Vitrification and sinter-crystallization of fly ash with glass cullet

Abstract
The synthesis of a new glass-ceramic obtained by sinter-crystallization has been investigated by using soda-lime-silicate glass waste and fly ashes from a coal power thermal station located in Andorra (Teruel, Spain). An original glass as frit with composition of 50wt% recycled soda-lime glass, 25wt% bottom ash, 15wt% fly ash and 10wt% CaCO₃ has been melted. After sinter-crystallization at 850°C, it has precipitated two main crystalline phases: sodium anorthite and the hedenbergite. The linear shrinkage is 1/3 of the type of conventional porcelanized stoneware and the water absorption of this glass-ceramic is similar to a conventional porcelanized stoneware tile (less than 1wt%), being the apparent density slightly higher than this type of tiles (2.6g/cm³ instead of 2.4g/cm³). Flexural strength is near twice than porcelanized stoneware (around 950kg/cm² instead of 550kg/cm²) (95MPa in the new glass-ceramic with respect to 55MPa for the above mentioned as reference material).

Keywords: recycling, industrial wastes, glass-ceramics, sinter-crystallization; fly ashes

Abbreviations: XRF, X-ray fluorescence; TTT, time-temperature-transformation diagram; EDS, energy dispersive X-ray spectrometer; SEM, scanning electron microscopy

Introduction
Nowadays, in this second decade of this century, the industrial residues they continue generating environmental problems. Such of these waste (glass cullet and bottom or fly ashes from the coal power stations) are still abundant and not definitive applications or solutions for its immobilizing have been given, though a considerable amount of research has been conducted in the last years. One of the promising applications proposed by laboratories research has been their potential for being used in the construction industry. With respect the glass cullet from the conventional glass there is abundant research carried out in the last decades of 20th and beginning of this 21st century. Vitrification process has been demonstrated is an adequate processing method for inertize toxic and abundant residues and even to facilitate their recycling as secondary raw materials in ceramics and glasses industries. Transforming of starting glasses after vitrification into glass-ceramics by controlled thermal treatment is possible to reach immobilizing of a wide range of industrial wastes (mineral residues, sludges from dumps, slags, ashes...). Besides, the low cost and great availability of waste make these glass-ceramics materials very attractive from an economical and technological point of view, so synthetic high-performance materials with broad applications in construction and civil engineering can be obtained from residues. Therefore, it has been the aim of this research to explore the synthesis of a new type of glass-ceramic by the sinter-crystallization process from soda-lime-silicate glass and several ashes from a coal power thermal station located in Andorra (Teruel, Spain).

Materials and methods
The batch original composition for melting of an “mother or original” glass able for being transformed in a glass-ceramic was: 25 wt% bottom ash (from Andorra), 15wt% fly ash (from Andorra), 10wt% calcium carbonate (industrial CaCO₃) and 50wt% of glass cullet (from the recycling glass sector). The chemical composition determined by XRF analysis of wastes and raw materials is shown in Table 1. X-ray fluorescence (XRF) dispersion wavelength equipment was the model S4 Pioneer – Bruker. The same Table 1 includes the XRF analysis of the final glass-ceramic. This mixture has been melted at 1500ºC during 1 hour in a lift furnace by using a refractory crucible and then, the glass composition has been quenched in cold water to obtain glassy granules, ready to be wetly milled in an alumina ball mill under 45μm. The dried glassy powder (24h at 110ºC) has been moistened at about 10 wt% to be pressed with a Nanneti® uniaxial press up to a pressure of 30MPa in the shape of rectangular glassy pieces. The pieces were subjected to the thermal firing cycle in a muffle kiln (Nannetti®), which has been achieved by varying the maximum temperature (700°C-1000°C) and the residence period (30min/10h) in order to elaborate the corresponding Time-Temperature-Transformation diagram (TTT). The different heat treated samples at successive temperatures and time for drawing the TTT diagram was characterized by XRD and SEM/EDS. Crystalline phases were identified X-ray diffraction (XRD) and by using a Bruker-AXS D4 Endeavor equipment (using Ni-filtered Cu-Kα radiation with scanning speed of 2°/ (2θ) per minute and registering the diffraction pattern in the 10°-80° Bragg angle range. The microstructure was observed by scanning electron microscopy (SEM) JEOL 7001F with energy dispersive X-ray spectrometer (EDS) operating in the 15-20kV interval.
The physical and technological properties, such as apparent density (hydrostatic balance method), flexural strength (determined by a HOYTOM® plasticinometer under load cell of 5000 N and 16N initial force), linear shrinkage (Mitutoyo® digital caliper) were determined in the original glass and glass-ceramics. Linear shrinkage was determined, as is usual by considering the initial length (Li), the length of the dried piece and the final length (Lf) of the heat treated sample: %\textit{Sh} = (L_i - L_f)/L_i \times 100. Water absorption has been determined according to the Quality Normative for Ceramic Tiles (UNE-EN-ISO 10545-3).

