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Improved Transparency and Hardness in α-alumina Ceramics

Fabricated by High-pressure SPS of Nanopowders

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Abstract

Nanocrystalline alumina powder with an average crystallite size of $\leq 50$ nm has been consolidated by spark plasma sintering (SPS) and hot pressing (HP) with a view to achieving dense, fine grained alumina bodies that display transparency. When as-synthesised powder was densified directly, excessive grain growth resulted from both the SPS and HP techniques and hence a large final grain size was observed. Attempts to improve the uniformity of the green microstructure prior to densification were unsuccessful when spray freeze dried granules were used, whether pre-pressed into a compact or not. The use of 53% dense slip cast green compacts, however, enabled final density of $\sim 99.96\%$ and a mean grain size of $\sim 0.32$ $\mu$m to be achieved when SPS conditions of 1200°C and 500 MPa were
applied for 20 minutes. These samples offered in-line transmittance values of up to ~80% and microhardness values of 22 GPa.

**Keywords:** Nanocrystalline; Alumina; SPS; Hot press; Transparent

1. **Introduction**

Nanocrystalline ceramics typically have a mean grain size of <100 nm. The motivation for manufacturing and studying such ceramics lies in their potential to yield unusual and useful properties such as superplasticity, high hardness and extremely low thermal conductivity [1-5]. Owing to its combination of high hardness, corrosion resistance, thermodynamic stability and relatively low cost alumina is the most common engineering ceramic. It is used for a very wide range of applications, including armour, hip implants, thread guides, cutting tools, valves, water tap mixers and many other applications [6]. One particular advantage offered by very fine grain size alumina is that it is transparent when the final density is ≥99.95% of theoretical and the mean grain size is below ~1 μm [7]. It should be noted, however, that, unlike ceramics with cubic crystal structures, polycrystalline alumina is birefringent [8]. Hence mean grain sizes smaller than the incident light wave length are needed to limit scattering losses and achieve true transparency [8]. For example, Krell, et al. [9] fabricated alumina with 99.9% of theoretical density and a mean grain size of ~0.5 μm, with 0.8 mm thick samples exhibiting real in-line transmissions (RIT) of 60% at a wavelength of 650 nm [9].

Highly dense, submicron, transparent ceramics have traditionally been obtained using either hot pressing or hot isostatic pressing of fine, submicron powders. More recently, spark plasma sintering has been investigated as a tool for achieving even finer microstructures. Essentially similar to hot pressing, the method of heating used – the passage of an electric current through a graphite die set – results in heating rates as high as 10⁶ K s⁻¹, lower sintering temperatures and reduced time at temperature [10-12]. For example, fully dense
alumina bodies with a mean grain size of 0.5 μm were obtained at only 1200°C with dwell times of 3-10 minutes by SPS [13,14]. Grasso et al. [15,16] fabricated transparent, pure alumina bodies by densifying commercial alumina powder with a crystallite size of 130-200 nm at temperatures as low as 1000°C under 500 MPa. Fully dense, fine grained alumina with porosity <0.05% was obtained with an in-line transmission of about 64% at a wavelength of 645 nm [17]. Use of dopants such as La\(^ {3+} \) has also been investigated to fabricate highly dense transparent ceramics with 71% real in-line transmission (at 640 nm) using slip cast samples prepared from commercial ultrafine alumina doped with La\(^ {3+} \) using SPS to achieve densification [18].

In this work, it is shown that high-pressure spark plasma sintering using an in-house synthesised nano α-alumina powder, with an initial particle size of ≤50 nm, can lead to the formation of dense transparent alumina bodies with transmittance values of ~80%. The effect of employing different green forming methods, including slip casting and spray freeze granulation, has been investigated. Hot pressing, an alternative pressure-assisted technique, has also been used to compare the densification processes.

2. Materials and methods

The powder used was pure α-alumina nanopowder with a specific surface area of ~35 m\(^2\)g\(^{-1}\) and average crystallite size of ≤50 nm. Details of the synthesis method are provided elsewhere [19,20], but, in brief, the powder was synthesised from an aluminium chloride and ammonia solution using a precipitation-based method followed by a two-step heat treatment route to prevent the high degree of agglomeration occurring during conversion of the precursors into metal oxides [21].

