Densification of Bi$_{0.5}$Na$_{0.5}$ZrO$_3$ ceramic using liquid-phase sintering method

Panupong Jaiban$^a$, Sukanda Jiansirisomboon$^{a,b}$, Anucha Watcharapasorn$^{a,b,*}$

$^a$ Department of Physics and Materials Science, Faculty of Science, Chiang Mai University, Chiang Mai 50200, Thailand
$^b$ Materials Science Research Centre, Faculty of Science, Chiang Mai University, Chiang Mai 50200, Thailand

*Corresponding author, e-mail: anucha@stanfordalumni.org

ABSTRACT: Lead-free bismuth sodium zirconate (BNZ) ceramics with formula Na$_{0.5}$Bi$_x$ZrO$_{3-x}$/2Bi$_2$O$_3$ with $x$ = 0, 2, 3, 4, and 6 wt% were prepared by liquid-phase sintering method. The specimens were sintered at 850 and 900°C. Phase identification was investigated using X-ray diffraction technique. BNZ/4 wt% Bi$_2$O$_3$ and BNZ/6 wt% Bi$_2$O$_3$ ceramics sintered at 900°C showed impurity phase of Bi$_{3.62}$Zr$_{0.62}$O$_{12.31}$ compound due to excess additive reacted with zirconium in system. Scanning electron microscopy and energy-dispersive X-ray spectroscopy were employed to study microstructure and measure chemical composition of ceramics, respectively. The results revealed creation of bismuth oxide liquid phase at BNZ grain boundaries inhibited grain growth and decreased pore size. This caused the relative densities of the modified samples to increase.

KEYWORDS: lead-free material, bismuth sodium zirconate, sintering aid, microstructure

INTRODUCTION

Sintering process is important for fabrication of ceramic materials. One of the well-known processes is liquid-phase sintering. Usually, the purpose of the mentioned method is to enhance densification rate and to decrease the fabrication temperature of a system requiring too high temperature in a conventional solid-state sintering. The microstructures of ceramics produced by liquid-phase sintering consist of two phases: (1) the crystalline grains and (2) the grain boundary phase resulting from the solidified liquid. Nowadays, the use of liquid-phase sintering method is increasing in ceramic components such as 96% alumina substrates for computer packages, bearing, silicon nitride extrusion dyes, etc. Moreover, it is also used in fabrication of electronic devices. Examples are actuators, sensors, transducers, etc.

Recently, Bi$_{0.5}$Na$_{0.5}$ZrO$_3$ ceramic (BNZ) was successfully fabricated$^{3,4}$. The novel material promotes interesting dielectric properties including the diffuse phase transition at high temperature ($T_c$ ≈ 425°C). A material having a high Curie temperature is presumed to be a promising candidate for various electronic devices such as multilayer capacitors, detectors, MEMs, sensors, actuators etc. On the other hand, its dielectric constant at room temperature was rather low (300; 0.1 kHz) but the electrical conductivity was quite high (9 × 10$^{-6}$ S/m; 0.1 kHz). Similarly, in firing process, the green body seemed to require a high sintering temperature of 1100°C. As it does not conserve energy, it is considered too expensive.

To resolve the mentioned problem, several research attempted to apply liquid-phase method in order to enhance microstructure evolution and reduce sintering temperature of many systems i.e., bismuth sodium titanate (Bi$_{0.5}$Na$_{0.5}$TiO$_3$)$^3$ and lead zirconate titanate Pb(Zr,Ti)O$_3$$^6,7$ ceramics that are well known and have been studied extensively$^8$. Because it provides high Curie temperature and has possible applications, Bi$_{0.5}$Na$_{0.5}$ZrO$_3$ ceramic is considered to be interesting for our studying. Meanwhile, investigation which involves using liquid-phase sintering technique with Bi$_{0.5}$Na$_{0.5}$ZrO$_3$ ceramic has not been reported. Therefore, the aim of this work is to study densification of Bi$_{0.5}$Na$_{0.5}$ZrO$_3$ ceramic at low firing temperature (850–900°C) by employing liquid-phase sintering process with Bi$_2$O$_3$ since it has low melting point and is known to be one of the widely used sintering aids for improvement of ceramic microstructure$^9,10$.

