An Assessment of the Influence of Dental Porcelain Slurry Preparation on Flexural Strength of Different Feldspathic Porcelains

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Abstract: Chipping remains a big challenge during the clinical application of glass ceramics in dentistry. The fabrication procedure used affects the mechanical properties of dental feldspathic porcelain and is associated with technical failures. This study aimed to compare the effect of the use of manufacturers' liquids versus H2O on the flexural strength of glass ceramics. Specimens (n = 120, n = 15 per group) (25 × 4 × 1.2 mm) were obtained using four porcelain powders (Creation CC, IPS InLine, Noritake EX-3, and Vita VM 13). Four groups were produced using porcelain powder and modeling liquid, and four groups using distilled water. The specimens were fired, sintered, and polished. Flexural strength was measured using a universal testing machine. Statistical analyses were conducted using post hoc Tukey's, two-way ANOVA, and Weibull analysis. Flexural strength values (mean ± SD) of the ceramic-manufacturer’s liquid mixture ranged between 67.2 ± 10.2 and 85.8 ± 12.8 MPA (NR < VT < IV < CR), while flexural strength values of the ceramic–water mixture were between 72.2 ± 6.9 and 95.2 ± 12 MPA (CR < NR < VT < IV). While the choice of the ceramic type significantly affected flexural strength, the use of water vs. manufacturers’ liquid showed in almost all cases no significance. To achieve better flexural strength results, InLine should be used with distilled water mixtures, while all ceramic powders except for Noritake can be used with the manufacturer’s liquid mixtures.

Keywords: ceramic; feldspathic porcelain; flexural strength; lithium disilicate; liquid

1. Introduction

Dental feldspathic porcelains are silica-based ceramics that consist of an amorphous glassy matrix (potassium aluminosilicate) with dispersed leucite particles [1,2]. The use of feldspathic porcelain in dentistry has been associated with superior aesthetics, high wear resistance, and biocompatibility [3–5]. Various porcelain-based restorations (e.g., PFM and veneers) have demonstrated safe treatment outcomes, with long-term survival rates. However, porcelain fracture or chipping were among the most reported complications [5]. Fabrication of dental porcelain, especially feldspathic veneering, is technique-sensitive, in addition to the material’s brittle nature and low strength [5,6]; therefore, thorough and precise adherence to technical fabrication protocol is crucial to clinical performance [7–9]. Moreover, several manufacturing factors can influence the final porcelain product, such as ceramic powder particle size, distribution, and chemical composition [9,10]. During the fabrication of porcelain prostheses, a mix of porcelain powder and liquid is made to produce the working ceramic slurry, paste for layering, and veneering to the full counter of the restoration. Ceramic slurry paste preparation conditions, ratios, and characteristics can
affect the physicomechanical properties, and therefore clinical outcomes [7–10]. As dental ceramic manufacturers and technicians have different recommendations and techniques on handling ceramics, it is not necessarily proven if slurry paste preparation is clinically advantageous to the sintered porcelain quality. For instance, Zhang et al. reported that the porcelain powder/liquid ratio influenced the density and total porosity of ceramic specimens [7]. Furthermore, the effect of porcelain powder–liquid mixing technique on bi-flexural strength was investigated. The results showed that when porcelain powder was incrementally mixed into liquid, a higher bi-flexural strength was achieved [8]. For proper porcelain condensation, porcelain powder has its particles distributed in a characterized size to yield a maximum density of particles. Such variation in particle size is critical to slurry application and sintering in terms of volumetric changes, and structural properties [9,10]. In the laboratory phase of porcelain slurry preparation, fine porcelain powder is mixed with liquid to form porcelain slurry paste. Dental technicians use different techniques to mix porcelain powder and liquid to produce porcelain slurry paste. Some technicians mix the porcelain powder with its own manufacturer’s porcelain modeling liquid. On the other hand, distilled water has been used with porcelain powder to produce porcelain slurry [10–12]. There is limited knowledge on the effect of using different liquids to mix with porcelain powder. Therefore, the aim of this study was to investigate the effect of using different liquids—distilled water and commercial porcelain liquids—on the flexural strength of sintered porcelain blocks. The null hypothesis tested was that mixing ceramic powders, with either the corresponding manufacturer’s liquid or water, would not have a significant effect on the flexural strength of leucite ceramics.

