Structure and Electrical Properties of Microwave Sintered BTS-BCT-xBF Lead-Free Piezoelectric Ceramics

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Abstract: Barium titanate (BT)-based ceramics are one of the promising piezoelectric materials for environment-friendly electro-mechanical transformation. However, high performance materials are often sintered at high temperatures, resulting in volatile components and increased energy consumption. Here, 0.82Ba(Ti0.39Sn0.11)O3–(0.18–x)(B0.7C0.3)TiO3–xBiFeO3 (BTS-BCT-xBF) piezoelectric ceramics were prepared by microwave sintering (MWS) method, and the structure and properties were emphatically studied, aiming to reveal the regulatory mechanism of MWS on the structure and properties. Compared with conventional solid sintering (CS), the phase structure presents a similar evolution in MWS ceramics as a function of BF, while the more refined grain size and the denser structure are observed in MWS ceramics. The electrical properties (e.g., $d_{33}$, $\varepsilon_r$, $\tan \delta$, etc.) of MWS ceramics are superior to the CS ceramics owing to the refined grain size and denser microstructure. It is worth noting that the energy storage performance (e.g., energy storage density, energy storage efficiency) significantly outperformed expectations due to the slender hysteresis loop resulting from the smaller grain and high cubic phase. Therefore, the MWS sintering mechanism can further drive practical application of BT-based ceramics.

Keywords: lead-free; BT-based ceramics; microwave sintered; structure; electrical properties

1. Introduction

As the representative of piezoelectric material, barium titanate (BT)-based ceramics are widely used in electronic components due to their excellent electric properties, such as micro-capacitors, ferroelectric memory, etc. [1,2]. However, the disadvantages, including high sintering temperature, poor temperature stability, and low breakdown field strength, seriously hinder the application in complex environment. In order to further seize the market, many means are used to overcome the shortages of BT-based materials, especially the various ways of preparation [3–5].

It is reported that the preparation of piezoelectric ceramics has many options in terms of sintering processes, such as spark plasma sintering (SPS) [6], atmosphere sintering [7], hot pressed sintering [8], microwave sintering (MWS) [9,10], etc. Among those options for sintering, MWS possesses the unique advantage of rapid sintering for materials at low temperature during the preparation process, which have been widely used to synthesize new materials [11,12]. It can also enhance the density of materials significantly, and improve the electrical properties. Takahashi H et al. reported a densified BT-based material system with high $d_{33}$ prepared by MWS [13]. Bafandeh et al. prepared KNN-based ceramics by conventionally sintering and microwave-sintering, and systematically compared the microstructure, ferroelectric, and piezoelectric properties of those samples, showing that MWS can inhibit grain growth and enhance densification in this ceramic, which results in improvements in the electric properties (e.g., piezoelectric properties, ferroelectric properties,
strain behavior, etc.) [14]. The effects of different sintering methods on the phase boundary and microstructure of BT-based ceramics were studied by Gao et al. [15]. Compared with conventional solid sintering (CS), the phase structure of microwave sintering ceramics remains almost stable, and the samples present a smaller grain size and uniform grain distribution. Another advantage of MWS is that it can significantly reduce temperature and time during the sintering process. However, the effects of MWS on the structure and properties have not been investigated sufficiently in BT-based ceramics with high performance. Revealing the regulatory mechanism of MWS on the structure and properties of BT-based ceramics is an urgent aim, which can further enrich strategies for tuning their performance.

Recently, a series of BT-based ceramics with excellent performance have been obtained by constructing multiphase boundaries through chemical modification (e.g., Ca, Sr, Zr, Sn, Hf, etc.). For example, Wang et al. [16] reported (Ba_{0.85}Ca_{0.15})(Ti_{0.9}Zr_{0.1})O_3 (BCTZ) ceramics with large \(d_{33}\) (~650 pC/N), and Zhu et al. observed a high \(d_{33}\) in (Ba, Ca)TiO_3–0.45Ba(Sn,Ti)O_3 ceramics. In particular, Zhao et al. reported high piezoelectric properties (\(d_{33}=700 \pm 30 \) pC/N) in 0.82Ba(Ti_{0.89}Sn_{0.11})O_3·0.18(Ba0.7Ca0.3)TiO_3 materials, which are comparable to that of lead-based and lead-free based piezoelectric materials. However, the ceramics with optimal piezoelectric properties were always sintered at a relatively high temperature, which could lead to the compositional volatile and increased the energy consumption during the sintering process. Therefore, it is expected that a lower temperature sintering could be used for fabrication, which will cut down the processing costs especially in production.