*Green forming:* 60 wt% solids content nanoalumina suspensions were produced by mixing appropriate quantities of deionised water and an anionic polyelectrolyte (Dolapix CE-64,
Zschimmer & Schwarz GmbH, Lahnstein, Germany) as a dispersant and stirring for 1 hour. The nanoalumina powder, weighed to 0.01 g, was added subsequently and the mixture ball-milled for 12 hours using alumina balls to achieve an homogeneous suspension. This was subjected to ultrasonic agitation (SoniPrep, MSE Ltd, London, UK) for 3 minutes whilst placed in a water bath, using 150 W at 15 kHz in clean glass beakers, whilst the after which the latter were sealed using Parafilm to prevent water evaporation and left for 1 hour to cool and attain equilibrium. Finally, the samples were held at a reduced pressure of 60 mm mercury for 10 minutes to remove as many air bubbles as possible.

Slip casting was performed using a 15 mm thick acrylic sheet with an array of nine 15 mm diameter drilled holes, clamped to a ~50 mm thick flat plate of Plaster of Paris, PoP (Prestia 23 casting plaster, Lafarge, Cambridgeshire, UK), Figure 1. The latter was produced using a PoP-to-water ratio of 60:40. Lubricant (silicone spray; Ambersil, CRC Industries UK Ltd, Bridgwater, UK) was applied to the inside surfaces of the holes in the acrylic sheet to ease green body removal followed by the suspension being cast into them. This approach allowed up to 9 samples to be slip cast simultaneously, the dewatering process occurring into the plaster plate beneath the acrylic sheet. Drying of the cast green bodies was carried out under ambient conditions of ~20ºC and 50% humidity for 72 hours before samples were removed from the acrylic sheet with great care. Following this preliminary drying, all the samples were placed on a Teflon sheet for further drying at 60ºC in an oven; the Teflon sheet assisted with sample shrinkage by providing minimal resistance.

Granulated powders were also produced from a similar slurry to that used above by spraying the suspension into liquid nitrogen followed by drying the frozen granules in a freeze dryer. This suspension included 2 vol. % Freon 11 (Spex CertiPrep. Inc, UK) as a foaming agent to weaken the granules and 0.5 wt% PVA (Fluka, UK) as a pressing binder. Spraying was performed using a twin fluid atomiser (BÜCHI Labortechnik AG, Postfach, Switzerland); the suspension flow rate was controlled via a compressed air flow from a VT 150 air compressor.
(Bambi Air Compressors Ltd, Birmingham, UK) using a flowmeter. Once the excess nitrogen had evaporated off, the frozen granules were freeze dried in a benchtop freeze drier (Virtis® Benchtop SLC, New York, USA). The process of freeze drying took at least 48 hours, depending on the amount of sample required to be dried. As a final step, sieving was undertaken to extract the fraction between 125 and 250 μm to maximise the flowability [22].

Dry pressing was carried out using a Lloyds mechanical testing machine (L10000 Tensometer, Lloyds Instruments, Fareham, UK) in a single-action, hardened steel die with a diameter of 10 mm. About 0.5 g of powder was used for each pellet, which was pressed at 480 MPa for 1 minute.

Removal of the organics present in both slip cast and die pressed green bodies was achieved by heating at a rate of 0.5°C min⁻¹, with dwells at 100°C, 200°C, 300°C and 400°C for 30 minutes each, and finally at 700°C for 2 hours before being cooled down to room temperature at a rate of 0.5°C min⁻¹. After the binder removal stage, all the samples were visually inspected for the presence of any defects such as cracks. The relative density of both slip cast and die pressed bodies was measured after binder removal by both the Archimedes technique using mercury and by measurement of mass and volume.

