon the addition of zirconia at more than 50 wt%. Moreover, in the case of aluminous porcelain as matrix, fracture toughness increased to a maximum value of 2.6 MPa m$^{1/2}$ by addition of 23 wt% zirconia, twice the toughness of glass alone. On the other hand, no increase of bending strength was observed in this case. Deflection and bowing of cracks as well as microcracking effects were related to these increases of mechanical properties in zirconia dispersed composite porcelains.

Key words: Porcelain, Zirconia, Mechanical property

INTRODUCTION

Porcelain has many applications in restorative dentistry including denture teeth, jacket crowns, porcelain-fused-to-metal bridge work, and inlays. Porcelain possesses excellent esthetics and resists mechanical wear and chemical degradation extremely well. However, conventional dental porcelain is weak with respect to shearing as it essentially is a brittle material. In order to compensate for this brittleness, various methods, including baking on metal and production in or addition of crystal to the glass matrix have been proposed. Garvie et al. reported high strength zirconia ceramics made from room-temperature-stable tetragonal zirconia in 1975. Since then fracture resistant ceramics which can overcome the disadvantages of brittleness have been the subject of considerable interest. In particular, significant attention has focussed on partially stabilized zirconia ceramics which absorb breaking energy by the transformation of tetragonal zirconia to monoclinic zirconia and possess high fracture toughness. In addition, much attention has been devoted to zirconia dispersed composite ceramics in which zirconia particles are dispersed in alumina and silicon nitride ceramics. It has been observed that deflection and curving of cracks as well as microcracking can be caused by the phase transformation of zirconia and can absorb breaking energy.

In 1984, we reported on the mechanical properties of composite porcelains containing dispersed zirconia particles in glass matrix as a model of dental porcelain. R. Morena et al. also reported the fracture toughness and bending strength of dental porcelains to increase 30 to 35 % upon the addition of zirconia.
In the present investigation, composite porcelains containing various kinds of zirconia dispersed into 2 kinds of matrices will be discussed as models of dental porcelain. A technique for producing controlled microcracks was applied to the glass matrix and the fracture toughness and bending strength of various ceramics are reported.

MATERIALS AND METHODS

Specimen Preparation

Two kinds of glass were employed as matrix materials for porcelain models: borosilicate (4 wt% Na₂O-13 wt% B₂O₃-81 wt% SiO₂) and soda-lime glass (14 wt% Na₂O-11 wt% CaO-73 wt% SiO₂) as summarized in Table 1. In addition, 3 different mixtures of borosilicate glass and electro-fused alumina were used as models of aluminous porcelain (Table 1). Five kinds of electro-fused zirconia powders* with a mean particle size of 0.4 μm were used as dispersion particles (Table 2). Glass and alumina powders used as matrices were crushed to a fine powder** in an alumina cell and their mean particle sizes were measured by an automatic particle size distribution measuring system***. Matrix materials and zirconia were then mixed in a ball mill for 1 h without a dispersing agent. Pressures of up to 20 MPa were applied to dry specimens in metal dies using a mechanical press. After press

<table>
<thead>
<tr>
<th>Table 1</th>
<th>Various matrices for zirconia dispersed composite system</th>
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<tr>
<td>Code</td>
<td>Sodium borosilicate glass (wt%)</td>
</tr>
<tr>
<td>----------</td>
<td>---------------------------------</td>
</tr>
<tr>
<td>G₁</td>
<td>100</td>
</tr>
<tr>
<td>G₂</td>
<td>0</td>
</tr>
<tr>
<td>GA₅</td>
<td>50</td>
</tr>
<tr>
<td>GA₆</td>
<td>60</td>
</tr>
<tr>
<td>GA₇</td>
<td>70</td>
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</tbody>
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<table>
<thead>
<tr>
<th>Table 2</th>
<th>Various zirconia specimens used in this study</th>
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<tr>
<td>Zirconia</td>
<td>Code</td>
</tr>
<tr>
<td>ZrO₂</td>
<td>Zm</td>
</tr>
<tr>
<td>ZrO₂ + Y₂O₃ (6.5 wt%)</td>
<td>ZYc</td>
</tr>
<tr>
<td>ZrO₂ + Y₂O₃ (4.1 wt%)</td>
<td>ZYt</td>
</tr>
<tr>
<td>ZrO₂ + CaO (5.5 wt%)</td>
<td>ZCc</td>
</tr>
<tr>
<td>ZrO₂ + CaO (3.5 wt%)</td>
<td>ZCt</td>
</tr>
</tbody>
</table>