**Results**

**Chemical composition**

The final composition of original glass and corresponding glass-ceramic was: SiO₂ (52.75wt%), Al₂O₃ (17.05wt%), CaO (12.34wt%), Fe₂O₃ (7.69wt%) and Na₂O (6.83wt%).

**Experimental time-temperature-transformation diagram (TTT diagram)**

The experimental TTT diagram which has been represented in the Figure1 shows the “nose curve” dividing the three main zones: a) the amorphous area outside the curve, where the glass stay vitreous without depicting diffraction peaks; b) the zone inside the nose curve where the crystallites nucleate and growth and c) the superior outside nose curve, where takes place the softening of glass and melting. Besides, X-ray diffractograms corresponding to samples fired for 30min are displayed in Figure1B.

**Microstructural morphology**

The microstructural morphology of heat treated samples during 30 minutes has been observed by SEM (Figure 2).

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**Table 1** Semiquantitative analysis (XRF) of industrial wastes and the glass-ceramic

| wt%    | Na₂O | MgO | Al₂O₃ | SiO₂ | P₂O₅ | K₂O | CaO | TiO₂ | Fe₂O₃ | LOI* |
|--------|------|-----|-------|------|------|-----|-----|------|-------|------|
| Wastes |      |     |       |      |      |     |     |      |       |      |
| Glass  | 12.59| 3.75| 0.85  | 73.16| 0.01 | 0.30| 8.94| 0.05 | 0.10  | 0.10 |
| Slag   | 1.13 | 23.85| 43.19 | 0.31 | 1.13 | 1.13| 5.22| 0.72 | 23.85 | 0.20 |
| Fly ash| 0.20 | 1.23 | 26.63 | 44.44| 0.41 | 1.23| 5.53| 0.92 | 18.43 | 0.15 |
| CaCO₃  | 0.00 | 0.00 | 0.00  | 0.00 | 0.00 | 0.00| 55.98| 0.00 | 0.00  | 44.01|
| Glass-ceramic | 6.83 | 2.10 | 17.05 | 52.75| 0.78 | 12.34| 3.36 | 7.69  | 0.03  |

*LOI, loss on ignition

![Figure 1](image_url) **Figure 1** A) TTT Diagram of the composition. B) X-ray diffractograms from samples treated during 30 minutes at successive temperatures from 800 to 1000°C.
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Figure 2 Microstructure observed by SEM from different samples heat treated during 30 minutes at A) 800°C, B) 850°C, C) 900°C, D) 950°C, E) 975°C, F) 1000°C.

Ceramic properties

The variations on apparent density and flexural strength vs temperature are displayed in Figure 3a, while the linear shrinkage and water absorption vs temperature are plotted in Figure 3B. These parameters are essentials to control the quality of the final product, because it has a ceramic application, being susceptible to cover floors and walls.