**Hot pressing (HP):** The as-synthesised nanoalumina powder was directly hot pressed using the facility at the University of Exeter, UK (8538 UEO, Systeme GmbH, Frankenblick, Germany). The powder was poured into the die, tapped to a level surface and then heated initially under vacuum until 900°C and then in argon; the heating rate was 10°C min⁻¹ for all the experiments. When the temperature reached the desired sintering value, in the range 1400-1700°C, see Table 1, an external pressure of 40 MPa was applied to both ends of the die using graphite plungers; this remained constant throughout the sintering process, which lasted 5 hours. Subsequently, the load was removed and the samples were left to cool down.

Samples of 50 (length) × 50 (breadth) × 2.4 (height) mm were obtained after densification.
Spark plasma sintering (SPS): Work started with the direct spark plasma sintering of as-synthesised nanoalumina powder using the facility at Queen Mary University, UK (FCT HP D 25, FCT Systeme GmbH, Frankenblick, Germany). The conditions used are summarised in Table 1. Two sample sizes were sintered; those densified at 70 MPa were 20 mm in diameter, whilst when 500 MPa or 1000 MPa was used the sample size had to be reduced to 5 mm in diameter. All samples were ~2.5 mm thick after densification. The die used for the two higher pressures was composed of outer and inner graphite dies with SiC punches. The powder was heated under vacuum at a rate of 100°C min\(^{-1}\) to an intermediate dwell point of 700°C, where the pressure was applied. Subsequently, the temperature was raised to the desired sintering temperature, followed by a dwell time of 30 minutes (70 MPa) or 20 minutes (500 MPa and 1000 MPa). Optical pyrometry was used to measure the temperature; it was focused on a hole that penetrated halfway through the inner graphite die. The sintering kinetics were monitored from the variation of the total height of the graphite die assembly with temperature. The applied pressure was removed during the cooling cycle, which involved a rate of 100°C min\(^{-1}\). Additional experiments were performed using the SFD powder which, it was expected, would compact to a higher density when the pressure was applied during the SPS cycle. Subsequently, die pressed and (thoroughly dried) slip cast samples were SPSed, Table 1.

Characterisation: To eliminate the layers affected by heavy carbon contamination from the dies used for both HP and SPS, the surfaces of the sintered samples were removed using a grinding disc with a grit size of 80 and their densities were measured by the Archimedes technique using water. Microstructures were investigated by FEG-SEM (Leo 1530VP FEGSEM, LEO Elektronenskopie GmbH, Oberkochen, Germany) after polishing and thermally etching the samples for 10 minutes at ~100°C below the sintering temperature for each sample. As a typical polishing cycle, mounted samples were initially plain ground using a grinding disc with a grit size of 220 (MD-Piano, Struers Ltd, Solihull, UK), followed by
polishing with 600 and 1200 grit-sized discs (MD-Piano, Struers Ltd, Solihull, UK) using water as the lubricating and cooling media. Subsequent polishing continued using polishing cloths, MD-Plan and MD-Dac (Struers Ltd, Solihull, UK), and stable diamond suspension containing a mixture of high performance diamonds and cooling lubricants, DP-Plan and DP-Dac (Struers Ltd, Solihull, UK), consisting of particles of 9 μm and 3 μm in size, respectively.

At least 300 grains were measured in each of three micrographs to determine the mean grain size for each alumina sample. The average grain size was calculated using the equation:

\[ D = \frac{L \times A_t}{M \times N} \]  

Eq. 1

where D is the equivalent average grain diameter, L is the length of the superimposed lines, A_t is a shape correction factor of 1.56 used to convert the two dimensional grain size to three dimensional [23], M is the magnification and N is the number of intercepts or intersections.

All the disks were polished to a mirror finish on both sides for optical characterisation using the conditions outlined above. The in-line transmission of the samples was measured using a Shimadzu Fourier Transform Infrared spectrometer (FTIR 8400S, Shimadzu, Tokyo, Japan) for the infrared range of 2.5-10 μm and a UV-VIS double-beam spectrometer (Lambda 35, Perkin Elmer, Shelton, USA) for the visible range of 300-800 nm.