2. Materials and Methods

2.1. Specimen Preparation

Four commercially available dental porcelain powders (Creation CC (Creation Willi Geller International GmbH, Meininge, Austria), IPS InLine (Ivoclar Vivadent, Schaan, Liechtenstein), Noritake EX-3 (Kuraray Noritake Dental Inc., Tokyo, Japan), and Vita VM 13 (VITA Zahnfabrik H. Rauter GmbH & Co. KG, Bad Sackingen, Germany) were tested in this study (Table 1).

### Table 1. Chemical compositions of different ceramics and related modeling liquids, according to manufacturers. Sources: [https://www.ivoclarvivadent.com](https://www.ivoclarvivadent.com) (accessed on: 31 July 2019); [https://www.creation-willigeller.com](https://www.creation-willigeller.com) (accessed on: 31 July 2019); [https://apollonia-indent.net](https://apollonia-indent.net) (accessed on: 31 July 2019); [https://www.kuraraynoritake.e](https://www.kuraraynoritake.e) (accessed on: 31 July 2019) and [https://www.vita-zahnfabrik.com](https://www.vita-zahnfabrik.com) (accessed on: 31 July 2019).

| Brand     | Manufacturer                                      | Ceramic Chemical Composition                                      | Ceramic Modeling Liquid Chemical Composition (Weight %) |
|-----------|---------------------------------------------------|------------------------------------------------------------------|--------------------------------------------------------|
| Creation CC (C) | Creation Willi Geller International GmbH, Meininge, Austria | Leucite-containing feldspathic porcelain SiO₂ (55–65%); Al₂O₃ (12–18%); K₂O (12–16%); Na₂O (3–5%); TiO₂ (<1%); ZrO₂ (<1%); CaO (1–3%); LiO (<1%); MgO (<1%); B₂O₃ (1–2%); BaO (1–2%); SnO₂ (<1%); P₂O₅ (<1%); CeO₂ CeF₃ oxides (<0.1%); Pigments (0.1–0.3%) | Purified water (>99%); Propylene glycol (0–4%), Zinc chloride (<0.1%) |
| IV IPS InLine (IV) | Ivoclar Vivadent, Schaan, Liechtenstein | Leucite-containing feldspathic porcelain SiO₂ (59.5–65.5%); Al₂O₃ (13.0–18.0%); K₂O (10.0–14.0%); Na₂O (4.0–8.0%); other oxides (<4.0%); pigment (<2.0%) | Water and/or glycol 90–99%, polymer (1–10%) |
Table 1. Cont.

| Brand                      | Manufacturer                        | Ceramic Chemical Composition                                      | Ceramic Modeling Liquid                      |
|----------------------------|-------------------------------------|------------------------------------------------------------------|----------------------------------------------|
| Noritake super porcelain EX | Kuraray Noritake Dental Inc., Tokyo, Japan | Leucite-containing feldspathic porcelain                           | Polyethylene glycol                           |
|                            |                                     | SiO<sub>2</sub> (64.5%); Al<sub>2</sub>O<sub>3</sub> (14.4%)       | Water                                         |
|                            |                                     | CaO (<1.0%); MgO (<1.0%); K<sub>2</sub>O (8.7%); Na<sub>2</sub>O (9.2%) | Additives                                     |
|                            |                                     | Li<sub>2</sub>O (<1.0%); B<sub>2</sub>O<sub>3</sub> (0%); pigments (<1%) | % is not disclosed by the manufacturer        |
|                            |                                     |                                                                  |                                              |
| VITA VM13 (V)              | VITA Zahnfabrik H. Rauter GmbH & Co. KG, Bad Sackingen, Germany | Leucite-containing feldspathic porcelain                           | Purified water (>99%); Inorganic components |
|                            |                                     | SiO<sub>2</sub> (55–72%); Al<sub>2</sub>O<sub>3</sub> (12–16%); K<sub>2</sub>O (8–10%); Na<sub>2</sub>O (4–6%); TiO<sub>2</sub> (<1%); CeO<sub>2</sub> (<1%); ZrO<sub>2</sub> (<1%); CaO (1–2%); B<sub>2</sub>O<sub>3</sub> (1–2%); BaO (1–3%); SnO<sub>2</sub> (<1%); Mg, Fe, and P oxides (<0.1%) | (<1%) |

