Crack Branching and Fracture Mirror Data of Glasses and Advanced Ceramics

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SUMMARY

The fracture mirror and crack branching constants were determined from three glasses and nine advanced ceramics tested under various loading and specimen configurations in an attempt to use the constants as a data base for fractography. The ratios of fracture mirror or crack branching constant to fracture toughness were found to be approximately two for most ceramic materials tested. A demonstration of how to use the two constants as a tool for verifying stress measurements was presented for silicon nitride disk specimens subjected to high-temperature, constant stress-rate biaxial flexure testing.

I. BACKGROUND

Fractographic analysis of glasses and ceramics can play an essential role in not only locating and characterizing fracture origins but identifying failure mechanisms. The commonly observed features of brittle fracture are mirror, mist, hackle, and crack-branching patterns which are normally formed in fracture surfaces particularly for silicate glasses, as shown in figure 1 (ref. 1). As a crack propagates catastrophically from its origin, it leaves behind sequentially these distinctive surface features. These fracture features with their boundaries can give significant information as well as empirical practicality in failure analysis. Fracture mirrors have been utilized to pinpoint fracture origins, to predict the sizes of failure-initiating flaws, to estimate fracture stresses, and to determine residual stresses presented in specimens (refs. 1 to 7). In terms of analytical approach, however, an enormous complexity has been encountered in understanding and formulating the mechanics of these fracture features with respect to dynamic crack propagation.

The distances from the fracture origin to the boundaries (fig. 1) have been empirically related with the fracture strength as follows (refs. 1 and 8)

\[ \sigma_f r_i^{1/2} = A_i \]  

where \( \sigma_f \) is the fracture strength (typically in MPa), \( r_i \) is the radius of the \( i \)th boundary from the fracture origin (typically in meter) and \( A_i \) is the constant for the \( i \)th boundary. Note that the units of \( A, \text{ MPa}\sqrt{m} \), are the same as those of stress intensity factor. The parameter \( A_m \) is known as the fracture mirror constant for the boundary of the mirror and the mist regions, and the parameter \( A_b \) is the crack branching constant for the boundary of the hackle.

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and the branching regions. In some cases, particularly for polycrystalline ceramics, fracture mirror size \((r_m)\) and crack branching length \((r_b)\) are not clearly distinguishable so that little difference between the fracture mirror and the crack branching constants has been attained.

Many investigations have been made to determine the fracture mirror or crack branching constant for silicate glasses since these materials generally exhibit the four distinctive fracture features, allowing reasonably accurate measurements of mirror sizes or crack branching lengths. The fracture mirror constant or crack branching constant has appeared to range from \(A = 1.7\) to \(2.7\) MPa\(\sqrt{m}\) with an approximate mean of \(A = 2\) MPa\(\sqrt{m}\), regardless of flaw history such as modes of loading (flexure or tension, uniaxial or biaxial), residual contact stresses, and environment (refs. 1 and 4 to 11). Because of this somewhat consistent result, the constant has been combined with fracture toughness of the material, drawing a conclusion that the constant would be a unique material constant. However, the scatter of mirror constants available in the literature is not comparable with that of mechanical properties such as fracture toughness, hardness and Young’s modulus. The mirror constants have shown a greater variation depending on load/specimen configurations and even on investigators than fracture toughness, hardness or Young’s modulus. By contrast, for example, fracture toughness of soda-lime and fused silica glasses do not exhibit any appreciable variation in terms of test method which is either by indentation, double torsion, double cantilever beam, compact tension, single edge precracked beam or by any other test techniques, with a range of \(K_{IC} = 0.75\) to 0.79 MPa\(\sqrt{m}\), as pointed out by Ramulu (ref. 12). Particularly for polycrystalline ceramics in which the respective boundaries are not generally well defined, the variation in \(A\) has become more significant (refs. 1, 4, 10, and 13).

