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NOTE ON THE RELATION BETWEEN THE COMPRESSIVE STRENGTH OF DEBASED ALUMINA AND ITS USE AS HOT-PRESSING DIE MATERIAL

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Resumé - La durabilité des céramiques d'alumine employées dans les matrices de pressage à chaud est très variable. C'est pourquoi la résistance à la compression de diverses céramiques d'alumine a été déterminée en fonction de la température. Les résultats sont analysés dans leurs rapports avec la microstructure et avec la morphologie des fractures. Il n'y a aucune corrélation entre les données obtenues et la durabilité des matériaux au pressage à chaud. Cette propriété semble être déterminée exclusivement par l'homogénéité de la céramique et par son absence de défauts.

Abstract - Durability of alumina ceramics when used as die material in compressive loading varies widely. The compressive strength of various alumina ceramics was therefore determined as a function of temperature. The results are discussed in terms of the microstructure and fracture morphology. The (hot-pressing) durability of the materials does not correlate at all with the compressive strength data obtained. Instead, this property seems to be determined entirely by the homogeneity and lack of flaws in the ceramic.

I - INTRODUCTION

In our laboratory during hot-pressing experiments in the temperature range of 800 to 1200°C and at pressures of 0.05 to 0.15 GPa, major differences in durability were found between various alumina ceramics. The prime property to study then seems to be the compressive strength of these alumina ceramics. Contrary to the tensile (or bending) behaviour of ceramics the compressive strength has received only scanty attention. A general discussion on the interpretation of the compressive strength of ceramics has been given by Rice (1). He relates compressive strength to hardness. Data for our particular ceramics were lacking, however, and therefore the compressive strength was measured as function of temperature.

II - EXPERIMENTAL

Although the measurement of the compressive strength, $S_C$, is conceptually simple, experience with this type of measurement is not wide spread. A broad discussion on the experimental problems is given by Sines and Adam (2).

For three different types of debased alumina specimens were cut of length 18 mm and diameter 6 mm. These were tested between alumina die's (diameter 40 mm) at a strainrate of about $0.3 \times 10^{-4}$ s$^{-1}$ at various temperatures in air. No interface mate-
rial was used. The measurements were done on a Tinius Olsen Electromatic universal testing machine in a Kanthal wound alumina tube furnace temperature controlled by a microcomputer. The ceramics were also tested at room temperature in water, in order to check on effects of the environment on the compressive strength, $S_c$. The fracture surfaces of the compression specimens were examined by scanning electron microscopy (SEM).

The microstructure was revealed by optical microscopy (OM) and SEM after polishing (4-7 μm diamond) and thermal etching in air at 1250°C for 4 hours. The grain size distribution was determined by linear intercept measurements from approximately 700 grains. The overall chemical composition was determined by chemical analysis while X-diffraction was used to determine the nature of the second phase. The density, $q$, was determined from weight and geometric data. The longitudinal wave velocity, $v_L$, at 10 MHz and the transverse wave velocity, $v_T$, at 20 MHz were determined at room temperature by the pulse-echo technique using Panametrics 5223 equipment. Young's modulus, $E$, and Poisson's ratio, $\nu$, were calculated from $q$, $v_L$, and $v_T$ according to the usual formulae for isotropic materials. The fracture toughness, $K_{IC}$, at room temperature was measured in a three-point bending set-up in dry air (dew point - 42°C) using specimens of size $3 \times 9 \times 45$ mm$^3$ at a cross-head speed of 1.0 mm/min. A notch with a relative depth of 0.15 and a width of about 150 μm was sawn in each specimen. Precracking was done by a Knoop indentation (5 N load) at the notch root on both sides of the specimen.

Blocks of the different materials normally used at hot-pressing dies' typically of 100 mm diameter and 50 mm height, were tested ultrasonically using Sonic Mark IV equipment. Test frequencies used were 5, 10 or 15 MHz with transducers having a diameter of 1/4 or 1/2 inch.

III - RESULTS

In table 1 the main characteristics of the various ceramics are presented. Major differences between the materials are found in the amount and nature of the additives and in the grain size. In all cases the grain size distribution was multi-modal log-normal and the overall modal grain size, $D$, is indicated. The differences between the materials are also reflected by the values of $K_{IC}$. A significant decrease of the compressive strength in water was observed.