The porcelain powders used were composed of dentin–feldspathic porcelain containing leucite crystals. In preparation for firing, 0.7 g of each porcelain powder was mixed once with distilled water (15 specimens/porcelain type), and once with their corresponding porcelain’s modeling liquid (15 specimens/porcelain type). A homogeneous slurry was obtained according to manufacturer’s recommendations and inserted, in one increment, into a metallic mold. The mold was overfilled with slurry, and condensed by one dental technician (K.V.) on a vibrating table for 90 s. The slurry was then transferred into standardized cylindrical metallic pistons (25 mm length, 4 mm width, and 1.2 mm thickness), and absorbent paper was used to remove excess liquid. The disc-shaped specimens were thereafter removed from the matrix and placed in specific furnaces (Vacumat 40, VITA Zahnfabrik) for firing cycles, according to the manufacturer’s recommendations (Table 2).

Table 2. Firing procedures of the dental ceramics tested. ST: starting temperature, DT: drying time, FT: final temperature, TRI: temperature rate of increase, HT: holding time.

| Ceramic | ST (°C) | DT (min) | TRI (°C/min) | Vacuum Pressure | FT (°C) | HT (min) |
|---------|---------|----------|--------------|-----------------|---------|----------|
| CR      | 580     | 6        | 55           | Yes             | 920     | 1        |
| IV      | 403     | 4        | 60           | Yes             | 910     | 1        |
| NR      | 400     | 8        | 65           | Yes             | 980     | 1        |
| VT      | 400 °C  | 6        | 55           | Yes             | 880 °C  | 1        |

After the sintering process, the surface of each specimen was manually polished sequentially, using up to #600-grit silicon–carbide paper (Struers, Willich, Germany) under water-cooling, until a flat surface was obtained. Dimensions were verified using a digital micrometer (Mitutoyo, Kanagawa, Japan). The three dimensions (length, width, and height) of each specimen were measured. Specimens were inspected for cracks and replaced if needed. The final dimensions of the discs were 25 mm in length, 4 mm in width, and 1.2 mm in thickness (ISO 6872) [1].

2.2. Flexural Testing

Upon production of fifteen ceramic discs per group (n = 120, 8 groups), flexural strength measurements were conducted according to ISO 6872:2015 [1]. A three-point bending test was performed using a universal testing machine (Universal Testing Machine, ZwickRoell, Ulm, Germany) at a crosshead speed of 1 mm/min on a 10 mm span. Each specimen was loaded until ceramic failure occurred, and fracture load data were obtained.
The recorded values of the fracture load were used to calculate the biaxial flexural strength. The expected flexural strength (\( \sigma \) in MPa) of a rectangular sample under a load in a three-point bending setup is calculated using the following formula [13,14]:

\[
\sigma = \frac{3 \cdot F_{\text{max}} \cdot L}{2 \cdot b \cdot d^2},
\]

where \( F_{\text{max}} \) is the maximum force (Newton) applied to achieve fracture of the specimen.

2.3. Statistical Analysis

Statistical analyses were performed using the Social Sciences statistical software package (SPSS Software V.20, Chicago, IL, USA). Data on the flexural strength parameters of 4 different ceramic systems (Noritake, IV-Line, Creation and Vita) with two different liquids (manufacturer’s liquid and H\(_2\)O) were analyzed using two-way ANOVA comparisons. Interaction terms were analyzed using Tukey’s tests. In addition, maximum likelihood estimation without a correction factor was used for a 2-parameter Weibull distribution. Weibull distribution was performed to interpret the predictability and reliability of adhesion (Minitab Software V.16, State College, PA, USA). A value of \( p < 0.05 \) was considered statistically significant in all tests. No ethical approval or informed consent was needed for this study.

3. Results

The flexural strength values (Mean ± SD) of the ceramic-manufacturers’ liquid mixtures ranged between 67.2 ± 10.2 and 85.8 ± 12.8 MPa (NR < VT < IV < CR), while the flexural strength values (mean ± SD) of the ceramic–water mixtures were between 72.2 ± 6.9 and 95.2 ± 12.7 MPa (CR < NR < VT < IV) (Table 3).