Because of the inconsistency in \(A\), some investigators have pointed out that fracture mirror or crack branching constant is not merely a material constant. In his dynamic fracture mechanics analysis using a dynamic finite element method, Ramulu (ref. 12) has shown that the formation of fracture mirror for soda-lime glass is attributed to the oscillations of dynamic stress intensity factor \((= K_{ID})\) as a crack propagates rapidly, suggesting that the constant be a product of mechanics response. Although several criteria such as velocity (ref. 14), strain energy density (refs. 8 and 9) and static stress intensity factor (refs. 15 to 18) have been proposed and each criterion is suitable to a certain specific condition, none of the criterion can explain satisfactorily all the detailed mechanics on the formation of fracture mirror and subsequent features (ref. 1). This is due to limited understanding of dynamic crack propagation in the nature of catastrophic failure associated with stress wave interaction, which has been considered as a ‘formidable’ problem.

Notwithstanding the difficulty in understanding the exact nature (mechanics) of fracture-mirror formation, the fracture mirror or crack branching constant has been used widely as a tool to predict the flaw size as well as the fracture stress associated with failure, as aforementioned. In some cases, critical flaws are too small in size to measure even microscopically and/or are not discernable enough to characterize their configurations. In this case, the fracture mirror sizes and mirror constant can give a useful information to estimate the flaw sizes at failure. Based on the fracture mechanics approach together with equation (1), a ratio of mirror size to critical flaw size can be obtained for an infinite body as follows:

\[
\frac{r_m}{c_f} = \left(\frac{K_m}{K_{IC}}\right)^2
\]  

(2)

where \(K_m\) is the ‘stress intensity factor’ at mirror formation, expressed as \(K_m = Y \sigma_f r_m^{1/2} = YA_m\) with \(Y\) being a crack geometry factor, and \(K_{IC} = Y \sigma_f c_f^{1/2}\) with \(c_f\) being a critical crack size. For example, a ratio of mirror size to flaw size was found to be about \(c_f/r_m = 10\) to 13 for silicate glasses (ref. 2). In other cases, a situation may occur where a ceramic component has failed in service without the working stress being known. If it is feasible to determine fracture mirror size or crack branching length through fractography, the approximate operating-stress that had caused the component to fail can be estimated based on equation (1) with the fracture mirror or crack branching constant available. Consequently, any pertinent changes in design or operating conditions would be made through this post-failure analysis. Since a failure-initiating flaw is generally surrounded symmetrically by mirror, mist and hackle regions (fig. 1), the location of the flaw also can be easily identified. In summary, the data on fracture mirror and/or crack-branching, sizes and constants, can be utilized very usefully in fractographic analysis of brittle materials. An ASTM standard (C 1322) (ref. 19) for fractographic analysis of advanced ceramics describes this technique in details, and many review articles on fractography allocate a large portion for the features of fracture-mirror or crack-branching sizes and constants, which reflect both the importance and usefulness of the technique.
A large quantity of glass and ceramic specimens have been tested under various testing programs at the Ceramics Lab of NASA Lewis Research Center. The measurements on fracture mirror sizes and/or crack branching lengths have been made for some of these glass and advanced ceramic specimens tested in constant stress-rate or strength testing at either room or elevated temperatures. This paper summarizes the measurements of fracture mirror sizes and/or crack branching lengths as a function of fracture stress for three glasses and nine advanced ceramics to determine the corresponding fracture mirror and/or crack branching constants. Since, in general, the fracture mirror or crack branching constants have shown to vary depending on materials, test methods and/or specimen configurations, a large amount of data covering a variety of test matrices is needed. The determined constants can then be used as a data base for a wide range of materials and/or specimen/loading configurations. Finally, an example of how fracture stresses were able to be remedied by the use of fractographic analysis based on equation (1) is presented for silicon nitride specimens subjected to uniaxial and biaxial flexure testing at elevated temperature.