The microhardness of the sintered bodies was determined after polishing using the approach described in ASTM E 384 (HM-124 micro hardness tester, Mitutoyo Corporation, Kawasaki, Japan). A square-based pyramidal diamond indenter with face angles of 136° was used by applying a load of 1 kg for 15 seconds. After removing the load, the lengths of the diagonals of the indentation on the sample were measured via a microscope and the Vickers hardness of the samples calculated using:

\[ HV = 0.0018544 \times \frac{P}{d^2} \]  

Eq. 2
where HV is the Vickers hardness in GPa, P is the applied load in N, and d is the length of the long diagonal of the indentation in mm.

3. Results and Discussion

Figure 2 shows the fracture surfaces of green slip cast and die pressed bodies with density values of ~53% of theoretical density. The latter was pressed using granulated powders at 480 MPa. Lower level of agglomeration was observed on both fracture surfaces particularly for the slip cast compact; this could be due to the improved dispersion of alumina nanopowder in the suspension via breaking up the agglomerates using an appropriate dispersant as well as a further sonication step.

3.1. Hot pressing

The density and mean grain size values of the sintered bodies made using the as-synthesised powder are shown in Table 2. Although hot pressing at both 1600°C and 1700°C yielded translucent compacts with relative densities of ~99.9%, the bodies possessed coarse microstructures, with average grain sizes of ≥7 μm and grains of over 10 μm frequently being observed, Figure 3. Further work on hot pressing was therefore discontinued.

3.2. Spark plasma sintering

Table 2 also shows the relative density and mean grain size values of the bodies sintered using SPS and produced directly from the as-synthesised nanoalumina powder. As expected, by increasing the pressure up to 500 MPa, higher relative densities and lower mean grain sizes were achieved and at lower temperatures compared to samples compacted at 70 MPa. The use of 1400°C/70 MPa (SPS-70-14) and 1200°C/500 MPa (SPS-500-12) both led to alumina bodies with densities of ~99.9% of theoretical density and, consequently, these compacts displayed a degree of translucency, as shown in Figure 4a & b. Note, however, that sample SPS-70-14 had a mean grain size of ~6 μm and some grains of >10 μm, Figure
4c, whilst for SPS-500-12 it was approximately one tenth this size, Figure 4d. Nevertheless, it will be observed that some larger grains were still visible. Unevenness in the compact colour, which is usually caused by carbon contamination occurring during spark plasma sintering, was observed for SPS-70-14 as for the hot pressed samples.

The experiment that involved the use of 1 GPa led to a final density of only ~99.5% of theoretical; this was due to the temperature being limited to only 900˚C by the properties of the graphite used for the die. As a result, the body was opaque, but the mean grain size, at 200 nm, was extremely fine, Figure 5.

The unevenness in the grain size of these samples could be induced by the presence of agglomeration in the as-synthesised powder [19]. It is well known that agglomerates, which are denser than the surrounding matrix, progress up the sintering curve faster leading to excessive grain growth and the solution is to ensure that the packing of the particles is as uniform as possible [24]. Hence spray freeze dried, SFD, granules were produced for subsequent densification using SPS, both as a loose granulated powder and as pre-compacted bodies. However, use of SFD powders resulted in lower final densities compared to the densification of the as-synthesised powder, Table 2 and hence this approach was discontinued.

Densifying die pressed bodies made from the SFD granules was also unsuccessful; once again the final density was <99% of theoretical and, consequently, the sintered bodies were opaque. The use of slip casting, however, was more successful owing to the superior green microstructures achieved. Sample SPS-500-12-SC achieved 99.9% of theoretical density, whilst also retaining a very fine mean grain size of ~300 nm; half the size of that for sample SPS-500-12. The grain size of the densified body made from the slip cast green compact was also fairly uniform, Figure 6. Consequently, the sample was transparent. Decreasing the
sintering temperature to 1100˚C resulted in translucency since, although the mean grain size, at 250 nm, was finer, the density was lower at ~99.6% of theoretical.