To further reveal the mechanism of MWS on the structure and properties BT-based ceramics with high performance, the Ba(Ti_{0.89}Sn_{0.11})O_3·(Ba_{0.7}Ca_{0.3})TiO_3 material system was chosen to verify the efficiency of this strategy in BT-based ceramics. In addition, BiFeO_3 (BF), a typical ferroelectric material with large \(P_t\) and high \(T_c\), was also introduced to this material system, not only aiming to reduce the sintering temperature but also further modify the structure and properties [17–19]. For better comparison the effects of MWS and CS on the structure and properties, two sets of 0.82Ba(Ti_{0.89}Sn_{0.11})O_3·(0.18-x)(Ba_{0.7}Ca_{0.3})TiO_3·xBiFeO_3 (BTS-BCT-xBF) samples were prepared by MWS and CS, aiming to reveal the regulatory mechanism of MWS and CS on the structure and electrical properties of lead-free piezoelectric ceramics, which may promote the practical application of BT-based ceramics.

2. Experimental Procedure

The 0.82Ba(Ti_{0.89}Sn_{0.11})O_3·(0.18-x)(Ba_{0.7}Ca_{0.3})TiO_3·xBiFeO_3 (BTS-BCT-xBF, \(x=0.002, 0.004, 0.006, 0.008, 0.010, 0.012\)) piezoelectric ceramics were prepared by microwave sintering and normal sintering. Raw materials, including BaCO_3 (99.0%), CaCO_3 (99.0%), SnO_2 (99.5%), TiO_2 (98%), Bi_2O_3, and Fe_2O_3 (99%), were mixed with ZrO_2 balls and the medium of ethanol for 12 h in planetary ball mill. After being dried in oven and calcined at 1200 °C for 2 h, these powders were mixed with polyvinyl alcohol (PVA, 7–8 wt%) and pressed into pellets of ~10 mm in diameter and ~1 mm in thickness. After the PVA binder burned out, the pellets were divided into two groups. One group was sintered at 1350 °C for 3 h by conventional sintering method, and the other group was sintered at 1290 °C for 2 h by microwave sintering method. Silver paste was used as electrodes for electrical measurements, and was fired at ~650 °C for 30 min. These samples were poled in a silicone oil bath under a dc field of ~3.0 kV/mm for 30 min.

X-ray diffractometer (XRD) with a CuK\(_\alpha\) radiation (DX-2700, Haoyuan Instrument, Dandong, China) was used to confirm phase structure of the samples under ~40 kV, ~30 mA. Their microstructure and chemical compositions were measured by a field-emission scanning electron microscope (FE-SEM) (JSM-7500, JEOL, Showa, Japan). The dielectric constant (\(\varepsilon_r\)) varying with temperature (~150–200 °C) was measured by using an LCR analyzer (HP 4980, Agilent, Palo Alto, USA) in connection with a temperature-controlled instrument. The polarization versus electric field (P–E) hysteresis loops and the strain-electric field (S–E) curves were measured at 1 Hz with a ferroelectric tester (Radiant Premier II, Radiant
3. Results and Discussion

3.1. Phase Structure

Figure 1a,b plots room temperature XRD patterns of BTS-BCT-xBF ceramics in the 2θ range of ~20–60° sintered with MWS and CS, respectively. Compared with the standard cards’ diffraction peaks of R (PDF # 85-1797), O (PDF # 81-2200), and T (PDF # 05-0626) in BaTiO₃ materials, all ceramics show a pure perovskite structure without any other phase, indicating that BTS-BCT and BF formed multiphase ceramics within the study range of 0.002 ≤ x ≤ 0.012. In addition, the effect of BF content on the phase structure of BTS-BCTxBF ceramics sintered with different sintering methods was analyzed by the enlarged XRD pattern with multi-peak fitting via Lorentz method (see Figure 1c,f) [20,21]. There are four separated peaks ((002)ᵣ, (200)ᵣ, (022)ₒ and (200)ₒ) in the XRD pattern of the ceramics with x = 0.002, and three characteristics peaks ((020)ᵣ, (002)ₜ and (200)ₜ) in the ceramics with x = 0.004, indicating that the sintering method did not change the phase structure of the ceramics.

