Research Article

An Investigation of Thermal Shock in Porous Clay Ceramics

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The thermal shock resistance of porous ceramic materials is often characterized by the Hasselman parameters. However, in other scenarios, the room-temperature residual strengths after thermal shock are also used to quantify the damage due to thermal shock. This paper attempts to link the measured residual strengths to the dominant crack features that are introduced due to thermal shock in porous clay ceramics produced by the sintering of clay powders with well-controlled size ranges. Residual strength estimates from bend tests are compared with fracture mechanics predictions. The implications of the residual strength results are then discussed for the characterization of damage due to thermal shock.

1. Introduction

The classical theory on the thermal shock behaviour of brittle ceramics was established by Hasselman in 1969 [1, 2]. He showed that opposing property requirements prevail, depending on whether the material is required to be resistant to crack initiation (for which high strength and low stiffness are essential), or resistant to strength degradation, following a severe thermal shock (in which case, low strength and high stiffness are beneficial). Furthermore, the room-temperature residual strengths after thermal shock are also used to quantify the damage due to thermal shock [2–5]. Since Evans and Charles [6] applied fracture mechanics to the modeling of thermal shock in 1975, resistance-curve (R-curve behavior) has been used extensively to model stable crack growth behavior in the ceramic community [7–11]. Lu and Fleck [12] have also developed a rigorous theoretical framework for the modeling of thermal shock while Soboyejo et al. [13] and Baker et al. [14] have explained crack growth resistance under thermal shock loading with crack-tip shielding concepts.

Porous clay ceramics form an important class of refractory materials that are used widely as thermal insulators in high-temperature applications such as in kilns, furnaces, and domestic charcoal stoves (jikos). In such applications, these materials are often exposed to cyclic thermal stresses and, at times, even severe thermal shocks that can result in significant strength degradation [15, 16]. There is, therefore, a need to relate the thermal shock behavior to the room-temperature strength degradation that is often observed in clays after thermal shock. This paper presents the results of a combined experimental and theoretical study that presents some new insights into how to relate damage due to thermal shock to the degradation of room-temperature strength of clay ceramics with well-controlled porosities. The implications of the results are discussed for the design of porous clay ceramics for refractory applications.
2. Experimental Procedures

2.1. Materials Processing. The clay raw materials used in this study were obtained from Mukono District in Uganda. The chemical composition of the clays is presented in Table 1. Four (4) types of test samples (with particle sizes ranges of 45–63 μm, 63–90 μm, 90–125 μm, and 125–255 μm) were prepared by ball milling and sieving. The sieved powders were then compacted into rectangular test specimens with dimensions of 6.5 ± 0.1 mm × 6.5 ± 0.1 mm × 52 ± 0.1 mm under 150 MPa pressure using a stainless die. The test samples were then fired to 1,300 °C at a heating rate of 5 °C/min, at a rate of 5 °C/min, and volume fraction porosity of the sintered samples were determined under three-point bend test using an Instron Model 8872 servohydraulic testing machine (Instron, Canton, MA) with cross-head speed of 0.5 mm min⁻¹ and a loading span (S) of 35 ± 0.1 mm. MOR was evaluated from the following expression [17]:

\[ \text{MOR} = \frac{3PS}{2B^2W^4} \]  

where \( P \) is the load at failure, \( S \) is the span; \( B \) and \( W \) are the respective breadth and width of the rectangular specimens. Fracture toughness (\( K_{\text{IC}} \)) tests were performed on single-edge notched-bend (SENB) specimens that were loaded monotonically to failure at a loading rate of 0.5 mm min⁻¹, with a loading span of 35 ± 0.1 mm. An initial notch, with a notch-to-width ratio (\( a/W \)) of ∼0.25, was introduced at the center of the specimens. Fracture toughness, \( K_{\text{IC}} \), was determined from (2) [18, 19]

\[ K_{\text{IC}} = \frac{P_0S}{BW^{3/2}} \times f \left( \frac{a}{W} \right), \]  

where \( P_0 \) is the failure load, \( a \) is the crack length (depth of the notch), and \( f(a/W) \) is the compliance function.