II. EXPERIMENTS

Types of testing, and loading and specimen configurations for each test material where the measurements of fracture mirror size or crack branching length were attained, are summarized in Table I. The table also includes hardness, fracture toughness, fracture mirror or crack branching constant, ratio of fracture mirror or crack branching constant to fracture toughness of each material. Due to the variety of test matrices, detailed experimental procedures for each material were omitted here and should be referred to the corresponding reference. Brief descriptions regarding test conditions and specimens were made in the Result and Discussion section. The mirror sizes and crack branching lengths were determined using low power microscopes, typically ranging in magnification from 10 times to 100 times. In general, the measurements of fracture mirror sizes were made for the specimens tested in uniaxial tension or uniaxial flexure, while the measurements of crack branching lengths were made for the disk or plate specimens in biaxial flexure. Exception to this was soda-lime glass plates (microslides) where fracture mirror sizes and crack branching lengths were determined, respectively, from biaxial and uniaxial flexure.

III. RESULTS AND DISCUSSION

1. Fracture Mirror and Crack Branching

(a) NC132 silicon nitride in uniaxial and biaxial flexures

Figure 2 shows the plots of fracture stress (log $\sigma_m$) as a function of fracture mirror size (log $r_m$) (fig. 1(a)) and of crack branching length (log $r_b$) (fig. 1(b)). This figure is for NC 132 silicon nitride uniaxial (four-point beam) and biaxial (ring-on-ring disk) flexural specimens subjected to constant stress-rate (or called “dynamic fatigue”) testing at 1100 °C in air (ref. 20). The mirror and branching constants were determined to be $A_m = 9.40 \pm 1.19$ MPa$\sqrt{m}$ and $A_b = 5.66 \pm 1.49$ MPa$\sqrt{m}$, respectively, from the functional-fit analysis based on equation (1). Contrary to common observations, the crack branching constant was rather small (40 percent less), as compared with the mirror constant. This was due to the error in fracture-stress measurements in biaxial flexure, which will be discussed in the later section. A typical fracture surface of a uniaxial four-point bar specimen as well as a typical crack branching pattern of a biaxial disk specimen is shown in figure 3.

(b) GN10 silicon nitride in pure tension and biaxial flexure

Figure 4 depicts both the fracture stress plotted versus fracture mirror size for GN10 button-head, pure tension specimens (ref. 21) and the fracture stress as function of crack branching length for GN10 particle-impacted, biaxial disk specimens (ref. 22). The mirror constant in pure tension was $A_m = 11.78 \pm 1.41$ MPa$\sqrt{m}$, and the crack branching constant in biaxial flexure was $A_b = 10.32$ MPa$\sqrt{m}$. Despite different vintage, a reasonable agreement between the two constants exists, indicating that the constants for GN10 are almost independent of flaw history with either machined surface with monotonic tension loading or particle-impacted surface with biaxial flexure.
loading. The independence of fracture mirror or crack branching constant on flaw history such as flaw type, loading condition and environment has been observed for many glasses and advanced ceramics (refs. 1 and 4 to 11). It should be noted that the demarcation between mirror and hackle regions is not usually well-defined in polycrystalline ceramics so that the difference between $A_m$ and $A_b$ is insignificant (refs. 10 and 13).

(c) *Hexoloy silicon carbide in biaxial flexure*

The crack branching results for Hexoloy silicon carbide (α-SiC) plates tested in ring-on-ring biaxial flexure (ref. 23) are shown in figure 5. The branching constant was $A_b = 5.45 \pm 0.30$ MPa$\sqrt{\text{m}}$ for a total of 111 data points resulting in an excellent data fit to equation (1) with a coefficient of variation of 5 percent in $A_b$. A typical crack branching pattern and a corresponding fracture surface of a tested specimen are shown in figure 6. It is noted that the surface flaw yielded a through-the-thickness crack that extended in a plane to a length of $2r_b$ before branching into several segments, as observed in biaxial disk fracture by Shetty et al. (ref. 24).