Discussion

About the chemical composition, it is expected the devitrification of crystalline phases belonging to the more common simplified CaO-Al<sub>2</sub>O<sub>3</sub>-SiO<sub>2</sub> ternary system, in which the fly ashes are usually located and mineral substitutions due to transition element oxides. Referring to the TTT diagram, the crystallization occurs from 850°C and is independent of time of heat treatment (Figure 1A). Besides, 850°C is a relatively low temperature that together with the short time of operation, become an economically profitable industrial process. The X-ray diffractograms of samples treated at 800-1000°C during 30 minutes of residence time are shown in Figure 1B. The sample tested at 800°C is completely amorphous, whereas the samples subjected to temperatures from 850°C to 950°C show the most intense peaks of the solid solution of sodium anorthite (JCPDS card No. 86-1650), a mineral phase of anorthite where some CaO has been substituted by Na<sub>2</sub>O, which is present in the glass-ceramic (Table 1) in an approximate content of 7wt% and hedenbergite (JCPDS card No. 70-1876), a diopside-pyroxenic phase with iron substitutions. While this sodium-anorthite presents its maximum intensity at 900°C, the hedenbergite shows the maximum crystallinity at 950°C. Both phases exhibit competitive mechanical properties.
About microstructural morphology, at 800°C, the corresponding SEM micrograph shows the presence of glassy particles partly sintered and without signs of crystallization, while at 850°C the vitreous particles show heterogeneities of crystals nuclei. Since 900°C to 1000°C micrographs exhibit the presence of crystals inside the sintered particles. The average chemical analysis by EDX of the crystals in these crystallized samples shows the following oxides: 6.0 wt% Na₂O, 0.9 wt% MgO, 15.5 wt% Al₂O₃, 61.2 wt% SiO₂, 0.8 wt% K₂O, 8.8 wt% CaO, 6.7 wt% Fe₂O₃, showing that there is a mixture of both phases, the sodium anorthite, and the hedenbergite. Figure 3A shows density values with a maximum value (2.60 g/cm³) is reached 850°C, been more or less constant until 1000°C, in samples heat treated at 30 minutes. This value is higher and is obtained at lower temperatures than in the case of porcelain ceramic tiles, which reaches an only apparent bulk density of 2.40g/cm³ at 1200°C. The flexural strength increases with temperature, achieving the maximum (around 95MPa) at 950°C when the hedenbergite shows the higher value. At higher thermal treatment temperature (1200°C), the resulting tile shows a conventional value of 60MPa. The linear shrinkage (Figure 3B) shows the same behavior than the apparent density and reaching the maximum (around 2.5%) at 850°C, remaining almost constant until 1000°C. In contrast, the water absorption (Figure 3B) exhibits a large reduction at 850°C, where the maximum linear shrinkage is obtained, holding at similar values of 1% until 975°C. After 975°C continues its reduction from 1% to 0.5 %, because the vitreous phase is growing whilst crystals are dissolving, a phenomenon which gives rise to the porous sealing, as can be seen in the SEM micrographs (Figure 2F). Compared to a conventional porcelainized stoneware tile, this glass-ceramic material exhibits a much lower shrinkage (2.5% with respect the 7.5% of porcelain tile) and similar water absorption at maximum apparent density (in both materials less 1%). Therefore, a conclusion it is evident the advantages of using glass cullet for obtaining a new type of glass-ceramic tiles.

![Figure 3: Variation of the glass-ceramic technological properties heat treated during 30 minutes: A) apparent density and flexural strength vs temperature and B) linear shrinkage and water absorption vs temperature.](image-url)
Conclusion

It has been obtained a high-resistance glass-ceramic tile formulated from industrial residues (50% glass cullet, 25% bottom ash, 15% fly ash and 10% CaCO₃), which exhibits better technological properties (2.60 g/cm³ apparent density and 95 MPa flexural strength) than conventional porcelainized stoneware tiles.

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Conflicts of interest

The authors declare no conflict of interest.

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