2.2. Porosity and Strength Measurements. The bulk density and volume fraction porosity of the sintered samples were measured using Archimedes’ immersion technique, which involved boiling the samples in water for 3 hours. The flexural strengths (MORs) of the rectangular test specimens were determined under three-point bend test using an Instron Model 8872 servohydraulic testing machine (Instron, Canton, MA) with cross-head speed of 0.5 mm min⁻¹ and a loading span (S) of 35 ± 0.1 mm. MOR was evaluated from the following expression [17]:

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2.3. Thermal Shock. The thermal shock resistance was determined by measuring the retention of the flexural strength of water-quenched (cold-shocked) samples. The samples were first heated in a furnace to 1,000 °C at the rate of 10 °C min⁻¹, A holding time of 30 min was then applied to the samples before quenching (cold-shocking) them into a container of water that was maintained at 5.0 °C. The samples were then dried in an oven for 30 mins at 110 °C. The dried samples were either tested in bending to determine their residual strengths or quenched further to establish the effects of quenching on their residual strengths. This was continued for 5 cycles.

The crack/microstructure interactions associated with cold-shock were observed in a scanning electron microscope (Philips XL30 FEG-SEM) equipped with energy dispersive X-ray spectroscopy (EDS) capabilities. The latter was used to characterize the local chemical compositions of the microstructural constituents. Some samples were also etched in concentrated hydrofluoric acid (HF 48%) for 3 minutes to reveal the underlying microstructures. These were rinsed thoroughly in water and ultrasonically cleaned in acetone before scanning electron microscopy (SEM) analysis. To avoid charging effects from the nonconducting ceramic surface, the samples were coated with a thin layer (~0.045 kA) of gold.

3. Results

3.1. Microstructure. The porous structure and the microstructure of the sintered clay samples are presented in Figures 1(a)–1(d). The samples produced by the sintering of small particles with diameters of ∼45–63 μm show nearly spherical pores (Figure 1(a)) with an overall porosity of ~8.8% (Table 2). However, the samples produced from the larger particle sizes (125–255 μm) show a network of interconnected pores (Figure 1(b)) with a higher level of porosity (18.3%), as shown in Table 2. Sample porosity also increased with increasing particle size range (Table 2).

Etched sections of the samples (Figures 1(c) and 1(d)) show large and well-defined mullite needles in a matrix of silica. The mullite needles have lengths of ∼1.5 ± 0.3 μm in samples processed from finer particle sizes (45–63 μm) and lengths of ∼5.5 ± 1.5 μm in samples processed from coarser particle sizes (125–255 μm) as indicated in Table 3.

3.2. Porosity and Room-Temperature Mechanical Properties. The room-temperature mechanical properties of the clay samples are summarized in Table 2. Figure 2(a) shows the dependence of the modulus of rupture (MOR) on porosity. The MOR decreases exponentially with increasing porosity. Similarly, the fracture toughness also decreases exponentially with increasing porosity (Figure 2(b)). Similar trends have been observed by other researchers [20, 21], and these trends of behaviour are generally attributed to stress concentration phenomena [22, 23].
Figure 1: Structure of sintered clay samples: (a) pore structures after sintering of powders with size ranges between 45–63 μm, \( P = 8.8\% \); (b) size ranges between 125–255 μm showing bimodal and interconnected pore structure, \( P = 18.3\% \); (c) etched sections of samples from particle size ranges 45–63 μm showing small mullite needles (1-2 μm) and (d) etched section of samples from particle size ranges 125–255 μm showing large mullite crystals (5–7 μm).

Table 2: Mechanical properties of clay ceramics.

<table>
<thead>
<tr>
<th>Particle size range (μm)</th>
<th>Porosity (%)</th>
<th>Bulk density, ( \rho ) (g cm(^{-3}))</th>
<th>Fracture toughness, ( K_{\text{IC}} ) (MPa√m)</th>
<th>MOR (MPa)</th>
<th>( E ) (GPa) ±5%</th>
</tr>
</thead>
<tbody>
<tr>
<td>45–63</td>
<td>8.8 ± 0.6</td>
<td>2.5 ± 0.1</td>
<td>2.1 ± 0.1</td>
<td>19.4 ± 4.0</td>
<td>54.06</td>
</tr>
<tr>
<td>63–90</td>
<td>9.9 ± 0.4</td>
<td>2.4 ± 0.1</td>
<td>1.2 ± 0.3</td>
<td>11.1 ± 3.2</td>
<td>53.15</td>
</tr>
<tr>
<td>90–125</td>
<td>14.0 ± 1.5</td>
<td>2.3 ± 0.1</td>
<td>0.8 ± 0.2</td>
<td>6.9 ± 1.0</td>
<td>49.99</td>
</tr>
<tr>
<td>125–250</td>
<td>18.3 ± 2.6</td>
<td>1.9 ± 0.1</td>
<td>0.6 ± 0.1</td>
<td>5.8 ± 1.5</td>
<td>46.95</td>
</tr>
</tbody>
</table>