MATERIALS AND METHODS

The specimen was fabricated according to the formula (Bi$_{0.5}$Na$_{0.5}$)ZrO$_3$/10 wt% Na$_2$CO$_3$. Na$_2$CO$_3$ content
was used to help complete calcination reaction. The powders were prepared by a conventional mixed-oxide method. The starting materials used in this study were ZrO$_2$ (99%, Riedel-de Haën), Bi$_2$O$_3$ (98%, Fluka) and Na$_2$CO$_3$ (99.5%, RdH). The mixtures of oxides were ball milled in ethanol for 24 h. The mixed powder was dried at 150 °C for 24 h and calcined in a closed alumina crucible at a temperature of 800 °C for 2 h. Then, the calcined powder was ball milled again for 6 h and was calcined again at the same temperature and time.

After obtaining BNZ powder, BNZ/$x$Bi$_2$O$_3$ mixture with $x = 0, 2, 3, 4$, and 6 wt% were prepared by ball milling for 5 h. All powders were dried at 150 °C for 24 h. After sieving, a few drops of 3 wt% PVA (polyvinyl alcohol) was added to the mixed powders as a binder before being pressed into pellets with a diameter of 10 mm using a uniaxial press with 1.0 ton weight for 15 s. Binder removal was carried out by heating the pellets to 500 °C for 1 h. These pellets were subsequently sintered at 850 and 900 °C for 2 h dwell time under its own atmosphere in a closed alumina crucible.

All samples were prepared for X-ray diffraction analysis. The sintered pellets were polished by abrasive paper and cleaned by ultrasonic technique. Phase identification of BNZ and BNZ/Bi$_2$O$_3$ ceramics were investigated using an X-ray diffractometer (XRD, Phillip Model X-pert) in a 2θ range of 20–90°. For microstructure observation, sintered samples were polished employing abrasive paper and alumina particle. After that, all of specimens were etched at 800 °C for 15 min. Scanning electron microscopy (SEM) and energy-dispersive X-ray spectroscopy were used to observe microstruture and measure chemical composition, respectively. Average sizes of grain and pore were measured using lineal analysis. The relative densities of all samples were calculated using their measured bulk densities determined by Archimedes' method (AD-1653) and their theoretical densities from simulated X-ray diffraction pattern.

**RESULTS**

X-ray patterns of BNZ and BNZ/Bi$_2$O$_3$ ceramics sintered at 850 and 900 °C compared with BNZ powder are shown in Fig. 1a and b, respectively. **Powder Cell** software was used to simulate X-ray pattern resembling the pattern of BNZ powder obtained from experiment. The analysis resulting from the program revealed that the material possessed orthorhombic perovskite structure. This was in agreement with the previous report. The space group of BNZ was Pnma and its lattice parameters were $a = 5.7742$ Å, $b = 8.1443$ Å, and $c = 5.7037$ Å. The unit cell volume and theoretical density were 266.833 Å$^3$ and 6.353 g/cm$^3$, respectively. In addition, non-perovskite phase of Bi$_{7.38}$Zr$_{0.62}$O$_{12.31}$ compound (PDF no. 43-0445) was also found with BNZ/4 wt% Bi$_2$O$_3$ and BNZ/6 wt% Bi$_2$O$_3$ materials sintered at 900 °C.

**Fig. 1** X-ray diffraction patterns of BNZ powder and BNZ/Bi$_2$O$_3$ ceramics (a) sintered at 850 °C and (b) sintered at 900 °C.

Microstructures of BNZ/$x$Bi$_2$O$_3$ ceramics sintered at 850 °C where $x = 2, 3, 4$, and 6 wt%, respectively. It indicated that all consisted of crystalline grains and pores having various sizes. Average grain and pore sizes are given in Table 1. Both values tended to decrease when Bi$_2$O$_3$ concentration was increased. The presence of Bi$_2$O$_3$-based liquid phase at grain boundary area of modified material could be seen apparently in BNZ system containing maximum additive content (BNZ/6 wt% Bi$_2$O$_3$).
Table 1  Grain and pore size of BNZ/Bi$_2$O$_3$ ceramics sintered at different temperatures.