(d) 96 wt% alumina in uniaxial and biaxial flexures

The results for the crack-branching length and fracture-mirror size measurements for 96 wt% alumina in four-point uniaxial, ring-on-ring biaxial and ball-on-ring biaxial flexures (ref. 25) are shown in figure 7. The constants were found to be $A_m = 7.64 \pm 0.53$ and $A_b = 7.24 \pm 0.66$ and $7.39 \pm 0.55$ MPa$\sqrt{\text{m}}$, respectively, for four-point uniaxial, ring-on-ring biaxial and ball-on-ring biaxial flexures. Little variation in $A$’s among the three different test conditions indicates again the independence of $A$ on flaw, specimen and/or loading configurations.

(e) Y-TZP, AS44 and SN220 silicon nitrides in biaxial flexure

The post-impact strength as a function of crack branching length for Y-TZP, AS44 and SN220 silicon nitride disk specimens are depicted in figure 8. Each disk specimen was subjected to a specified particle-impact test and then its subsequent strength was determined by using a ring-on-ring biaxial fixture at room temperature (ref. 22). The crack branching constants were $A_b = 11.48 \pm 1.46$, $10.85 \pm 2.71$, and $8.13 \pm 2.36$ MPa$\sqrt{\text{m}}$, respectively, for Y-TZP, AS44 and SN220. It is noted that the scatter in the data was significant at high $r_b$, particularly for AS44 and SN220 silicon nitrides. It is questionable whether the crack branching data with $r_b > 1$ mm can be included, in view of equation (1) which is based on the assumptions of an infinite body and a uniaxial stress-state. However, the values of crack branching constant excluding such data ($r_b > 1$ mm) would not be appreciably different from those for all data included.

(f) SiC whisker-reinforced composite silicon nitride and monolithic silicon nitride in uniaxial flexure

The previously published, fracture stress versus mirror size data on 30 vol% SiC whisker-reinforced composite silicon nitride and similar monolithic silicon nitride (both fabricated by Norton Co.) (ref. 13) are presented in figure 9. The strengths of both the composite and monolithic (flexure beam) specimens indented with indentation loads (with Vickers microhardness indenter) ranging from 1 to 10 N were determined in four-point flexure at room temperature. The fracture mirror constant was $A_m = 6.63 \pm 0.11$ and $5.88 \pm 0.14$ MPa$\sqrt{\text{m}}$, respectively, for the composite and monolithic silicon nitrides.

(g) Glasses in uniaxial and biaxial flexures and in pure tension

The results of fracture stress vs. mirror size for different glasses are shown in figure 10. The figure consists of the data on: (a) fused silica rods (as-received) in four-point uniaxial flexure (ref. 26); soda-lime glass square plates (annealed and etched) in ring-on-ring biaxial flexure (ref. 27); and indented fused silica optical glass fibers (ref. 7). The mirror constant was obtained to be $A_m = 2.20 \pm 0.33$, $1.81 \pm 0.28$, and $2.10$ MPa$\sqrt{\text{m}}$, respectively, for fused silica rods, soda-lime square plates, and optical glass fibers. The values of fracture mirror constant are in good agreement with the published data which range from $A_m = 1.7 - 2.7$ MPa$\sqrt{\text{m}}$ with a mean of $A_m = 2$ MPa$\sqrt{\text{m}}$ (refs. 1 and 4 to 1). This indicates again that the mirror constant for glass is almost independent of the flaw history.
such as loading (pure tension, or uniaxial or biaxial flexure), flaw types (natural, machining or indentation flaws) and environment (moisture or inert), as long recognized (refs. 1 and 4 to 11). A typical fracture surface of a fused silica glass rod showing a well-defined mirror formation is shown in figure 11.

