Preparation and Characterization of Mullite-Alumina Structures Formed "In Situ" from Calcined Alumina and Different Grades of Synthetic Amorphous Silica

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1. Introduction

Mullite (3Al₂O₃·2SiO₂ or Al₆Si₂O₁₃), one of the main raw materials for the ceramic industry⁴,⁵,⁶, is an input for the production of refractory materials⁴,⁵,⁶, catalyst support⁷,⁸,¹⁰,¹¹, electronic and optic devices¹²,¹³, thermal insulators⁸,¹⁴, hot gas filters (above 600°C)¹⁶,¹⁷, and biological and biomedical scaffolds¹⁸. Its formation is a complex set of high-temperature (800-1400°C) solid state reactions¹²,¹³ based on the mutual diffusion of Al³⁺, Si⁴⁺ and O²⁻ ions amongst particles. Various studies have pointed out the characteristics of the silica source (crystallinity, particle size distribution, presence of impurities) and processing conditions (heating rate, temperature of the thermal treatment) affect directly the yield, kinetics and morphology of the products formed¹⁴. For instance, reactions between coarse alumina and micronized quartz particles (45 μm average size) generate porous mullite at temperatures above 1300°C, whereas the combination of finer nanoparticles of alumina and amorphous silica produced by alcoxii-based precursors reduced the range to 800-1100°C and promote greater densification⁵. Therefore, by changing the characteristics of the silica source, ceramics of different microstructures, porosity levels and thermomechanical properties can be designed.

Amongst the several raw materials available for mullite production, many recent studies have explored the use of synthetic amorphous silica (SAS) grades because they are high-purity particles (SiO₂-content above 97 %) of great homogeneity and reproducibility. During the first heating and above the glass transition temperature (typically in the 900-1100°C range), their viscous flow behaves as sintering aid enhancing particle densification. SASs particles can be produced in a wide range of characteristics, as shown elsewhere⁵. Its technological importance suggests that it is important to understand how these different physical properties could affect mullite formation. Therefore, in this study, four grades of SAS⁸,¹⁵,³⁸,³⁹ (Table 1, Fig. 1) were tested regarding their ability in generating dense structures of in situ mullite after a reaction with fine calcined alumina. They were chosen because they exhibit similar average particle sizes and purity and significant differences in other physical properties, such as a specific surface area and levels of intra-particle mesoporosity.

SASs promote early mullite formation (below 1300°C) and greater densification during sintering. This paper reports the "in situ" formation of mullite-alumina structures from fine calcined alumina (α-Al₂O₃) and four grades of SAS of different particles' morphology, specific surface area and internal porosity. After sintering assisted by dilatometry (up to 1500°C), the samples' total porosity, density and flexural strength were measured. The relative density and strength levels of some of these structures were greater than or comparable to other studies in which similar compositions were sintered at higher temperatures (1600-1700°C). Their microstructure assessment indicated that the specific surface area and internal porosity of SAS particles showed a major influence in the development of these physical properties.

Keywords: Synthetic amorphous silica, mullite, microstructure, physical properties, calcined alumina, densification.
Table 1. Characteristics of the raw materials tested.