* Tateho Chemical Industries Co., Okayama, Japan
** T-100 model Vibrating sample mill, Heiko Seisakusyo, Ltd., Fukushima, Japan
*** CAPA-300 Centrifugal automatic particle analyzer, Horiba, Ltd., Kyoto, Japan
drying, compacts were sintered at a rate of 5°C·min⁻¹ in atmosphere.

**Dense Temperature**

The linear shrinkage of pressed samples of 8φ × 10 mm was measured by constant heating at a rate of 5°C·min⁻¹ to 1300°C. Dense temperature was determined by linear shrinkage vs the temperature curve. The density of the sintered porcelains was measured by Archimedes' method.

**SEM Observations**

The microstructure of the destroyed surface of sintered materials was also observed with a scanning electron microscope.

**Mechanical Properties**

Test pieces for the measurement of mechanical properties were made by grinding the surface of a test piece flat and smooth using #700 diamond rap. The sizes of the test pieces were 2mm × 6mm × 15mm and 4mm × 6mm × 15mm for the bending strength and fracture toughness tests, respectively. In measuring fracture toughness (K_Ic), 10 specimens at each ZrO₂ content were notched to a depth of 1.0 mm and thickness of 0.2 mm with diamond wheel at a slow speed. Unnotched specimens were used to measure bending strength. Both fracture toughness and bending strength specimens were tested for 3-point bending over a 15 mm span at a crosshead speed of 0.5 mm·min⁻¹. The fracture toughness of the sintered materials was calculated using the Single-Edge-Notched-Beam (SENB) method.

**X-Ray Diffraction**

The X-ray powder diffraction patterns (XRD) of sintered materials were recorded. The diffractometer system employed used Ni-filtered CuKα radiation. The 2θ range from 10° to 70° was covered at a scanning speed of 1.0 degree min⁻¹ at 30 kV and 10 mA.

**RESULTS AND DISCUSSION**

**Zirconia Dispersed Glass Porcelain**

The dense temperatures of borosilicate and soda-lime glass were 800°C and 730°C, respectively. The sintering temperature of these glasses rose as the amount of added zirconia increased. In the case of composite porcelains containing 50 wt% zirconia, dense temperatures were from 880°C to 1050°C. On the other hand, composite porcelains containing 60 wt% zirconia showed rather high dense temperatures, from 1100°C to 1170°C. Zircon (SiZrO₄) was produced by the reaction between zirconia and a glass component, i.e. silica, at high sintering temperatures. In the case of stabilized and partially stabilized zirconia with calcia (ZCc, ZCt), the amount of cubic-zirconia decreased with sintering. On the other hand, partially stabilized zirconia with yttria (ZYt) in the cubic-zirconia phase increased with sintering. The binding strengths and fracture toughesses (K_Ic) of the composite ceramics containing 50 wt% zirconia are summarized in Table 3. In the case of borosilicate glass, the dense temperatures and binding strengths were higher than those of soda-lime glass.