Table 3: Microstructural details of the clay ceramics.

<table>
<thead>
<tr>
<th>Particle size range (μm)</th>
<th>Length (l) of mullite needles (μm)</th>
<th>Mullite width (w) (μm)</th>
<th>Aspect ratio (l/w)</th>
</tr>
</thead>
<tbody>
<tr>
<td>45–63</td>
<td>1.50 ± 0.30</td>
<td>0.16 ± 0.08</td>
<td>9.03 ± 0.38</td>
</tr>
<tr>
<td>63–90</td>
<td>2.60 ± 0.40</td>
<td>0.25 ± 0.05</td>
<td>10.4 ± 0.45</td>
</tr>
<tr>
<td>90–125</td>
<td>3.25 ± 0.25</td>
<td>0.35 ± 0.15</td>
<td>9.28 ± 0.40</td>
</tr>
<tr>
<td>125–250</td>
<td>5.50 ± 1.50</td>
<td>0.60 ± 0.20</td>
<td>9.17 ± 1.70</td>
</tr>
</tbody>
</table>
Table 4: Measured residual strengths of clay ceramics after thermal shock.

<table>
<thead>
<tr>
<th>Particle size range (μm)</th>
<th>Residual strength, $\sigma_{\text{meas}}$ (MPa)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>$N = 1$</td>
</tr>
<tr>
<td>45–63</td>
<td>12.21</td>
</tr>
<tr>
<td>63–90</td>
<td>8.39</td>
</tr>
<tr>
<td>90–125</td>
<td>6.64</td>
</tr>
<tr>
<td>125–250</td>
<td>5.15</td>
</tr>
</tbody>
</table>

Figure 2: The effects of porosity on room-temperature mechanical properties: (a) Flexural strength (MOR) and (b) Fracture toughness $K_I$.

3.3. Residual Strengths and Cold Shock. The dependence of residual flexural strengths on number of shock cycles is presented in Table 4 and shown graphically in Figure 3. The largest changes in strength in all samples were observed after the first cycle. Thereafter, smaller incremental changes in the residual strength were observed as the number of shock cycles increased. The highest initial strengths and the highest subsequent drops in strength were observed in the samples processed from particles with diameters of 45–63 μm. These failed consistently after 3 shock cycles. Furthermore, the residual strengths decreased with increasing particle size ranges, though the fractional changes in strength were lower as the number of shock cycles increased. The samples produced by the sintering of the larger particle sizes generally survived for up to 5 cold-shock cycles.

3.4. Crack/Microstructure Interactions. Typical crack/microstructure interactions observed in the thermally shocked samples are presented in Figures 4(a)–4(c). These show the growth of dominant cracks, with uncracked bridging ligaments behind the crack tip. Viscoelastic crack bridging [24] acts to restrict the opening of the cracks, and thereby promote the shielding of cracks. After 1 cold quench cycle, clay samples with finer particle sizes (45–63 μm) were observed to have more viscous bridging ligaments than those produced from larger particles sizes (125–255 μm). This is shown clearly in Figures 4(a) and 4(b); however, the incidence of ligament bridging decreases with increasing number of shock cycles, as shown in Figure 4(c).