### Table 3. The mean flexural values (MPa ± standard deviations) of IV, NR, VT, and CR, and their related modeling liquids. 95% confidence interval of mean values of IV, NR, VT, and CR, and their related modeling liquids. The same superscript lowercase letters in the same column indicate no significant differences and uppercase letters based on the experimental group (\( p < 0.05 \)).

| Ceramic Type | Flexural Strength (Mean ± SD) Water (W) | Flexural Strength (Mean ± SD) Liquid (L) | Min–Max FS (95% CI) Water (W) | Min–Max FS (95% CI) Liquid (L) |
|--------------|----------------------------------------|----------------------------------------|------------------------------|-------------------------------|
| CR           | 72.2 ± 6.9 \( ^{a,A} \)                | 85.8 ± 12.8 \( ^{a,A} \)               | 67.3–107.9 (68.5–75.9)       | 63.1–86.7 (79.0–92.6)       |
| IV           | 95.2 ± 12.7 \( ^{b,c,B} \)             | 83.1 ± 16.9 \( ^{a,c,B} \)             | 68.5–121.1 (88.4–102.1)      | 45.6–132.8 (74.1–92.1)      |
| NR           | 76.3 ± 6.3 \( ^{a,d,C} \)              | 67.2 ± 10.2 \( ^{b,d,e,C} \)           | 45.9–89.0 (72.9–79.7)        | 38.5–76.7 (61.8–72.6)       |
| VT           | 81.7 ± 12.1 \( ^{a,c,D} \)             | 79.7 ± 14.6 \( ^{a,c,e,D} \)           | 56.7–108.7 (75.3–88.1)       | 60.2–98.9 (71.9–87.5)       |

When the H\(_2\)O mixture of each ceramic system was compared to its corresponding manufacturer’s liquid mixture, there were no significant differences for all four ceramic types.

Within the H\(_2\)O mixtures, IV showed significantly higher flexural strength values compared to the other ceramics; namely, Creation (\( p = 0.00007 \)), Noritake (\( p = 0.006 \)), and Vita (\( p = 0.018 \)). Within the liquid mixtures, NR had the significantly lowest flexural strength values compared to the other ceramics, namely Creation (\( p = 0.003 \)) and InLine (\( p = 0.0431 \)), and no difference with Vita (\( p = 0.122 \)) (Figures 1–3).

When flexural strength was evaluated, Weibull distribution presented lower shape values for the groups IV (8.3 vs. 4.3), VT (9.0 vs. 6.0), and CR (11.0 vs. 7.8) when mixed with H\(_2\)O compared to their corresponding liquids, and higher shape values for the group NR (10.0 vs. 10.6). Weibull analysis showed more reliable flexural strength for the groups IV, VT, and CR when mixed with H\(_2\)O compared to their corresponding liquids (Figure 4).
When flexural strength was evaluated, Weibull distribution presented lower shape values for the groups IV (8.3 vs. 4.3), VT (9.0 vs. 6.0,) and CR (11.0 vs. 7.8) when mixed with H2O compared to their corresponding liquids, and higher shape values for the group NR (10.0 vs.10.6). Weibull analysis showed more reliable flexural strength for the groups IV, VT, and CR when mixed with H2O compared to their corresponding liquids (Figure 4).

**Figure 1.** Boxplot-best of the groups CR, IV, NR, and VT, mixed with H2O or their corresponding modeling liquids.

**Figure 2.** Mean of MPA interaction of the various tested ceramic groups CR, IV, NR, and VT, mixed with H2O or their corresponding modeling liquids.