<table>
<thead>
<tr>
<th>Ceramics</th>
<th>sintered at 850 °C</th>
<th>sintered at 900 °C</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Grain size (µm)</td>
<td>Pore size (µm)</td>
</tr>
<tr>
<td>BNZ</td>
<td>4.77 ± 0.96</td>
<td>1.58 ± 0.94</td>
</tr>
<tr>
<td>BNZ/2Bi$_2$O$_3$</td>
<td>4.69 ± 0.69</td>
<td>0.85 ± 0.19</td>
</tr>
<tr>
<td>BNZ/3Bi$_2$O$_3$</td>
<td>3.59 ± 0.50</td>
<td>0.48 ± 0.28</td>
</tr>
<tr>
<td>BNZ/4Bi$_2$O$_3$</td>
<td>3.52 ± 0.38</td>
<td>0.47 ± 0.09</td>
</tr>
<tr>
<td>BNZ/6Bi$_2$O$_3$</td>
<td>3.47 ± 0.49</td>
<td>0.47 ± 0.20</td>
</tr>
</tbody>
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Fig. 2  SEM micrographs of BNZ/xBi$_2$O$_3$ ceramics sintered 850 °C where $x = (a)$ 0, (b) 2, (c) 3, (d) 4, and (e) 6 wt%.

Fig. 3  SEM micrographs of BNZ/xBi$_2$O$_3$ ceramics sintered 900 °C where $x = (a)$ 0, (b) 2, (c) 3, (d) 4 and (e) 6 wt%.

Demonstration of Bi$_2$O$_3$-based liquid phase existing at BNZ/6 wt% Bi$_2$O$_3$ ceramics sintered at 850 and 900 °C was carried out using energy-dispersive X-ray spectroscopy. The chemical composition results of selected areas between crystalline grain and grain boundaries inserted by liquid phase of both specimens are given in Table 2 in unit of atomic percentage. It revealed that all areas are composed mainly of bismuth, sodium, zirconium, and oxygen elements. Nevertheless, it could be noticed that the Bi amount...
Relative density values of BNZ and BNZ added Bi$_2$O$_3$ ceramics sintered at 850 °C and 900 °C are shown in Fig. 4. With increasing Bi$_2$O$_3$ content, the relative density value was maximized close to 90% with BNZ/6 wt% Bi$_2$O$_3$. This increasing trend was not different from BNZ and BNZ/Bi$_2$O$_3$ ceramics sintered at 850 °C.

**DISCUSSION**

According to the X-ray diffraction results of pure BNZ ceramics sintered at temperature of 850 and 900 °C, it could be seen that the patterns were in agreement with its calcined powder. Sintering process at both temperatures indicated that it could produce phase pure Bi$_{0.6}$Na$_{0.4}$ZrO$_3$ material and did not cause stoichiometry deviation during fabrication. In case of the Bi$_2$O$_3$-added systems sintered at 850 and 900 °C, all samples did not possess XRD diffraction pattern characteristics that were different from those of un-doped BNZ specimens sintered at the same temperature. Also, bismuth oxide phase was not detected. It was assumed that XRD technique was not sensitive enough to confirm a presence of this additive phase in BNZ system although adding concentration was more than 5 wt%. On the other hand, a weak impurity peak of Bi$_{13.8}$Zr$_{0.62}$O$_{12.31}$ compound existing approximately at 2θ = 28° was found with BNZ/xBi$_2$O$_3$ ceramics sintered at 900 °C where x = 4 and 6 wt%. Excess Bi$_2$O$_3$ content was presumed to react with ZrO$_2$ in the system at high temperature to form mentioned phase. The result was similar to a study by Sood et al., who reported that this non-perovskite phase occurred at temperature above 850 °C and increased with increasing of Bi$_2$O$_3$. Hence in samples sintered at 850 °C, this second phase was therefore not found in Bi$_2$O$_3$-doped BNZ ceramics.