After cold-shock, the other toughening mechanisms observed in the room-temperature flexural specimens are presented in Figures 5(a)–5(c). These show evidence of toughening by crack-tip blunting (cracks being arrested by pores, Figure 5(a)), crack deflection and crack branching (Figure 5(b)), and secondary microcracking (Figure 5(c)).
4. Discussions

4.1. Fracture Modes and Residual Strength Predictions. Typical fracture modes observed in the flexural specimens after cold-shocking and room-temperature fracture (under three-point loading) are presented in Figures 6(a)–6(d). These show clearly that final fracture initiated largely from corner cracks that were introduced into the specimens during thermal shock. Such cracks are found in a large number of engineering structures and components [7], where stress

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Figure 4: Schematic of observed toughening mechanisms due to viscous bridging in clay samples: (a) 45–63 μm, N = 1, showing viscous bridging ligaments at the crack tip; (b) 125–255 μm, N = 1, showing less ligament bridging at the crack tip compared to (a); (c) 45–63 μm, N = 2, showing shattered bridging ligaments due to thermal shock.

Figure 5: Schematic of observed toughening mechanisms in clay samples due to (a) crack blunting by a pore, (b) crack deflection and crack branching, and (c) fractured surface showing intergranular fracture and secondary crackings.
concentrations at corners can promote their formation and growth. In the current work, most of the cracks had profiles that could be approximated as quarter circular cracks (Figures 6(a)–6(d)). Hence, the stress intensity factor, $K$, along the crack front can be estimated from [25]:

$$K = \frac{2}{\pi} \sigma \sqrt{\pi a}, \quad (3a)$$

where $\sigma$ is the remote applied stress and $a$ is the crack length. It is important to note here that this is an approximate solution for a quarter-circular crack in a half space. Hence, for semielliptical cracks or larger cracks that interact significantly with the specimen geometries, more complex solutions are needed, such as those published by Newman and Raju [24, 26] and Scott and Thorpe [27]. Finite width correction factors by Holdbrook and Dover [28] may also be applied to larger cracks. In any case, Equation (3a) may be rearranged to obtain estimates of the residual strength. This gives the following:

$$\sigma_f = \frac{K_c}{2\sqrt{\pi a/\pi}}, \quad (3b)$$

where $\sigma_f$ is the fracture strength, $a$ is the crack length (due to thermal shock), and $K_c$ is the stress intensity factor. The residual strengths of the thermally shocked samples were predicted using (3b). The values of the crack length, $a$, were obtained from SEM images of the fractured surfaces (Figure 6). It is important to note here that the crack lengths observed after the first quench cycle ($N = 1$) were larger ($a = 1.3$ mm) in samples processed from finer particle sizes (63–90 $\mu$m) than the crack lengths ($a = 1.0$ mm) observed in the samples processed from coarser particle sizes (90–125 $\mu$m) for $N = 1$. Further, the extent of shock-induced crack growth was greater in samples produced from finer particle sizes compared to the crack growth in samples produced from coarser particle sizes. This is shown in Figure 7 in which the measured crack lengths are plotted against the number of shock cycles for the porous clays that were examined in this study.

Figure 8 shows a plot of the predicted residual strengths ($\sigma_f$) against the measured residual strengths obtained using (1). It can be seen that the predictions from (3b) are clearly in very good agreement with the measured residual strengths.
4.2. Thermal Shock Resistance Parameters. Hasselman $R'''$ parameter [1, 29] characterizes the ability of a material to resist crack propagation under loading and further damage and loss of strength on thermal shocking. It is, therefore, of interest to explore the dependence of the $R'''$ parameter on pore size and the number of shock cycles. The $R'''$ parameter is related to the thermomechanical and elastic properties ($E, \nu$) via [29]:

$$R''' = \frac{E}{\sigma_f^2(1 - \nu)},$$  \hspace{1cm} (4)

where $\sigma_f$ is the flexural strength (MOR), $E$ is Young's modulus, and $\nu$ is Poisson’s ratio.

In the current work, Young’s modulus, $E(P)$ at porosity $P$, was estimated from the analytical expression derived by Boccaccini et al. [22]. This accounts for the effects of pore shape and pore orientation on Young’s modulus. It is given by [22]:

$$E(P) = E_o \left(1 - P^{2/3}\right)^s,$$  \hspace{1cm} (5)

where $E_o$ is Young’s modulus for zero porosity and $s$ is given by

$$s = 1.21 \alpha^{1/3} \sqrt[3]{\left[1 + \left(\frac{1}{\alpha^2} - 1\right) \cos^2 \phi\right]}.$$  \hspace{1cm} (6)

In the above expression, $\alpha$ is the mean aspect ratio (shape factor) of the pores and $\cos^2 \phi$ represents the orientation factor of the pores, with $\phi$ being the angle between the rotational axis of the pores and the stress direction. The case of random statistical orientation is obtained by setting $\cos^2 \phi = 0.33$.