Table 1: Characteristics of the various alumina ceramics

<table>
<thead>
<tr>
<th>Material</th>
<th>A</th>
<th>B</th>
<th>C</th>
</tr>
</thead>
<tbody>
<tr>
<td>$q$(g/cm$^3$)</td>
<td>3.85</td>
<td>3.86</td>
<td>3.80</td>
</tr>
<tr>
<td>X-ray (2nd phase)</td>
<td>spinel</td>
<td>-</td>
<td>spinel</td>
</tr>
<tr>
<td>$D$(μm)</td>
<td>15</td>
<td>31</td>
<td>6.4</td>
</tr>
<tr>
<td>wt % Si</td>
<td>0.050</td>
<td>0.10</td>
<td>1.9</td>
</tr>
<tr>
<td>wt % Ca</td>
<td>0.046</td>
<td>0.020</td>
<td>0.31</td>
</tr>
<tr>
<td>wt % Mg</td>
<td>0.58</td>
<td>0.037</td>
<td>0.27</td>
</tr>
<tr>
<td>$E$(GPa)</td>
<td>369</td>
<td>369</td>
<td>347</td>
</tr>
<tr>
<td>$v$</td>
<td>0.236</td>
<td>0.235</td>
<td>0.240</td>
</tr>
<tr>
<td>$K_{IC}$(MPa.m$^{1/2}$)</td>
<td>4.10(0.22)</td>
<td>383(0.08)</td>
<td>5.66(0.20)</td>
</tr>
<tr>
<td>$S_c$(GPa), air</td>
<td>2.11(0.11)</td>
<td>1.90(0.13)</td>
<td>1.83(0.21)</td>
</tr>
<tr>
<td>$S_c$(GPa), water</td>
<td>1.33(0.14)</td>
<td>0.96(0.20)</td>
<td>1.67(0.33)</td>
</tr>
</tbody>
</table>

Sample standard deviations are given in parentheses.

At low temperatures the specimens of all materials essentially exploded forming powder, many tiny chips and some larger pieces. In the temperature range of 1000 to 1200°C the specimens were mostly sheared in the middle.
The data for ceramic care­
ually the average of data from two
nominally the same· materia1s but
produced as rod and plate. As
should be the case, the micro­
structure, chemical composition,
mechanical properties and frac­
ture morphology were the same
within experimental accuracy.
Consequently the data obtained
were averaged.

In fig. 1 the compressive
strength of the various ceramics
is presented. At room temperature
the compressive strength corre­
lates well with the fracture
toughness. Apart from the maximum
in the compressive strength val­
ues for one of the materials, no
large differences in temperature
dependence between the various
ceramics are observed. The rela­
tive change with temperature \( R = \frac{1}{S_c} \cdot \frac{dS_c}{dT} \)
was about \( 1.1 \times 10^{-3} \)
K\(^{-1}\) in the temperature range of
400 to 1200°C.

The data for ceramic C are actu­
ally the average of data from two
nominally the same materials but
produced as rod and plate. As
should be the case, the micro­
structure, chemical composition,
mechanical properties and frac­
ture morphology were the same
within experimental accuracy.
Consequently the data obtained
were averaged.

Fig. 1. Compressive strength of the various
ceramics as a function of temperature.

For all ceramics clear twinning phenomena were observed upto 800°C. These phe­
nomena were occasionally observed at much higher temperatures. In general the frac­
ture mode was mainly transegranular below about 800°C, changing to mainly intergran­
ular above that temperature.

On the fractographs various microfracture phenomena were observed. Microcracks due
to twinning or originating from grain boundary or triple junction voids but also
from pores within a single grain. In a number of cases the linking of microcracks
could be identified. In fig. 2 some of these phenomena are shown.

Fig. 2. Microfracture phenomena as detected on the fracture surfaces.
Left : microfracture originating from a pore inside a grain.
Right: twinning linked to microfracture originating from a grain boundary
pore.
From the scarce literature data one can distinguish two classes of alumina ceramics as far as $S_C$ is concerned. Rather pure materials with an $S_C$ value typically above 3 GPa and an $R$ value of about $5 \cdot 10^{-4} \text{ K}^{-1}$ (3-8) and the less pure materials with $S_C$ typically around 2 GPa and an $R$ value of about $1 \cdot 10^{-3} \text{ K}^{-1}$ (4-9). (The material of Nash (9), although not explicitly stated, probably contains also SiO$_2$ since he used commercially available materials).