The crack branching data on soda-lime glass rectangular plates subjected to constant stress-rate testing in four-point flexure were purposely collected and are presented in figure 12. The specimens were indented with low indentation loads (P = 2 and 5 N with Vickers microhardness indenter), and tested in as-indented or annealed condition (ref. 28). The crack branching constant was found to be $A_\alpha = 3.54 \pm 0.64$ MPa$\sqrt{m}$. This value of crack branching constant is considerably higher than those shown in figure 10. Also note a poor data fit to equation (1) with a coefficient of variation of 18 percent in $A_\alpha$. The figure shows that the lower $r_b$’s yield the lower branching constant, and vise versa. This clearly indicates that there exists a lower limit of branching length (in conjunction with specimen thickness) below which only a reasonable value of crack branching constant is obtained. Therefore, the use of equation (1) to determine the crack branching constant for branching lengths greater than 1 mm for thin (1.2 mm thick) microslides is not appropriate, primarily due to the inapplicability of the infinite-body assumption. Typical crack branching patterns and corresponding fracture surfaces for the two specimens failed at low and high fracture stresses are shown in figure 13. Similar to figure 6, the surface flaws extended in a plane to almost a length $2r_b$ to form through-the-thickness cracks before branching.

2. The Ratio of Fracture Mirror or Crack Branching Constant to Fracture Toughness, $A/K_{IC}$

A summary of the ratios of fracture mirror or crack branching constant to fracture toughness, $A/K_{IC}$, is presented in Table I. The plots of $A$ as a function of $K_{IC}$ are also shown in figure 14. Here, the soda-lime glass uniaxial flexure data (fig. 12) were not included since some of crack branching lengths were much greater than the specimen thickness. The NC132 fracture mirror constant was corrected with a correction factor of 1.38 (see Section 4). Glasses exhibit the ratios ranging from $A/K_{IC} = 2.4$ to 2.8. However, most other data except for glasses seem to follow approximately along the line of $A/K_{IC} = 2$, with an overall average value of $A/K_{IC} = 2.05 \pm 0.34$. This is somewhat inconsistent with the results obtained by Rice et al. (refs. 2 and 6), where the data follow reasonably well with a ratio of $A/K_{IC} = 3$. However, it should be noted that a relatively large range of the ratios from $A/K_{IC} = 1.1$ to 3 has been reported for polycrystalline ceramics, depending on specimens, loading configurations and even researchers (refs. 10, 13, and 19). This implies that the ratio is not solely a material-independent constant for advanced ceramics, as pointed out previously (refs. 10, 12, and 13), which in turn means fracture mirror or crack branching constant is not a uniquely-defined material constant. This is in contrast with other physical properties such as Young’s modulus and fracture toughness. It is believed that the discrepancy may be further amplified particularly for the fracture mirror constant, due to the arbitrary choice of mirror boundaries in polycrystalline ceramics since unlike glasses the boundaries are not completely evident (refs. 10 and 13).

3. Effect of Hardness on Crack Branching

Kirchner suggested that a critical crack-tip strain should be reached to initiate branching (ref. 29). The basic idea is that crack branching occurs when the total strain at the boundary of a yield zone at the crack tip is a constant that is related to the crack opening required for a branching crack in ceramics, and that hardness is a factor in determining the crack tip strain. With some ceramics data Kirchner showed a relationship between stress intensity factor and hardness of the material as follows:

$$K_B = MH + \sqrt{2K_{IC}}$$

(3)

where $K_B$ is the (static) stress intensity factor at branching, expressed as $K_B = Y\sigma_{r_B}^{1/2} = YA_B$, $H$ is hardness, and $M$ is a constant (or the slope in a plot of $K_B$ versus $H$). The crack geometry factor $Y$ does not change more than 10 percent in the range of $0 \leq r_b \leq 0.5t$ with $t$ being specimen thickness (ref. 30). Equation (3) also can be used by replacing $K_B$ with $A_B$ (or $A_m$) since the functional trend remains unchanged. The results of $A/K_{IC}$ versus $H/K_{IC}$ for glasses and ceramics are shown in figure 15. The soda-lime glass crack branching data shown in figure 12 was not included in the figure. The NC132 fracture mirror constant was corrected with a correction factor of 1.38 (see
Section 4). Unlike the data shown by Kirchner (ref. 29), a significant scatter is apparent in the data obtained from this study, as shown in the figure. Notwithstanding, overall trends suggested by Kirchner seem to be supported by much broader range of data, as also mentioned by Rice (ref. 1).