<table>
<thead>
<tr>
<th>Characteristics</th>
<th>α-Al₂O₃ source</th>
<th>Precipitated silica (PS)</th>
<th>Synthetic amorphous SiO₂ sources</th>
<th>Microsilica (MS)</th>
<th>Silica from rice husk (RH)</th>
<th>Silica from rice husk ash (RA)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Composition (wt%)</td>
<td>Al₂O₃: 99.8; Na₂O: 0.05; Fe₂O₃: 0.01; CaO: 0.03; MgO: 0.07; SiO₂: 0.04</td>
<td>SiO₂: 97.8; Na₂O: 2.2</td>
<td>SiO₂: 94.9; Na₂O: 0.2; K₂O:0.2; SO₃: 0.2; Fe₂O₃: 0.1; CaO: 0.2; MgO: 0.2; Al₂O₃: 4.0</td>
<td>SiO₂: 99.9; Na₂O: 0.1</td>
<td>SiO₂: 99.9; Na₂O: 1.0</td>
<td></td>
</tr>
<tr>
<td>Solid density (ρ, g.cm⁻³)</td>
<td>3.9</td>
<td>2.1</td>
<td>2.4</td>
<td>2.20</td>
<td>2.1</td>
<td></td>
</tr>
<tr>
<td>Specific surface area (SSA, m².g⁻¹)</td>
<td>9.4</td>
<td>118</td>
<td>24</td>
<td>178</td>
<td>126</td>
<td></td>
</tr>
<tr>
<td>Total pore volume (TPV, cm³.g⁻¹)</td>
<td>0.02</td>
<td>0.4</td>
<td>0.05</td>
<td>0.2</td>
<td>0.4</td>
<td></td>
</tr>
<tr>
<td>Internal porosity (IP, %, Equation 3)</td>
<td>7.8</td>
<td>43.2</td>
<td>10.7</td>
<td>32.7</td>
<td>48.7</td>
<td></td>
</tr>
<tr>
<td>Average pore diameter (nm)</td>
<td>9.4</td>
<td>12</td>
<td>8.5</td>
<td>5</td>
<td>15</td>
<td></td>
</tr>
<tr>
<td>Primary particles average size (nm)</td>
<td>500</td>
<td>30-80</td>
<td>50-200</td>
<td>20-30</td>
<td>20-30</td>
<td></td>
</tr>
<tr>
<td>Clusters average size (μm)</td>
<td>Not applicable</td>
<td>1-6</td>
<td>Not applicable</td>
<td>1-8</td>
<td>1-5</td>
<td></td>
</tr>
</tbody>
</table>

a) A1000SG, Almatis, Germany; b) Tioxil T300, Solvay-Rhodia Group S/A, Brazil; c) Microsilica 971, Elkem, Norway; d) References 40,43; e) Reference 45.

Figure 1. SEM images of the synthetic amorphous silica (SAS) grades tested: a) silica precipitated from sodium silicate (PS), b) silica obtained from rice husks (RH), c) silica obtained from rice husk ashes (RA), d) microsilica (MS).

of intra-particle mesoporosity suggests that these silica grades can be extremely reactive in contact with calcined alumina at high temperatures. Microsilica (MS, Fig. 1d) particles are spherical, dense and individualized and their broad particle size distribution ranges from 50 to 500 nm. They are obtained as a precipitated by-product during the production of elemental silicon and are widely used in civil construction for enhancing the packing of high performance concrete, and in refractories for the production of mullite-based bricks and castables1,7,8,15,38,39,48,49.

After physical characterization, the SAS particles were combined with calcined alumina to produce in situ mullite dense parts. Their sintering behavior was assisted by dilatometry and the products formed were identified by X-ray diffraction and scanning electron microscopy. The physical properties (solid and relative density, total porosity, flexural strength) of the resulting structures were related to the characteristics of the amorphous silica particles.

2. Materials and Methods

2.1. Synthesis and characterization of the particles

The following raw materials (Table 1) were used: a) calcined alumina (CA, A1000SG, Almatis, Germany), b) silica from the precipitation of sodium silicate (PS, Tixosil T300, Solvay-Rhodia Group S/A, Brazil), c) silica extracted from rice husk (RH43), d) silica extracted from rice husk ash (RA43), e) silica obtained by the precipitation of oxidized silicon vapor (MS, Microsilica 971, Elkem, Norway), f) organic binder (PVB; Poly(vinyl butyral), Butivar 98, Sigma-Aldrich, USA) and g) isopropyl alcohol (Synth, Brazil).

The RH particles40,43 were produced initially washing dried rice husks with a 10 wt% citric acid solution (C₆H₈O₇; Synth, Brazil) in autoclave (Phoenix Luferco, AV 100 model,
Brazil) for 1 h at 127°C ± 5°C and 150 kPa ± 20 kPa. After neutralization with distilled water and drying (100°C ± 5°C for 12 h), the husks were calcined at 700°C for 2 hours at 15°C/min heating rate. The calcined material was dry-milled for 2 hours (ball-mill operating at 60 rpm with ZrO$_2$ spheres of 5 mm diameter; spheres ratio of 10:1 in relation to the dried material).