![Figure 1. XRD patterns of BTS-BCT-xBF ceramics sintered with (a) MWS and (b) CS; enlarged XRD patterns with multi peak fitting of BTS-BCT-xBF ceramics with (c) x = 0.002, (d) x = 0.004 sintered by MWS; enlarged XRD patterns with multi-peak fitting of BTS-BCT-xBF ceramics with (e) x = 0.002, (f) x = 0.004 sintered by CS.](image)

Figure 2 shows the εᵣ-T curves (~-80–120 °C) of BTS-BCT-xBF ceramics prepared by MWS and CS, measured at 1~100 kHz, which aimed to further verify the phase structure of BTS-BCT-xBF ceramics sintered with MWS and CS. Three dielectric peaks are observed in the curve of the ceramics with x = 0.002, corresponding to the rhombohedral-orthorhombic (R-O) phase transition temperature (Tᵣₒ) at lower temperature, the orthorhombic-tetragonal (O-T) phase transition temperature (Tₒₜ) near room temperature, and the Curie temperature (Tₙₜ) at high temperature, respectively. Only two dielectric peaks are observed in the curves of the ceramics with x > 0.002, which is related to the rhombohedral-tetragonal (R-T) phase transition temperature (Tᵣₕₜ) at lower temperature and Curie temperature (Tₙₜ) at high temperature, respectively. One can see that the phase boundaries are very sensitive to BF content, as shown in Figure 2a–f. Compared with CS samples of x = 0.002,
the $T_{O-T}$ (MWS: $\sim 25 ^\circ C$, CS: $\sim 28 ^\circ C$) and $T_{R-O}$ (MWS: $\sim 6 ^\circ C$, CS: $\sim 10 ^\circ C$) moves towards low temperature, and the $T_c$ (MWS: $\sim 52 ^\circ C$, CS: $\sim 53 ^\circ C$) keeps almost stable in MWS ceramics, indicating that the O-T phase boundaries is obtained near room temperature. With increasing BF to 0.004, all ceramics show R-T phase boundaries because the O-T and R-O phases form a convergence zone near room temperature, and the $T_{R-T}$ of MWS sample is higher than the CS ceramics. With further increasing BF, the $T_{R-T}$ and $T_c$ decrease simultaneously, and the T-C phase boundaries are observed near room temperature in those ceramics. In addition, $T_c$ of MWS ceramics ($52 ^\circ C \rightarrow 22 ^\circ C$) changes faster than CS ceramics ($53 ^\circ C \rightarrow 39 ^\circ C$) in BTS-BCT-xBF ceramics. Figure 2g,h plots the phase diagram of MWS and CS BTS-BCT-xBF ceramics. The evolution of phase boundaries is highly matched with the XRD patterns, that is, O-T for $0.002 \leq x < 0.004$, R-T for $0.004 \leq x < 0.006$, and T-C for $x \geq 0.006$.

![Figure 2](image)

Figure 2. (a–f) $\varepsilon_r$-$T$ curves from $-80 ^\circ C$ to $120 ^\circ C$ of BTS-BCT-xBF ceramics sintered with MWS and CS. (g,h) Phase diagrams of BTS-BCT-xBF ceramics sintered with MWS and CS, respectively.

Figure 3 plots the modified Curie-Weiss law ($\ln(1/\varepsilon_r - 1/\varepsilon_m) - \ln(T - T_m)$) of BST-BCT-xBF ceramic, which is originated from the following formula:

$$1/\varepsilon_r - 1/\varepsilon_m = (T - T_m)^\gamma/C,$$

(1)

where $\varepsilon_m$ is the maximum value of relative dielectric constant and $\gamma$ is the degree of diffuseness. The higher $\gamma$ value is observed in MWS ceramics, indicating that the MWS ceramics show a stronger relaxation degree than the CS ceramics. This can be ascribed to the smaller grain sizes in MWS ceramics, which will be discussed later.
one can find that the density first increases and then decreases in the samples as a function of BF, and the density of MWS ceramics is higher than CS ceramics, indicating that MWS plays a more effective role in enhancing density.

