Influence of granule characteristics on fabrication of translucent alumina ceramics with high strength and reliability

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Translucent alumina ceramics with high strength and reliability were fabricated through dry pressing of industry-grade granules and subsequent vacuum-sintering prior to hot isostatic pressing (HIP). The raw materials consisted of two types of granules, containing acrylic binder with and without 500 ppm MgO. The internal structures of samples in each manufacturing step were evaluated by the liquid immersion method using an optical microscope. Large defects and their transitions were observed in the samples after each processing step. The fabricated ceramics had high relative density (>99.6%) and very fine microstructures with grain sizes smaller than 1 μm. In the granule system with the binder non-uniformly distributed, interstices in the granules were observed in the powder compacts and samples after vacuum sintering. Alumina ceramics derived from the granule system with uniformly distributed binder and MgO additive exhibited high translucency with 50% in-line transmittance and strength greater than ~700 MPa. This result suggests that characteristics of the granules considerably affect not only the packing structures in powder compacts but also the microstructure and properties of the sintered ceramics.

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1. Introduction

Advanced ceramics have been developed for various applications, and their microstructures have been studied and controlled to obtain functional properties. For example, translucent polycrystalline alumina with large grains and highly dense structures has been fabricated by using fine, high-purity (>99.99%) particles and a small amount of sintering additives such as MgO.1–5 Sintering has been conducted at temperatures greater than 1800°C under a controlled atmosphere containing H2. The use of finer primary particles promotes densification and elimination of pores during sintering. High-purity raw materials reduce the scattering of light due to contamination and prevent excess grain growth. Sintering additives such as MgO retard grain growth before densification and are effective at reducing the closed pores retained within the grains. A higher sintering temperature reduces grain boundaries due to grain growth up to several tens of microns. A H2 atmosphere contributes to the easy diffusion of gas molecules from within the pores to the outside of the samples.

For optical use, ceramics are required to exhibit both translucency and good mechanical properties.6–9 Translucent alumina ceramics manufactured with the above mentioned processing method have not exhibited high strength,6,9 because grain growth at high temperature reduces the grain boundary but simultaneously causes the growth of large pores, which degrades mechanical properties. These large pores could not be completely removed. To achieve translucency and good mechanical properties simultaneously, other strategies for microstructure control should be considered. One solution is to control the processing for obtaining a very fine microstructure.6–14 Morinaga et al. reported translucent alumina with submicron grains formed by injection molding and sintering with hot isostatic pressing (HIP).15 Krell et al. reported the fabrication of high-strength transparent alumina through the wet-shaping approach combined with sintering under normal and HIP conditions at 1400°C.16–19 For HIP, sintering is generally carried out under capsule-free condition by using pre-sintered samples that contain closed pores. Ikuse reported that for fabrication of YAG ceramics, the sample was presintered in vacuum prior to HIP.11–13 Shen et al. and other researchers reported the spark-plasma sintering of fine-grained alumina.20–23 They reported that densification must be achieved by sintering at low temperatures.

From an industrial viewpoint, although many routes exist for forming and sintering, conventional forming and sintering processing are preferred.24 Granules are used in the industry primarily to achieve high productivity for dry packing as well as high and homogeneous packing density. However, the characteristics of granules have unfavorable effects on the powder processing and properties of ceramics.25–28 For example, hard, elastic granules tend to form large defects in green compacts as well as in sintered ceramics.29 The binder also affects the structure through its distribution in the granules. PVA binder tends to segregate on the surface of the granules, leaving large processing defects in the compact after de-binding.29–31

Direct observation of the structure during processing provides important information for controlling the structure of ceramics.12,32 Observations using the liquid immersion method (LIM) revealed that the interstices between granules and dimples of granules changed to large defects in the ceramics.33 In the LIM, the granules or powder compact was immersed in an appropriate liquid having the same refractive index as that of the material. For alumina, methylene iodide or borononaphthalene are suitable as the liquid, and it results in transparent alumina powder compacts. Therefore, the internal structure is observed with an optical
microscope in transmission mode. Conversely, the internal structure of a sintered sample can be observed with an optical microscope in transmission mode by using a sample thinned to 100 μm because such a sample would show translucency.34,37 Thus, large defects can be identified at each step during manufacturing.

This present study aimed at fabricating alumina ceramics with high reliability and high density through powder processing using industrial granules by focusing on structural changes during sintering. We used granules with a binder layer as well as granules with a thin binder layer and a small amount of sintering additive, MgO.