It has been shown that the commonly referred three criteria of fracture mirror formation or crack branching, crack velocity, strain energy density and stress intensity factor, are generally interrelated. Rice mentioned that for example, crack velocity is generally related to both strain energy density and stress intensity factor; in turn, strain energy density can be directly related to stress intensity factor, and Kirchner’s strain intensity criterion is a direct algebraic derivative of the stress intensity factor criterion (ref. 1). This means that taken separately, none of the criteria can give a satisfactory explanation to all of the observations made earlier. In fact, one of the basic difficulties with previous explanations is their attempt to explain all phenomena by a single criterion. Instead, two or more criteria may be required for explaining the fracture phenomena, as detailed by Rice (ref. 1).

4. Application of Fracture Mirror and/or Crack Branching Constants for the Appropriateness of Stress Measurements

The uniaxial and biaxial flexural strengths of NC132 silicon nitride were previously determined in constant stress-rate testing at 1100 °C in air (ref. 20). The biaxial strength predicted from the uniaxial strength data, based on the Weibull statistics, was substantially higher (60 percent) than the actual biaxial data. This prompted a fractographic analysis to determine both the fracture mirror constant for the rectangular beam specimens tested in uniaxial flexure and the crack branching constant for the disk specimens tested in biaxial flexure. The idea came from the fact that regardless of flaw history (flaw type, stress state, or environment) the crack branching constant of a given material is not greatly different from the fracture mirror constant. Therefore, by comparing the two constants, the appropriateness of the stress measurements determined from each individual testing can be verified. The fracture mirror and crack branching constants were obtained as \( A_m = 9.40 \pm 1.19 \) and \( A_b = 5.66 \pm 1.49 \) MPa√m, respectively, which is already shown in figure 2. This value of fracture mirror constant is significantly higher (66 percent) than the crack branching constant. This indicates that in order to have a value of crack branching constant similar to that of fracture mirror constant, the stress determined in the biaxial specimens would need to have been increased by at least 66 percent (eq. (1)).

Several approaches were made to pinpoint the reason for discrepancy in the stress measurements occurring in biaxial flexure (ref. 20). They included analytical stress calculation, indentation strength method and strain-gaging experiments to determine the related stresses of the biaxial disk specimens. Details regarding these approaches can be found in a previous report (ref. 20). Based on the results of these three approaches, it was found that the desirable state of biaxial stress could not be achieved within the loading-ring diameter of a disk specimen for a given number of loading balls of five (for the upper fixture). The stress occurring at the tensile side of a disk specimen, responsible for fracture, was highest on the loading-ball site (region 'A'), intermediate at the center (region 'C') and lowest between the two adjacent loading balls (region 'B') (fig. 16). Therefore, the strengths obtained from the disk specimens were underestimated since the stress calculation was based on the magnitude occurring at the center of the specimen (ref. 24). Most of the disk specimens failed from the regions (region 'A') directly under the loading balls. Therefore, the obtained failure stress should be modified with an appropriate correction factor. A correction factor of 1.38 was determined from the results of these three approaches. The correction factor of 1.38 yielded a new fracture mirror constant of \( A_m = 7.81 \) MPa√m, which is in reasonable agreement with \( A_b = 9.40 \) MPa√m. This also gave a much better prediction in strength between the uniaxial and biaxial loading conditions. It was found that a reasonably accurate biaxial state of stress within the loading-ring diameter can be obtained by using an increased number of loading balls (at least or greater than 10). Furthermore, it was found that a continuous ring-on-ring configuration with a condition of minimized friction provides another alternative for a proper biaxial strength testing method. A fractographic analysis using the fracture mirror and crack branching constants thus can be utilized as an efficient tool to identify the problem associated with stress measurements, which was demonstrated with the uniaxial and biaxial flexural strength data obtained from high-temperature, constant stress-rate testing for NC132 silicon nitride.
IV. CONCLUSIONS

The fracture mirror and crack branching constants were determined for glasses and advanced ceramics tested under various loading and specimen configurations in an attempt to utilize the constants as a data base on fractography. The ratios of fracture mirror or crack branching constant to fracture toughness were found to be approximately two for most ceramic materials tested. A demonstration to use the two constants as a verification tool for the appropriateness of stress measurements was made for silicon nitride disk specimens subjected to high-temperature, constant stress-rate testing.