**Figure 3.** Mean and standard deviation for the various tested ceramics groups CR, IV, NR, and VT, mixed with H2O or their corresponding modeling liquids.
It was found that both the liquid mixing ratio and porcelain type had a significant effect on resulting porosity and apparent density, and therefore influenced the mechanical characteristics of tested porcelains [14]. In that respect, differences in densities between the manufacturers’ modeling liquids (0.9–0.94 g/cm³) and distilled water (1.0 g/cm³) lead to variations in mixed slurry density, which consequently leads to different porcelain outcomes. Different studies showed increased density and decreased porosity when modeling liquid was used for porcelain slurry preparation. Zhang et al. [15] investigated the effect of different power/liquid ratios on porosity and translucency [7]. It was found that both the liquid mixing ratio and porcelain type had a significant effect on porosity and apparent density, and therefore influenced the mechanical performance during the procedure of ceramic restoration preparation is unknown. Our findings demonstrated comparable outcomes in the flexural strength of each ceramic, made either with distilled water slurry or using the manufacturers’ modeling liquid. However, our results showed that different ceramics produced with distilled water had significantly higher flexural strength in the IV–water group. Conversely, among ceramics prepared using manufacturers’ modelling liquids, the NRL group had significantly lower flexural strength compared to others. Similarly, Sinmazısık et al. [11] reported a non-significant statistical difference between bi-flexural strength of porcelains prepared with either distilled water or manufacturers’ modeling liquid. Within this non-significant difference, a 5% increase in bi-flexural strength was found when porcelains were prepared with modeling liquids. Our study, however, showed variations, with both increases and decreases in flexural strength achieved by both mixing liquids. These differences in findings, although not statistically significant, could be attributed to multiple factors, such as the difference in brands used in both studies, despite porcelain type similarity [11]. Moreover, Sinmazısik et al. reported results similar to our study’s, where significant differences were found between bi-flexural strength of tested ceramics, irrespective of the mixing liquids [11]. Nonetheless, all flexural strength values measured in this study are well above the ISO 6872 standard, with a minimum requirement of 50 MPa for ceramics used in metal substructures [13].

Optimizing the technical aspects of porcelain-based restoration aligns with maximizing its clinical performance and success. Clinically, chipping and fracturing of ceramic veneers are among the most frequent technical complications reported [2,3]. Of relevance, technical aspects of ceramic fabrication could influence its mechanical properties. Zhang et al. examined the effect of different power/liquid ratios on porosity and translucency [7]. It was found that both the liquid mixing ratio and porcelain type had a significant effect on resulting porosity and apparent density, and therefore influenced the mechanical characteristics of tested porcelains [14]. In that respect, differences in densities between the manufacturers’ modeling liquids (0.9–0.94 g/cm³) and distilled water (1.0 g/cm³) lead to variations in mixed slurry density, which consequently leads to different porcelain outcomes. Different studies showed increased density and decreased porosity when modeling liquid was used for porcelain slurry preparation. Zhang et al. [15] investigated...
the effects of various water powder/liquid and powder/water ratios and attributed this to the fact that a larger volume of modeling liquid was needed compared to distilled water, to achieve similar weight in the tested groups, which resulted in an increase in the density [14]. Thus, it increased the wetting of porcelain powder with modeling liquid and, therefore, the density. It was concluded that the observed increase in flexural strengths could be a result of reduced porosity and improved density of porcelain specimens [11,15]. Flexural strength values might be influenced by material composition and microstructure, as all tested materials yielded comparable SiO$_4$ values ranging between 55% and 72%, and the microstructure of glass ceramics, e.g., crystal size, crystal volume, and homogeneous distribution, influences the mechanical properties and crack propagation matter of the material [7]. However, previous investigations using XDR and SEM showed that neither distilled water or manufacturer’s recommended modelling liquid had any effect in the resultant final microstructure [11]. This finding is also in accordance with the results of this study, as flexural strength was comparable. A limitation of this in vitro study is that ageing procedures simulating the human oral environment were not used. Therefore, future research should investigate the effect of ageing upon the longevity of ceramic mixtures containing both different veneering liquids, and H$_2$O.

5. Conclusions
   From this study, the following could be concluded:
   - The flexural strength of the ceramic Creation was better when mixed with the corresponding producer’s liquid while all other tested ceramics (Noritake, InLine, and Vita) achieved higher flexural results when mixed with H$_2$O.
   - When the commercial modeling liquids were used, NRL resulted in a significantly lower flexural strength.
   - When different ceramic powders were mixed with distilled water, IV showed significantly highest flexural strength.
   - Overall, regardless of which mixing liquid was used, within each ceramic type, there was no statistically significant difference in flexural strength.

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