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* S-700 Scanning electron microscope, Hitachi Co., Tokyo, Japan
** MC-411 Crystal cutter, Maruto Co., Ltd., Tokyo, Japan
*** AGS-500A Autograph, Shimazu Co., Kyoto, Japan
**** ADG-301, Toshiba Co., Tokyo, Japan
Table 3  Bending strength and fracture toughness values for glass-zirconia composite systems

<table>
<thead>
<tr>
<th>Glass (50wt%) Zirconia</th>
<th>G₁</th>
<th></th>
<th></th>
<th></th>
<th>G₂</th>
<th></th>
<th></th>
<th></th>
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</thead>
<tbody>
<tr>
<td></td>
<td>Zm</td>
<td>ZYc</td>
<td>ZYt</td>
<td>ZCc</td>
<td>ZCt</td>
<td>Zm</td>
<td>ZYc</td>
<td>ZYt</td>
</tr>
<tr>
<td>Bending strength</td>
<td>70</td>
<td>134</td>
<td>99</td>
<td>128</td>
<td>118</td>
<td>130</td>
<td>112</td>
<td>92</td>
</tr>
<tr>
<td>(MPa)</td>
<td>8</td>
<td>8</td>
<td>4</td>
<td>10</td>
<td>11</td>
<td>17</td>
<td>9</td>
<td>2</td>
</tr>
<tr>
<td>Fracture toughness</td>
<td>1.72</td>
<td>2.03</td>
<td>1.85</td>
<td>2.03</td>
<td>2.22</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>K₁c (MPa • m¹/²)</td>
<td>(0.18)</td>
<td>(0.10)</td>
<td>(0.11)</td>
<td>(0.03)</td>
<td>(0.49)</td>
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<td></td>
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</tr>
</tbody>
</table>

( ) : SD

matrix (G₁), dispersion of unstabilized zirconia (Zm) resulted in a minimum bending strength of 70 MPa, similar to that of G₁ glass alone, at 60 MPa. The highest values of bending strength were achieved with yttria or calcia stabilized zirconia (ZYc, ZCc), at about 134 and 128 MPa, respectively. These values are twice that of matrix alone. In the 50 wt% partially stabilized zirconia system, bending strengths were 99 and 118 MPa, respectively. These results show that the bending strength of zirconia composite porcelains of G₁ glass increase with increased in cubic phase zirconia.

Differing results were obtained when soda-lime glass (G₂) was used as matrix. In such cases, the maximum bending strength was observed in composite porcelains with unstabilized zirconia (Zm) at 130 MPa and bending strengths between 90 and 120 MPa were obtained with other zirconia composite porcelains. One reason mechanical strengths may differ between the 2 glass matrix groups depending on the kind of zirconia may be that the glass and zirconia components are different and differing reactions at firing temperature may, in turn, result in differentes of mechanical strength. Fracture toughness (K₁c) of all G₁ glass-zirconia composite ceramics was higher than in sintered materials of glass alone, at 1.32 ± 0.13 MPa • m¹/². In the case of composite ceramics containing 50 wt% zirconia, fracture toughness as high as 2 MPa • m¹/² was obtained. In the case of composite ceramics containing hard crystal, e.g. glass-alumina composite porcelains, high mechanical strengths and toughnesses were obtained by the deflection and bowing of main cracks induced by destruction. In this way, both bending strength and fracture toughness increased. In this experiment as well, both bending strength and fracture toughness was much greater than that of ceramics with glass alone. Thus, main cracks causing destruction may be deflected by dispersed particles, and the crack front proceeding around the latter, requiring more energy. The bending strength and fracture toughness of composite porcelains containing more than 50 wt% zirconia were lower than those of composite porcelains containing only 50 wt% zirconia (Table 4). The reason for this decreasing strength and toughness may have to do with the crystallization of zircon in composite porcelains.

Zirconia Dispersed Aluminous Porcelain

The bending strength of G₁ glass-alumina (GA) composite porcelains increases with increases in added alumina, reaching a maximum of 180 MPa at 50 wt% alumina. Fracture toughness changes in a similar fashion as bending strength, as shown in Fig. 1, reaching a maximum of 2.11 MPa • m¹/² when 50 wt% alumina is added. The shrinkage curves of powder compacts of aluminous porcelain containing 40 wt% alumina (GA₄) and various
additions of ZYt-zirconia are summarized in Fig. 2. Shrinkage curves and sintering temperatures as well increase with zirconia addition. Similar phenomena were observed for GA3 and GA5 matrix as well as for other zirconia. Crystals other than raw materials were not detected in composite porcelains fired below 1100°C, though zircon, produced by a reaction between zirconia and silica, was detected by X-ray diffraction analysis in composite porcelains fired above 1200°C. The amount of zircon was affected by the kinds of zirconia added. The porosity of the glass-alumina-zirconia composite porcelains increased with increases of added zirconia. The typical porosity of the composite porcelains are summarized in Fig. 3 for