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REFERENCES


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<th>Number</th>
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<tr>
<td>13</td>
<td>Norton comp</td>
<td>RT strength</td>
<td>4-pt flexure (4.8/20mm)/beams (3mm×4mm×25mm)</td>
<td>M</td>
<td>19.5</td>
<td>4.64$^3$</td>
<td>6.63(0.11)</td>
</tr>
<tr>
<td>14</td>
<td>Fused silica</td>
<td>RT strength</td>
<td>4-pt flexure (20/40mm)/rods (2mm and 4mm in dia.)</td>
<td>M</td>
<td>7.6</td>
<td>0.79$^4$</td>
<td>2.20(0.33)</td>
</tr>
<tr>
<td>15</td>
<td>Soda-lime glass</td>
<td>RT const. stress-rate test</td>
<td>ring-on-ring (10/32.2mm)/plates (1.5mm×50mm×50mm)</td>
<td>M</td>
<td>5.6</td>
<td>0.75$^4$</td>
<td>1.81(0.28)</td>
</tr>
<tr>
<td>16</td>
<td>Fused silica optical glass fibers</td>
<td>RT const. stress-rate test</td>
<td>tension/ fibers (150μm in dia.)</td>
<td>M</td>
<td>7.6</td>
<td>0.79$^4$</td>
<td>2.10</td>
</tr>
<tr>
<td>17</td>
<td>Soda-lime glass</td>
<td>RT const. stress-rate test</td>
<td>4-pt flexure (20/40mm)/plates (1.2mm×25mm×25mm)</td>
<td>B</td>
<td>5.6</td>
<td>0.76$^3$</td>
<td>3.54(0.64)</td>
</tr>
</tbody>
</table>

Notes: 
1. Fracture toughness evaluated with 1) SEPB method (ASTM PS070); 2) Indentation fracture method; 3) Indentation strength method; and 4) from reference 31.
2. Corrected using a correction factor of 1.38 (see Section III-4).
3. The number in parenthesis indicates ±1.0 standard deviation.
4. The number in parenthesis represents corresponding reference.
Figure 1.—Schematic of fracture mirror, mist, hackle, and branching patterns on typical ceramic fracture surface [1]: $c_f$ represents the critical flaw size, and $r_m$, $r_h$, and $r_b$ represent the fracture mirror size, hackle length, and crack branching length, respectively. A fracture surface and a crack branching pattern represent in A and B, respectively.

Figure 2.—Fracture strength as a function of fracture mirror size (a) and crack branching length (b) for NC132 silicon nitride tested in constant stress-rate testing at 1100 °C in air [20]: (a) four-point uniaxial flexure; (b) ring-on-ring biaxial flexure. The lines represent the best-fit lines based on Eq. (1).
Figure 3.—A typical fracture surface (a) and a typical crack branching pattern (b) of NC132 silicon nitride tested in constant stress-rate testing at 1100 °C in air [20]: (a) four-point uniaxial flexure; (b) ring-on-ring biaxial flexure.
Figure 4.—Fracture strength as a function of fracture mirror size (a) and crack branching length (b) for GN10 silicon nitride tested at room temperature: (a) pure tension [21]; (b) post-impact strength in ring-on-ring biaxial flexure [22].

Figure 5.—Fracture strength as a function of crack branching length for Hexoloy silicon carbide plates tested in ring-on-ring biaxial flexure at room temperature [23].
Figure 6.—A typical crack branching pattern and corresponding fracture surface of a Hexoloy silicon carbide plate tested in ring-on-ring biaxial flexure at room temperature [23].
Figure 7.—Fracture strength as a function of fracture mirror size (a) and crack branching length (b) and (c) for 96 wt % alumina (Alsimag) tested in constant stress-rate testing at room temperature [25]: (a) four-point flexure; (b) ring-on-ring biaxial flexure; (c) ball-on-ring biaxial flexure.