The values obtained for our rather impure ceramics (see table) at room temperature are typically 2 GPa and thus consistent with those of the second class. Also the $R$ value of our ceramics is similar to the rate observed by Dawihl and Dörre (4) for their silicate containing material. Consequently the conclusion must be that for these ceramics the flaw density is higher (lower strength value) and that cavitation is easier (larger rate of decrease).

Most studies of the compressive strength of alumina were done on translucent material (Lucalox), particularly by Lankford (5-8). A significant strain rate effect was observed (5) but an influence of the environment, so manifestly present in tensile testing, was not found. In testing these rather pure materials at room temperature, twinning was observed at least to some extent over a widely varying strain-rate range (5). The twinning led to microcracking starting mostly at grain boundaries. Microcracks also started at voids at triple junctions. After nucleating some growth of the microcracks took place. The coalescence of the microcracks then led to catastrophic fracture. At higher temperatures cavitation in the (supposedly glassy) grain boundary phase was said to take place. Growth and coalescence again led to catastrophic fracture. The observed maximum in compressive strength is explained by a decreased effectiveness of the twinning mechanism while the grain boundary phase is not yet easy flowing (6).

All the phenomena discussed extensively by Lankford for translucent alumina were observed on these debased alumina's as well. In particular the suggestion that pores are probably also a source of microfracture (7) has been found true.

A maximum in $S_C$ was observed for ceramic C only. This is probably related to the large amount of SiO$_2$ mainly present in the grain boundary phase. The fracture toughness of debased alumina shows a maximum at about 800°C as shown by Davidge and Tappin (10), which is explained by a reduction of the crack tip stress by viscous flow of the glassy second phase, effectively increasing energy dissipation. At slightly higher temperature viscous grain boundary sliding, introducing pores, results in a strength decrease. The viscous flow dissipation mechanism is likely to operate in compressive fracture as well, besides the already mentioned twinning and cavitation mechanisms, but only if enough glassy grain boundary phase is present. The absence of a maximum in $S_C$ for the other ceramics is probably due to the much easier cavitation as compared with translucent alumina.

Contrary to the results of Lankford (5), a significant environmental effect was observed by Nash (9). For a 97.5% pure alumina the value of $S_C$ dropped from 1.7 GPa when tested in air to 1.2 GPa when tested in a physiological salt solution. Similarly for 99.5% pure material, $S_C$ dropped from 2.2 GPa to 1.7 GPa. A significant decrease in $S_C$ was also observed in our case. This probably due to the fact that the specimens used by Nash (and ours) were ground on the outer surfaces while for the specimens of Lankford these surfaces were as-fired. The density of the flaws on the outer surfaces, sensitive to the environment, is much smaller in the latter case thus reducing the environmental effect.

V - RELATION WITH HOT-PRESSING PRACTICE

Comparing the values of $S_C$ with the pressures normally applied during hot-pressing it is clear that the latter are generally much lower. Nevertheless fracture occurs. The order of longer durability in (hot-pressing) practice is: A>B>C. This order does correlate weakly with the room temperature compressive strength
(order: $A > B = C$) but this correlation is irrelevant since hot-pressing is done at higher temperatures. It does not correlate with the high temperature compressive strength ($A = B = C$). Slow crack growth may be important since at room temperature a clear environmental effect was observed. More important seems to be, however, the homogeneity of the material when delivered in large blocks. For material $A$ the as-delivered blocks used for hot-pressing are essentially free of macroflaws (as tested by pulse-echo technique) but the other ceramics often show some large inhomogeneities probably due to extruding or pressing (Fig. 3). Failure of the blocks during hot-pressing is often originating from these defects. This makes the $S_C$ measurements irrelevant. The processing of the ceramics is thus, again, shown to be of vital importance even in the case of compressive loading which is usually considered as a safe type of loading.

Fig. 3. Ultrasonic pulse echo patterns (A-scans) of a particular block of ceramic $B$ taken at 10 MHz and gain 63 dB.

Left: proper pattern from the edge showing only the transmitting pulse (left) and backside echo (right).

Right: pattern from the centre showing also two defects.

VI - ACKNOWLEDGEMENT

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REFERENCES