![Image](image_url)
Fig. 2 Changes in firing shrinkage curves of powder compacts consisting of glass-alumina matrix (GA₄) and zirconia (ZYt), with temperature raised continuously.

Fig. 3 Relation between porosity and zirconia content of sintered specimens consisting of glass-alumina matrix (GA₅) and zirconia (Zm).
the GA₅-zirconia system and the bending strengths of the GA₅-Zm composite porcelains are summarized in Fig. 4. The bending strength of GA₅ matrix alone was about 180 MPa but decreased upon zirconia addition to about 110 MPa at 33 wt% zirconia. One reason bending strength decreases may be that porosity increases as zirconia is added³⁰) (Fig. 3). On the other hand, the fracture toughness of the GA₅-Zm composite porcelains was constant up to 16 wt zirconia and equal to that of matrix alone as shown in Fig. 5. At 23 wt% zirconia, a maximum value of 2.57 MPa·m⁰·⁵ was obtained for fracture toughness, and further addition of zirconia over 23 wt% caused fracture toughness to decrease (Fig. 5). In order to observe the microstructure of the surface of destroyed composite porcelains, scanning electron microscopy (SEM) was conducted. As shown in Fig. 6, deflection and bowing of cracks were seen to occur before destruction. However, microcracks due to transformation of zirconia phase were not observed by SEM. Increases in fracture toughness occur with the microcracking effect as maximal fracture toughness is observed at a zirconia content accompanied by a decrease in bending strength¹⁰,¹²). The microcracks in zirconia dispersed ceramics are formed by the expansion of zirconia particles during the tetragonal → monoclinic phase transformation. Toughening with the microcracking effect is explained by fracture energy absorption in a microcrack zone, as microcrack opening and branching of main cracks causing destruction⁷,⁸,¹⁰,¹²,²³,²⁷,³¹). The reason for the decreasing fracture toughness in the case of GA₅-Zm composite system containing more than 23 wt% zirconia may have to do with existence of large cracks that the microcracks join up between the particles in a composite

Fig. 4 Relation between bending strength and zirconia content of sintered composite systems consisting of glass-alumina matrix (GA₅) and zirconia (Zm).
porcelain\textsuperscript{10,12}. When yttria partially stabilized zirconia was used for the GA\textsubscript{4}-zirconia composite system, addition of ZYt had little or no effect on bending strength as shown in Fig. 7. The porosity of GA\textsubscript{4}-ZYt composite porcelains increased with increases in the amount of zirconia, but porosity values were lower than those of the GA\textsubscript{5}-Zm composite system. The fracture toughness of GA\textsubscript{4}-ZYt composite porcelains increased when compared to GA\textsubscript{4} matrix alone, the latter's value being 2.0 MPa·m\textsuperscript{1/2}, while the former's value reached 2.2 MPa·m\textsuperscript{1/2} at 16 wt% ZYt content (Fig. 8). The fracture toughness of GA\textsubscript{4}-ZYt composite system had a maximal point, smaller than that of GA\textsubscript{5}-Zm system (Fig. 5). This reason may as well be supplied by the microcracking effect, similar to the case of GA\textsubscript{5}-Zm system. These results may show that the dispersion of unstabilized zirconia is more effective than that of partially stabilized zirconia in an aluminous porcelain matrix.

CONCLUSIONS

Zirconia dispersed composite porcelains with glass and aluminous porcelain as matrix were prepared as models of dental porcelain in order to assay increases in the mechanical properties of dental porcelain. The bending strength and fracture toughness of glass matrix porcelains increased 20\% to 80\% upon dispersion of 50 wt\% zirconia.