Y-TZP; biaxial Post-impact

\[ \sigma_{r_{b1/2}} = 11.48 \text{ (MPa}\sqrt{\text{m})} \]

AS44 \( \text{Si}_3\text{N}_4 \) Biaxial; post-impact

\[ \sigma_{r_{b1/2}} = 10.85 \text{ (MPa}\sqrt{\text{m})} \]

SN220 \( \text{Si}_3\text{N}_4 \) Biaxial; post-impact

\[ \sigma_{r_{b1/2}} = 8.13 \text{ (MPa}\sqrt{\text{m})} \]

Figure 8.—Post-impact strength as a function of crack branching length for Y-TZP and AS44 and SN220 silicon nitride disk specimens tested in ring-on-ring biaxial flexure [22]: (a) Y-TZP; (b) AS44 silicon nitride; (c) SN220 silicon nitride.
Figure 9.—Fracture strength as a function of fracture mirror size for 30 vol % SiC whisker-reinforced composite silicon nitride (a) and similar monolithic silicon nitride (b) tested in four-point uniaxial flexure at room temperature [13].

Figure 10.—Fracture strength as a function of fracture mirror size for glasses tested at room temperature: (a) as-received fused silica glass rods in four-point uniaxial flexure [28]; (b) etched soda-lime glass plates in ring-on-ring biaxial flexure [27]; (c) indented fused silica optical glass fibers in pure tension [7].
Figure 11.—A typical fracture surface of a fused silica glass rod showing a well-defined fracture mirror formation.

![Fracture Surface Image](image)

Figure 12.—Fracture strength as a function of crack branching length for indented soda-lime microslide plates tested in constant stress-rate testing in four-point uniaxial flexure at room temperature [28].

\[ \sigma_f(r_b)^{1/2} = 3.54 \pm 0.64 \text{ (MPa}\sqrt{\text{m}}) \]
Figure 13.—Typical crack branching patterns and corresponding fracture surfaces of indented soda-lime glass plates tested in constant stress-rate testing in four-point uniaxial flexure at room temperature: (a) high failure stress; (b) low failure stress. T and S represent top view and side view (fracture surface), respectively.
Figure 14.—Plots of fracture mirror or crack branching constant as a function of fracture toughness for the materials studied in this work, except for the soda-lime glass crack branching data (Figure 12). The NC132 fracture mirror constant was corrected using a correction factor of 1.38 (see Section 4). T, B and U denote pure tension, biaxial and uniaxial flexure loadings, respectively.

Figure 15.—Plots of A/KIC as a function of H/KIC for the materials studied in this work, except for the soda-lime glass crack branching data (Figure 12). The line represents a best-fit line based on a relation of A/KIC vs. H/KIC, which is A/KIC = 6.76 x 10^-5 H/KIC + 1.8055 (units in all MPa and m). T, B and U denote pure tension, biaxial and uniaxial flexure loadings, respectively.

Figure 16.—Geometry of a disk specimen in ring-on-ring biaxial flexure used for NC132 silicon nitride disk specimens tested in constant stress-rate testing at 1100 °C in air [20]. The regions A, B and C denote at the tensile side of a disk specimen: region A - directly under the loading ball; region B - between the two adjacent loading balls; region C - at the center.
Crack Branching and Fracture Mirror Data of Glasses and Advanced Ceramics

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Unclassified - Unlimited

The fracture mirror and crack branching constants were determined from three glasses and nine advanced ceramics tested under various loading and specimen configurations in an attempt to use the constants as a data base for fractography. The ratios of fracture mirror or crack branching constant to fracture toughness were found to be approximately two for most ceramic materials tested. A demonstration of how to use the two constants as a tool for verifying stress measurements was presented for silicon nitride disk specimens subjected to high-temperature, constant stress-rate biaxial flexure testing.