With respect to the microstructural evidence of pure BNZ ceramic sintered at 850 and 900 °C, it could be seen that both materials contained large non-uniform crystalline grains in a size range of 4.77–4.81 µm and many open pores with size up to 1 µm. Creation of large BNZ grains was expected which was attributed to a rapid migration of the boundaries during final stage of conventional solid-state firing. The mentioned behaviour also led to coalescence of the pores causing the average pore size to increase. Influences of rapid grain growth and pore coalescence on microstructure evolution were found in another ceramic systems such as sintering of TiO$_2$ as well as UO$_2$. Minimum relative density data of un-modified samples sintered at both temperatures were correlated well with their microstructures containing large pores. On the contrary, the observed microstructures of Bi$_2$O$_3$-added BNZ ceramics less than 4 wt% Bi$_2$O$_3$ at 850 and 900 °C showed a slight decrease of BNZ grain and pore sizes to an approximate range of 3.5–4 µm and 0.2–0.5 µm, respectively. Their densities were therefore enhanced. However, SEM images could not demonstrate clearly the existence of bismuth-oxide based liquid phase at BNZ grain boundaries and this made it difficult to see effect of additive sintering aid on enhancing microstructural evolution. In case of microstructural feature of BNZ/6 wt% Bi$_2$O$_3$ fired at temperature of 850 and 900 °C, it was observed that average grain size of both cases were decreased to 3.5 µm. Doped solid particles were assumed to affect microstructural evolution of Bi$_{0.6}$Na$_{0.4}$ZrO$_3$ material due to the melting of additive to form liquid at temperature above 825 °C and provided a path for enhanced matter transport at grain boundaries. Usually, diffusion of atoms and ions through a liquid is much faster than in a solid. However, the distance
of diffusion path depended on the volume fraction of liquid. Increase of the volume fraction providing long diffusion distance leaded to decrease the rate of grain growth. From the contribution of liquid phase, size of grains of modified BNZ ceramic at maximum Bi$_2$O$_3$ concentration sintered at temperature of 850 and 900 °C were decreased up to 27% when compared with pure material fired at the same temperature. This behaviour was in agreement with several reports attempting to explain the dependence of grain growth on the volume fraction of liquid. Increase of the volume fraction providing a result of average pore size decreasing to less than 0.5 µm. The measured maximum relative density values (89–90%) in both samples were accompanied by a decrease of size and number of pores. This seemed to agree well with a study on adding Bi$_2$O$_3$ in fabrication of Bi$_{0.5}$Na$_{0.5}$ZrO$_3$ ceramic. In addition, SEM micrographs indicating solidified liquid at BNZ grain boundaries and numerical details of Bi concentration between grain as well as grain boundary area could confirm clearly that mentioned material was bismuth oxide although it was not detected with XRD analysis. Non-perovskite liquid of bismuth oxide at grain boundaries built on the path for diffusion further influenced microstructural development.

Based on this study, Bi$_{0.5}$Na$_{0.5}$ZrO$_3$ ceramic implied enhancement of densification employing liquid-phase sintering process as Bi$_2$O$_3$ sintering aid at firing condition below 1100 °C. However, addition of Bi$_2$O$_3$ more than 4 wt% induced unwanted Bi$_{1.38}$Zr$_{0.62}$O$_{12.31}$ compound at sintering temperature of 900 °C. This best condition found in this study was BNZ/6 wt% Bi$_2$O$_3$ sintered at 850 °C. Accordingly, sintering at temperature of 850 °C and added Bi$_2$O$_3$ content higher than 6 wt% were believed to be the main factors for obtaining the product of phase pure BNZ ceramic with relative density close to 100%. In the near future, preparation of BNZ ceramic with increased added Bi$_2$O$_3$ concentration will be investigated.

In conclusion, Bi$_2$O$_3$-doped Bi$_{0.5}$Na$_{0.5}$ZrO$_3$ ceramics were successfully fabricated using the liquid-phase sintering method at low temperature. Bi$_2$O$_3$ material produced liquid phase at grain boundaries of Bi$_{0.5}$Na$_{0.5}$ZrO$_3$ ceramics and increased diffusion distance. This influenced the microstructural evolution of BNZ system including a reduction of grain growth rate, size of grain, number of pores and pore size. An improved change of these parameters caused densification of pure ceramic to enhance. In consequence, it promoted an increase of relative density value. In firing at 900 °C, excess adding of Bi$_2$O$_3$ (> 4 wt%) induced formation of Bi$_{7.38}$Zr$_{0.62}$O$_{12.31}$ compound. The optimum sintering temperature for this material should therefore be less than 900 °C with Bi$_2$O$_3$ added as a sintering aid.

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