2. Experimental

2.1 Granules and their characterization

Two types of commercial alumina granules were used as raw materials (DS31 and DS 90M, Taimei Chemical K. K., Japan). The size of the primary particles in both granules was 0.15 μm as received from the supplier. DS31 contains an acrylic binder and DS 90M contains an acrylic binder and 50 ppm MgO as a sintering additive. Hereafter, DS31 and DS 90M are referred to as “granule B” and “granule M”, respectively. LIM was used to examine the internal structure of the granules. In this examination, the granules were immersed in bromonaphthalene (refractive index: 1.66) to make them transparent (refractive index of alumina: 1.77), and the internal structure was observed using an optical microscope in transmission mode (DX50, Olympus, Japan). A scanning electron microscope (JSM-5310L, JEOL, Japan) was used to examine the morphology and structure of the granules in detail. Micro-compaction equipment (MCTE-500, Shimadzu, Japan) was used to examine the compaction behavior of individual granules. For equilibration with moisture, the granules were placed in an ambient atmosphere for a day. A granule with a diameter of approximately 50 μm was selected using the optical microscope for examination. The temperature was controlled at 20°C. The tensile strength of the granule (S_t) was determined using the following equation proposed by Hiramatsu et al.:37

\[ S_t = 2.8P_f / \pi D^2 \]

where \( P_f \) is the applied load at fracture and \( D \) the average diameter of the granule.

The granules were uniaxially pressed in a die with double action by a universal mechanical testing machine (Autograph AG1, Shimadzu, Japan) at a crosshead speed of 1.0 mm/min. The test was performed at 20°C for both granules. A compaction curve was constructed from the load-displacement curve automatically recorded for each granule. The relative density (R.D.) of the compact was determined during the compression test by using the following equation:

\[ \text{R.D.} = 100 w / (A h \rho), \]

where \( w \) is the weight of filled granules, \( A \) is the base area of the compact (cross-sectional area of the pressing punch), \( h \) is the height of the compact in the die, and \( \rho \) is the theoretical density of alumina: 3.987 × 10^2 kg/m^3.

2.2 Forming and sintering

Granules were placed in a die (45 mm × 55 mm) and pressed at 20 MPa; subsequently, they were cold isostatically pressed at 200 MPa. After de-binding at 500°C for 30 min in air, the compact was heated to 1350°C for 2 h in vacuum and then cooled to room temperature. The sample sintered in vacuum (1 Pa of air) was subsequently sintered at 200 MPa and 1300°C for 2 h in an Ar atmosphere with HIP (Dr-HIP, Kobe steel, Japan).

2.3 Characterization and mechanical properties

LIM was applied to examine the internal structure of the powder compact, as with the granules. The powder compact, which was sliced and ground to a thickness of 100 μm, was immersed in bromonaphthalene, and the internal structure was observed with an optical microscope in transmission mode. The structure of the sintered body was also observed using an optical microscope in transmission mode. The samples were sliced, ground, and polished to a thickness of 100 μm to make them translucent. To examine the eliminating process of pores by HIP, the 100-μm-thick sample heat treated in vacuum was treated with HIP. The transmitted image after HIP was compared with that before HIP. The microstructures of the samples were observed using a scanning electron microscope. Before observation, the polished samples were etched thermally at 1250°C for 30 min to clarify the grain boundary. The surfaces of samples were then coated by gold sputtering.

The bending strength of the sintered body was measured by four-point bending tests at a cross-head speed of 0.5 mm/min by using a universal mechanical testing machine. The samples were 3 mm thick, 4 mm wide, and 40 mm long.

3. Results and discussion

Figure 1 shows SEM micrographs of the primary particles. Figure 2 shows SEM and LIM micrographs of the granules used in this study. Granule B appears to have a spherical shape with a dimple. In the LIM micrograph, however, its internal structure is not uniform and shows a rim at the outline of the granule, which is the binder segregated around the surface. It appears dark because the binder scatters light owing to the large mismatch in the refractive indices. Granule M appears to have a non-spherical shape with a dimple. Its internal structure is uniform and shows a thin binder shell. The shape of granules may be influenced by the insufficient dispersion of ceramic particles in the slurry during spray drying.

Figure 3 shows a Weibull plot of the strength of a single granule, which was measured through the compaction test of a single granule. Granules B and M have average strengths of approximately 0.8 and 0.3-0.4 MPa, respectively. The binder shell on granule B, as shown in Fig. 2, may support the applied stress. This is why granule B had higher strength, and the data could be explained by the Weibull plots.

Figure 4 shows typical compaction curves for die pressing using these granules. The relative density of the compacted granules gradually increased with increasing applied stress for both granule systems. The compaction curves depended on the mechanical property of a single granule, as reported in previous...
The relative densities at low stress corresponded to the filled density in the metal die. Changes in slope appear at two locations, ~1 and ~10 MPa, in each curve. The first change in slope corresponds to the starting point of granule deformation. The stress at the point at which there is a change in the slope of the curve for granule B is higher than that of granule M. This result is associated with the strength of a single granule. The second change corresponds to the starting point for particle rearrangement or densification within the granule in the compact body. A larger change in the relative density was observed in granule M than in granule B for an applied stress of up to 100 MPa. The transfer of stress from the punch into the sample for the hard granule system is generally higher than that for the soft granule system. It is therefore possible for the hard granule system to deform well all over the sample.

Figure 5 shows the structure of the powder compacts and de-bindered samples examined with LIM using the optical microscope in transmission mode. (a) Sample made from granule B; (b), (a) after de-binding. (c) Sample made from granule M; (d), (c) after de-binding.
A dimple is observed at the center of each granule. The frequency of dimples in the granule B system is greater than that in the granule M system.