To obtain RA particles, a suspension containing 0.1 wt% of rice husk ash in a 2 Mol.L$^{-1}$ sodium hydroxide (NaOH) solution was heated under reflux conditions for 2 h. The sodium silicate solution formed was filtered to remove the non-reacted ashes, and hydrochloric acid (HCl, 2 mol.L$^{-1}$) was added until pH = 3 for the formation of a gel of silica particles, according to Equation 1:

$$Na_2SiO_3 + 2HCl \rightarrow SiO_2(\text{Precipitated}) + 2NaCl(\text{Dissolved}) \ (1)$$

RA particles were attained after intensive washing with distilled water for the removal of sodium chloride salt, filtering and drying (2 h at 110°C).

Particles samples were previously degassed at 200°C for 3 h in a vacuum atmosphere and then characterized regarding the specific surface area (SSA, m$^2$.g$^{-1}$, BET method, ASTM C1069-09 standard "Standard Test Method for Specific Surface Area of Alumina or Quartz by Nitrogen Adsorption"), total pore volume (TPV, cm$^3$.g$^{-1}$, for pores thinner than 50 nm) and average pore diameter (PD), by adsorption-desorption isotherms of high-purity N$_2$ (Nova 1200e, Quantachrome Instruments, USA; the ratio between relative pressure, $P$, and atmospheric pressure, $P_0$, ranged from 0.05 up to 1). Their density ($\rho$, g.cm$^{-3}$) was measured by the Helium picnometer method (Ultrapyc 1200e, Quantachrome Instruments, USA).

The particles’ internal porosity (IP, %) is the ratio between the total pore volume inside the particles (TPV) and the total volume of the particles (which corresponds to the sum of TPV and solid volume)$^{26}$. The IP was calculated by Equation 2:

$$IP(\%) = 100\% \times \left[\frac{(\rho \times TPV)}{(\rho \times TPV + 1)}\right] \ (2)$$

2.2. Preparation and characterization of mullite samples

The stoichiometric composition of mullite (3Al$_2$O$_3$.2SiO$_2$)$^{22-24}$ was prepared combining the different silica grades (PS, MS, RH and RA) and calcined alumina (CA) (Fig. 2a-b). The deagglomeration and mixing processes of alumina and silica particles were performed in a ball-mill with a 2 wt% of PVB isopropyl alcohol suspension containing 33% solids (grinding specification: 5 mm diameter zirconia spheres, mass ratio of sphere/powder: 1:3, 2 liters polyethylene flask, 60 rpm for 2 hours; Fig. 2c). The isopropyl alcohol was completely evaporated (Fig. 2d) and the resulting powder was shaped by uniaxial compression (40 MPa, Fig. 2e). The sintering of green samples (8 mm length × 6 mm diameter cylinders) was monitored by a 402C dilatometer (NETZSCH, Germany), under a 5°C/min heating rate up to 1500°C.

Samples for porosity and mechanical property measurements (bars of 6 × 20 × 70 mm) were heat-treated at 1100°C and 1500°C (heating rate of 5°C/min, threshold of 2 hours at 400°C and 3 h at the final temperature, 10°C.min$^{-1}$ cooling rate) (Fig. 2f-g). Their relative density (RD, %) and the total porosity (TP, %)$^{15,50}$ were calculated using Equations 3 and 4:

$$RD(\%) = 100\% \times \left[\frac{M}{L \times W \times T \times \rho}\right] \ (3)$$

$$TP(\%) = 100 - RD \ (4)$$

M, L, W and T are, respectively, the samples' mass (g) and average length (cm), width (cm) and thickness (cm). The density of these materials ($\rho$) was measured by the Helium picnometer method in equivalent samples after milling ($D_{\text{Part}} \leq 100$ µm).

$$Na_2SiO_3 + 2HCl \rightarrow SiO_2(\text{Precipitated}) + 2NaCl(\text{Dissolved}) \ (1)$$

A universal testing machine (WDW-30E, China, 0.5 mm.min$^{-1}$ load application rate) and Equation 5 were used to determine their flexural strength (FS, MPa) by the Three-Point Bending Test (ASTM C78: "Standard Test Method for Flexural Strength of Concrete Using Simple Beam with Third-Point Loading"):

$$FS (MPa) = \frac{(3 \times F \times S)}{(2 \times W \times T^2)} \ (5)$$
F is the maximum applied load (N), S is the distance between the two supporting points (m), and W (m) and T (m) are the average width and thickness of the test sample, respectively. In all cases, five samples were tested for each condition.