In the case of aluminous matrix porcelains, fracture toughness increased 2 times, to 2.6
Fig. 6 Microstructure (SEM) of sintered composite systems consisting of glass-alumina matrix (GA₅) and zirconia (Zm).
Fig. 7  Relation between bending strength and zirconia content of sintered composite systems consisting of glass-alumina (GA₄) and zirconia (ZYt).

Fig. 8  Relation between fracture toughness (K_{IC}) and zirconia content of sintered composite systems consisting of glass-alumina (GA₄) and zirconia (ZYt).
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MPa·m$^{1/2}$, upon dispersion of 23 wt% zirconia. Bending strength, however, showed no change or even a decrease as zirconia was added.

Such increases in mechanical strength and fracture toughness seem to result from the deflection and bowing of cracks, as well as from the microcracking effect.

REFERENCES


歯科用陶材の破壊靭性と強さに及ぼすジルコニア添加の影響
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徳島大学歯学部歯科理工学教室

歯科用陶材のモデルとしてガラスまたはアルミナス陶材をユニプレックスに用い，ジルコニア分散陶材を作製した。その複合陶材の曲げ強さと破壊靭性（Kic）について検討した。

ガラスマトリックスに各種ジルコニアを50 wt %添加した複合陶材の曲げ強さと破壊靭性はガラス単独の場合に比較して20～80 %高くなった。50 wt %以上ジルコニアを添加すると強さおよび靭性とともに低下する傾向を示した。アルミナス陶材をマトリックスとした場合では曲げ強さの向上はみられなかったが，破壊靭性については向上するものが数種出現した。ジルコニア含有量16～23 wt %時に極大値を示し，靭性値は最高でガラス焼結体の2倍の約2.6 MPa・m1/2が得られた。

各種マトリックスにジルコニアを分散した複合陶材の強さや靭性が高くなるのはクラックの消滅と偏向またはマイクロクラック効果と考えられた。

焼付用陶材のガラス転移温度域での粘度について
浅岡和三，今 政幸，桑山則彦
徳島大学歯学部歯科理工学教室

陶材の粘度は金属焼付陶材の適合性，すなわち合金と陶材の好ましい物理的・機械的性質の組み合わせ，焼成方法と残留応力の関係を決める重要な因子である。一般に，ガラス転移温度域での陶材の粘度はアレニウス式により表示される。この粘度が，応力を加えながら一定速度で加熱したときに，陶材が膨張から収縮へ変化する温度（変形温度）を測定することにより求まることを，粘弾性モデルより導き，具体的な測定方法を明らかにした。

市販焼付用歯冠金制陶材6種，オペレック陶材6種について活性化エネルギーを上記の方法により測定した。その結果をもとに焼付用陶材の粘度，加熱時の変形とガラスの特性温度の関係を比較検討した。また，陶材の加熱速度と熱膨張係数の関係について調べた。膨張係数から求めたガラス転移域の下限温度が粘度より計算された重点と陶材の焼成温度での粘度がガラスの軟化温度の粘度に一致した。ここで示された測定方法が簡易で信頼性の高い方法であると結論された。

低屈折率を有するジメタクリレートの応用による光重合コンポジットレジンの光透過性の改善
平林 茂，平澤 忠
鶴見大学歯学部歯科理工学教室

物性の低下を招く事無く，光重合型コンポジットレジンの光透過性を改善するために，低屈折率を有し，かつ発蒸基またはフッ素置換ビスフェノール基のような高耐光性骨を有する4種のジメタクリレートを合成し，それらを含む6種の試作コンポジットレジンの硬化深さ並びに物性を調べた。UDMA，Bis-MEPPまたはBis-GMAを含有した3種の対照コンポジットレジンと比較検討した。

合成したモノマーを含有した実験グループの硬化深さはUDMAを含有するコンポジットを除いた対照グループに比較して深かった。硬化深さはマトリックスモノマーとフィラーの屈折率の差が小さくなるに従い増加し