3.3. IR and UV-Vis transmission measurements

Figure 7a shows the UV-Vis transmission spectra for a selection of the samples produced as measured using a UV/VIS spectrophotometer (Lambda 35, Perkin Elmer). The plot also shows the in-line transmission values of alumina compacts investigated in different studies. It can be seen that in the current work the transmittance values increased with increasing wavelength; samples SPS-70-14 and SPS-500-12-SC both had total transmissions of ~75% at 645 nm, whilst the other samples, with their lower densities, exhibited lower transmittances. It also can be seen that the transmittance values over both wavelength ranges are broadly comparable to data measured by other researchers. The data included in Figure 7b presents the IR transmission spectra for the manufactured alumina compacts and, in the inset, for ceramics such as alumina, spinel, yttria and YAG. It can be seen that, as expected, the transmittance was very low, <50%, for the HP samples owing to their very low densities. Samples SPS-70-12, SPS-70-14 and SPS-500-12 all showed very similar behaviour, with maximum transmittance values in the range of 73-75%. Sample SPS-500-12-SC, which had been slip cast, however, showed a higher transmittance of ~80% at ~5 µm, with the cut-off occurring at higher wavelengths. Note, however, that this sample was only ~0.5 mm thick.

3.4. Hardness measurements

Figure 8 shows Vickers microhardness data as a function of mean grain size for the SPS samples. Each data point is an average of at least 10 indents and all the indents were created by applying a 1 kg load.

As can be seen, the hardness values of the samples showed a small increase with decreasing grain size. The reason could be the presence of more grain boundaries in the samples with smaller grain size, restricting the free path of dislocations resulting in a
reduction in plastic deformation [29,30]. It may also be seen that the hardness value of sample SPS-500-12-SC, ~22 GPa, was the highest and matched the data in the literature for fully-densified alumina bodies prepared using sub-micrometer powders.

4. Conclusions

Although the use of hot pressing was not successful in the current work, samples made using SPS could be produced with densities up to 99.9% of theoretical. When as-synthesised powder was densified directly, excessive grain growth resulted and hence a large final grain size was observed; ~6.25 µm for sample SPS-70-14 and ~0.64 µm for sample SPS-500-12. The use of spray freeze dried granule powder as a more crushable alternative to as-synthesised powder, whether pre-pressed into a compact or not, failed to result in high enough densities, but the use of slip casting yielded green bodies of 53% density that could subsequently be densified to ~99.96% of theoretical density at 1200°C/500 MPa, whilst retaining a mean grain size of ~0.32 µm. These samples offered in-line transmittance values of up to ~80%; which approaches the transmission value of sapphire (86%). Microhardness values of 22 GPa were also obtained on reducing the grain size of sintered alumina by applying high pressure during the SPS process; this value is slightly improved in comparison with the values achieved in previous work on pure alumina. Attempts to increase the pressure to 1 GPa or drop the temperature to 1100°C at 500 MPa were unsuccessful, in both cases higher sintering temperatures were needed.

Acknowledgements

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the USA for funding. The authors would also acknowledge use of facilities within the Loughborough Materials Characterisation Centre (LMCC).
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Figure 7  (a) UV-Vis transmittance spectra of nanoalumina bodies sintered by SPS and HP [9,17,18,25-27] and (b) IR transmittance spectra; inset in both graphs are the equivalent transmittance spectra of some common ceramic materials [28]

Figure 8  Variation in hardness values with grain size in alumina compacts. Hardness values found in the literature were measured on fully-dense compacts prepared from sub-micrometer alumina powders [30,31].
<table>
<thead>
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<th>Sample</th>
<th>Type</th>
<th>T / °C</th>
<th>Time / min</th>
<th>Pressure / MPa</th>
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<td>Grain size / μm</td>
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<tr>
<td>SPS-500-12-DP</td>
<td>98.92 ± 0.1</td>
<td>0.66 ± 0.09</td>
<td></td>
<td></td>
</tr>
<tr>
<td>SPS-500-12-SC</td>
<td>99.96 ± 0.02</td>
<td>0.32 ± 0.07</td>
<td></td>
<td></td>
</tr>
<tr>
<td>SPS-500-11-SC</td>
<td>99.67 ± 0.08</td>
<td>0.25 ± 0.07</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>