The results attained were compared with other similar systems using relative flexural strength (RFS, %) versus relative density (RD, %) curves. RFS is the ratio between the experimental results of FS and the theoretical value obtained for a 100% dense sample (for mullite 2,4,16,30,34,51, the theoretical FS = 180 MPa). Such an approach is useful for investigating the mechanical behavior of porous (RD ≤ 50 %) and semi-porous structures (95% ≤ RD > 50 %). It is based on the fact that, ideally, a reduction in the material’s relative density (caused by the introduction of new internal surfaces, as pores) would be followed by a proportional reduction of its mechanical properties. Such a behavior is represented by the y = x function. In real cases, however, most commonly, the reduction of the mechanical properties caused by the introduction of porosity is far more intense due to the presence of tensile concentrators and lack of efficient sintering amongst particles.

The phases formed during thermal treatment were identified by X-ray diffraction (Rigaku, Rotaflex RV 200B, Japan, pulverized samples). The microstructure of the samples was assessed at fracture surfaces by field emission gun scanning electron microscopy (SEM, FEI, Inspect F50 model, Netherlands).

3. Results and Discussion

3.1 Samples sintered at 1100°C

Samples containing precipitated silica (M-PS₁₁₀₀), silica from rice husk (M-RH₁₁₀₀) and silica from rice husk ash (M-RA₁₁₀₀) showed lower total porosity and flexural strength levels (Fig. 3a) and earlier densification (Fig. 4a-c) than the silica-free reference sample (CA₁₁₀₀). Such behavior can be explained analyzing the same flexural strength results as a function of silica particles’ internal porosity (IP) (Fig. 3b and Table 1). The particles of these three SAS grades are comprised of nanoclusters strongly aggregated containing large quantity of mesopores amongst them (Fig. 1). These pores are smaller than 30 nm and are extremely unstable under heating. Due to this, even mild temperatures, such as 400-800°C can induce their collapse and early densification of the silica particles. The overall effect upon the samples’ structure is an intense macroscopic shrinkage and porosity reduction, not necessarily followed by gains of strength.

For microsilica-containing samples (M-MS₁₁₀₀), on the other hand, a different behavior was observed. Comparing to the other SAS grades, MS particles have a much lower internal mesoporosity (IP = 10.7 %) and specific surface area. Due to this, its reduces the amount of voids amongst particles at the compacting step. During the initial heating, no significant densification was observed up to 1000°C (Fig. 4d). After sintering at 1100°C, M-MS samples exhibited the lowest level of porosity and the highest flexural strength amongst the silica-containing-ones. It is also possible that the impurities of the microsilica have generated some small portion of low melting point the liquid phase. They were not detected after cooling in this study but such effect has been already described elsewhere.

No traces of silica crystallization or mullite formation were detected by XRD (Fig. 6) in the samples sintered at 1100°C. Their fracture surfaces (Fig. 5a-e) are comprised of porous granular packs of individualized particles containing very low necking points. These two aspects indicate that only physical rearrangements, such as pore coalescence or viscous flow, occurred up to 1100°C and that the physical properties of the structures were defined mainly by the calcined alumina matrix behavior.

3.2 Samples sintered at 1500°C

All silica-containing samples sintered at 1500°C showed porosity reduction and gains of strength in comparison to those...
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The densification rate of the M-MS sample was significantly reduced between 1300-1430°C (Fig. 4d). This behavior was also observed in other studies\(^8,9,15\) and can be related to the mullite generation reaction. In comparison to the calcined alumina matrix (\(\rho = 3.8-4\) g.cm\(^{-3}\)), the density of mullite is lower (\(\rho = 3.2\) g.cm\(^{-3}\)), therefore, its formation can be followed by a volumetric expansion, depending on the compacting level of the structure. In the present case, since MS improved the initial structure packing efficiency, the formation of mullite partially hindered the particles' densification. However, due to the presence of several alkali-based impurities in the MS composition, it is reasonable to assume the high flexural strength attained is also related to a certain level of liquid sintering.