Figure 6 shows the structures of the vacuum-sintered samples and HIP-treated samples observed using the optical microscope in transmission mode with a thinned specimen (200 µm). Many large defects are observed as black shadows in the vacuum-sintered samples. The relative density of samples is 95% for both systems. This suggests that sintering occurred easily in the dense part of the sample and that the loosely packed regions such as the granule dimple and boundary remained and developed into large defects. The large pores tend to gather the fine pores around themselves during sintering, according to classical sintering theory. Conversely, many large defects appear to have been eliminated by HIP treatment. The relative densities of HIP-treated samples with granules B and M increased to 99.6 and 99.8%, respectively. Grain growth occurred with HIP treatment. Defects of size 20–30 µm are indicated as grey shadows in the internal structure. Many defects are observed in the granule B system.

However, the circular cloud-like pores of outer size 50–100 µm are occasionally observed in both HIP-treated samples, as shown in Fig. 7. A crack can be seen at the center of the circle in Fig. 7(a). The gray shadow around the circle comprises very fine pores. For the granule M system, the cloud pores without center defects were observed, as shown in Fig. 7(b). To examine the effect of HIP on elimination of pores, particularly, to clarify the circular cloud-like pores, the structures at the same position before and after HIP were compared. Figure 8 shows micrographs taken using an optical microscope in transmission mode. An expanded view of the large pores after HIP is also shown. In Fig. 8, some large pores are observed at the same position after HIP treatment. The original pore (<30 µm) in the sample remained after HIP treatment. In the expanded view of the large pores after HIP [Fig. 8(c)], the outlines of pores are ambiguous and similar to the structure shown in Fig. 7(b). This result suggests that cloud-like pores are formed on the eliminating large pores through HIP treatment. The grains near the large pores move, as in creep deformation, to close large pores owing
to HIP, because the size of the grains is very fine, less than 1 μm, as shown in Fig. 9. As a result, the traces of the moved grains are seen as cloud-like regions with fine pores.

Figure 9 shows the microstructures of both samples. The grain sizes in samples made from granules B and M, which were measured by the intercept method, were 0.75 and 0.63 μm, respectively. The effect of MgO was seen in the higher densification and finer microstructure in samples made from granules M than in samples made from granules B. Figure 10 shows the appearances of both samples. The thicknesses were 800 μm. The samples made from granules B and M show translucency, and the in-line transmittances of HIP-treated samples were 50 and 15%, respectively, at a wavelength of 650 nm, as shown in Fig. 11.

Figure 12 shows the Weibull plots of the flexural strength of ceramics made from granule B and granule M. The average strength of ceramics using granule M was 691 MPa, which is greater than that of ceramics using granule B (565 MPa). In detail, although the maximum strength of the granule M system is the same as that of the granule B system, the minimum strength of the granule B system is considerably weaker. The Weibull modulus of the sample made from granule M is 18.0, whereas that of the sample made from granule B is ~6. While HIP treatment under a pressure of 200 MPa at 1300°C was effective for reducing the size of large defects, the results of the very weak strength of the sample made from granule B suggest the limitation of pore size. Particularly, as shown in Fig. 6(a), it seems that a few crack-like defects of size ~50–100 μm are present in the sample made from granule B with vacuum-sintering. Those defects may remain even after HIP.

Here, the equivalent crack length is calculated from the strength value according to the Griffith equation:

$$\sigma = \frac{K_{lc}}{\sqrt{\pi a}}$$

where $\sigma$ is the fracture stress, $K_{lc}$ is the fracture toughness, and $a$ is half of the length of a crack. The equivalent crack length, $2a$, decreases in inverse proportion to the square of the strength. If the strengths are 400 and 700 MPa and fracture toughness is 3.5 MPa m$^{0.5}$, the equivalent crack lengths are ~49 μm and ~16 μm, respectively. In the sample made from granule M of strength 700 MPa, defects of size ~20 μm were observed in Fig. 6(d). Conversely, large defects of size ~50 μm are shown in Fig. 7(a). These pores may govern the mechanical strength. If this property is to be improved further, the pore size in the powder compact...
must be reduced by controlling the granule characteristics.

4. Conclusions

Translucent alumina ceramics with high strength and reliability were fabricated by dry-pressing and subsequent sintering with HIP. The structure was observed during manufacturing by using an optical microscope in transmission mode. In the experiment, industrial granules containing an acrylic binder and a small amount of MgO were used as the raw material. The characteristics of the granules govern the final structures from the packing structure in the powder compact to the structure of sintered ceramics. In the granule system with binder, granule B, processing defects such as dimples and interstices of granules were remarkably observed in the powder compact and samples after vacuum sintering. Even after HIP treatment, many large defects remained in the sample. Conversely, the size of large pores in the granule system with the binder and the sintering additive MgO, granule M, are small compared with that of granule B. A very thin binder layer and MgO additive in granules play an important role in structure formation. We achieved 50% in-line transmittance, strength of ~700 MPa, and Weibull modulus of 18 in the ceramics formed with granule M. These properties should be improved further by controlling the granule characteristics.

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References