From the SEM micrographs (Figures 1(a) and 1(b)), it is clear that the pores in the clay samples were mostly randomly oriented. Hence, it is justified to use a value of $\cos^2 \phi = 0.33$ and $\alpha = 0.4$ [22, 30] in calculating the value of $s$. From the literature [30], the value of $E_o$ is 65.0 GPa while the Poisson’s ratio $\nu$ is 0.23. These were used in calculating $E(P)$. In this case, it was assumed that porosity and pore shape have a negligible effect on Poisson’s ratio such that $\nu(P) = \nu_o = 0.23$ [31].

The thermal shock parameter calculated from (4)–(6) are plotted in Figure 9 as a function of the number of quench cycles, $N$. The porous clay samples clearly show an increase in the $R'''$ parameter with increasing number of cold-shock cycles. Furthermore, samples produced from coarser particle sizes (125–255 μm) exhibit higher crack growth resistance than those obtained from finer particle size distributions (45–63 μm). This observation is in agreement with Figure 7.
in which it was observed that the shock-induced crack growth was greater in samples produced from finer particle sizes compared to the crack growth in samples with coarser particle sizes. Since the $R'''$ parameter characterizes the ability of a material to resist crack propagation, the current results suggest that the coarser particle sizes should result in greater thermal shock resistance. However, it is also important to note that larger particle sizes and increased porosity give rise to lower residual strength levels (Figure 3), although the number of shock cycles to failure generally increases with increasing porosity arising from the sintering of larger particle sizes.

4.3. Implications. The implications of the current work are quite significant. First, the results show clearly that the thermal shock resistance, $R'''$ and the room-temperature residual strengths change with increasing pore volume fraction, which was controlled in the current study by the sintering of different particle size ranges. However, the trends in the $R'''$ parameter (Figure 9) are opposite to those in the room-temperature residual strengths obtained after cold thermal shock (Figure 3). This is due presumably to the different damage and shielding mechanisms that operate in the different temperature and loading regimes (Figures 4 and 5). Hence, comparison of thermal shock resistance should consider the underlying damage and crack-tip shielding mechanisms.

Furthermore, the good agreement between the residual strength measurements and the residual strength predictions (Figure 8) suggest that simple fracture mechanics and fractographic approaches may be used to estimate the residual strength levels. Furthermore, nondestructive surface inspections for corner cracks may also provide useful approaches to the life assessment and management of kilns that are used to process molten metal and ceramics in the trillion-dollar materials processing industry. Since the stoppage of production (required for kiln repair) results in millions of dollars of losses in lost revenue, the application of the methods proposed in this study could provide the basis for the improved management and design of kilns. The methods can also be applied to the modeling of thermal shock in a wide range of thermostructural applications in which damage can occur due to thermal shock.

5. Conclusions

This paper presents the results of combined experimental and theoretical study of damage due to thermal shock in porous clays. Salient conclusions arising from the study are summarized below.

1. The Hasselman thermal shock resistance parameter, $R'''$, and the room-temperature residual strengths after thermal shock are strongly dependent on the volume fraction of porosity. The parameter $R'''$ increases with increasing porosity (volume fraction) and shock cycles while the room-temperature residual strengths decrease with increasing porosity fraction and shock cycles.

2. The room-temperature residual strengths can be predicted from a simple fracture mechanics model that idealizes the corner cracks formed due to thermal shock as quarter-circular cracks in a half space. However, actual crack geometries should be considered when the cracks are not approximately semicircular. The models should also account for finite width effects when the crack sizes are large compared to the structural sizes.

3. Viscoelastic crack bridging is the dominant crack-tip shielding mechanism under thermal shock loading of porous clays. The extent of viscoelastic bridging decreases with increased porosity and increasing shock cycles. Furthermore, the room-temperature residual strengths are dominated by the shielding due to ligament bridging. Lower shielding levels are associated with crack deflection and crack branching during room-temperature fracture and shock-induced crack growth.

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References