The comparison between the RFS values of this study to those in other reports (Fig. 7) has shown two important aspects. Firstly, for the range of relative density of 45-99 % tested, the RFS showed a linear variation, which indicates it is possible to tailor such property level depending on the application. For instance, compositions of higher RFS and low porosity, such as M-RA\(_{1500}\) and M-MS\(_{1500}\), could be used to produce refractory bricks or abrasive tools. On the other hand, those of lower RD (M-RH\(_{1100}\) and M-PS\(_{1100}\)) could be used as porous structures for thermal insulation or as biological scaffolds. Secondly, the overall level of the RFS values obtained in this work is close to those described in the literature for samples of similar compositions, but produced by more complex compaction techniques (isostatic pressing\(^3,10,12\)), longer sintering times (longer than 3 h\(^{16,17}\)) and

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**Figure 4.** Thermal linear variation for samples containing different synthetic amorphous silica (SAS) grades: a) M-PS, b) M-RH, c) M-RA, d) M-MS

thermally treated at 1100°C (Fig. 3a). Their microstructure (Fig. 5f-j) evolved to a solid block comprised of mullite (major constituent) and traces of un-reacted corindon and cristobalite (Fig. 6), where evidence of viscous flow can be seen. The dilatometric analysis of the M-PS\(_{1500}\), M-RH\(_{1500}\) and M-RA\(_{1500}\) samples (Fig. 4a-c) revealed total shrinkage levels similar or greater than the one observed for the silica-free reference (CA\(_{1500}\)). For such compositions, the larger the IP of the silica particles, the more intense the reduction in the structures’ total porosity and enhancement of their strength (Fig. 3b).
Figure 5. Fracture surfaces of samples treated at 1100°C and 1500°C for 3 h (SEM observation)

higher thermal treatment temperatures (above 1500°C\cite{28,29}) or with the aid of liquid phase sintering\cite{27,32}. Therefore, for similar levels of flexural strength, a significant energy saving in the processing operations could be achieved by using SASs.

Figure 6. X-ray diffraction patterns for samples treated at 1100°C and 1500°C for 3 h: a) M-PS, b) M-RH, c) M-RA, d) M-MS. Crystalline phases (JCPDS file): C = corundum (α-Al₂O₃, 42-1468), S = cristobalite (SiO₂, 39-1425), M = mullite (Al₆Si₄O₁₃, 15-0776)

Figure 7. Relative flexural strength (RFS) versus relative density (RD) for samples tested (other results from the literature are also shown on the same basis)
4. Conclusions

The internal porosity and specific surface area of the synthetic amorphous silica (SAS) particles tested significantly impacted the development of microstructure and physical properties of the alumina-mullite structures attained. Two opposite effects were observed.

Firstly, samples containing SAS grades of high internal porosity (silica precipitated from sodium silicate, silica obtained from rice husk, and silica obtained from rice husk ashes) experienced a significant shrinkage between 400-800°C. At this temperature range, the mesopores located inside the particles of silica collapsed increasing their density and the overall packing efficiency of the system. As a consequence, after sintering at 1500°C, structures of very low total porosity and high flexural strength were attained. The comparison with other reports in literature showed that the strength levels attained were similar to other systems sintered at higher temperatures for longer periods. This aspect is technologically important because these compositions could replace conventional materials and produce a significant energy saving during the thermal treatment. Secondly, for samples containing the silica grade of very low internal porosity and specific surface area (microsilica), the spherical shape of such particles improved packing efficiency during the pressing step. Due to this, comparing to the other SAS grades, the microsilica-containing structures presented lower porosity and higher strength after sintering at 1100°C. Besides this, such samples also showed lower densification rate and total linear shrinkage. This effect was attributed to the formation of mullite, which is can be an expansive reaction, when it occurs in a dense structure.

After sintering at 1500°C, all the samples exhibited a dense microstructure containing signals of viscous flow and were comprised mainly of mullite and small portions of un-reacted corindon (α-Al₂O₃). It is possible that the crystallization of the SAS particles may have hindered the total conversion of the reactants in mullite or would require longer thermal treatment periods. These aspects are currently under investigation in authors' research group.

5. Acknowledgements

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