3.2. Microstructure

Figure 4a–d and Figure S1 (see the Supplementary Materials) show the SEM surface morphology of MWS ceramics. A phenomenon of bimodal distribution of large and small grains is observed in MWS ceramics with 0.002 \( \leq x \leq 0.004 \). The average grain size of large grains is about 43.65–47.96 \( \mu \)m, and the small grains of 1.0–1.8 \( \mu \)m surround the large grains, resulting in the improved density by filling the pores around the large grains, and the average grain size decreases from 21.25 \( \mu \)m to 13.99 \( \mu \)m. With further increasing BF in MWS ceramics, the average grain size decreases from 21.25 \( \mu \)m to 13.99 \( \mu \)m, and the bimodal distribution of grain disappears in those ceramics. Figure 4e–h shows the SEM morphology of MWS ceramics. A phenomenon of bimodal distribution of large and small grains is observed in MWS ceramics with 0.002 \( \leq x \leq 0.012 \). The average grain size of BST-BCT-xBF ceramics with MWS and CS, (a) \( x = 0.002 \), (b) \( x = 0.004 \), (c) \( x = 0.012 \).

Figure 3. Plots of \( \ln(1/\varepsilon_r - 1/\varepsilon_m) \) versus \( \ln(T - T_m) \) of BST-BCT-xBF ceramics with MWS and CS, (a) \( \gamma = 1.94 \), (b) \( \gamma = 1.79 \), (c) \( \gamma = 1.92 \).
3.3. Dielectric and Ferroelectric Properties

Figure 5a,b shows the $P$-$E$ and $S$-$E$ loops for MWS and CS ceramics, measured at room temperature and 30 kV/cm. All samples present saturated $P$-$E$ loops and standard $S$-$E$ curves, indicating that all samples are normal ferroelectrics. Due to the high $P_r$ for BF, the $P_r$ increases with increasing the proper BF, and the ferroelectric properties would deteriorate in the ceramics with adding excessive BF, as shown in Figure 6a. The $P_r$ of MWS samples is smaller than that of CS ceramics, which results from the grain refinement leading to a decrease in $P_r$ due to the increased clamping effect of domain walls relative to grain boundaries [25,26]. Compared with the CS samples with $x > 0.004$, the $P_r$ decreases sharply in MWS samples, which is related to the evolution of phase boundary. With increasing BF content, the cubic phase of MWS ceramics gradually dominates in the phase structure near room temperature, and then results in deteriorating of the ferroelectric properties. The $E_c$ of two group samples keeps almost unchanged with increasing BF content, as shown in Figure 6b. The $S_{pos}$ (the strain between the maximum strain and the strain under 0 kV/cm) of MWS and CS ceramics are shown in Figure 6c, which first increases and then decreases with increasing BF, achieving maximum $S_{pos}$ value (MWS: ~0.102%, CS: ~0.105%) at the ceramics with $x = 0.004$, which can be ascribed to the low energy barrier in multiphase boundary. Moreover, the maximum strain in MWS ceramics is lower than CS due to the limitation of a large number of fine grain when $x \leq 0.004$. Figure 6d plots the dielectric constant ($\varepsilon_r$) of CS and MWS ceramics, measured at 10 kHz. The $\varepsilon_r$ increases continuously in MWS ceramics, while first decreasing and then increasing in CS ceramics with increasing BF. The higher $\varepsilon_r$ of MWS ceramics results from the finer grains because $\varepsilon_r$ increases with decreasing grain size when the average grain size is greater than 1.2 $\mu$m [27,28]. Of course, the high density of MWS ceramics also contributes positively to this. The dielectric loss ($\tan \delta$) of two sets of samples is very close, and remains around 0.02–0.04, as shown in Figure 6e. This suggests that the $\tan \delta$ is not sensitive to the sintering methods, and this phenomenon has been reported in other BT-based ceramics [29]. The $d_{33}$ of MWS and CS ceramics presents a similar change trend, as shown in Figure 6f. That is, $d_{33}$ first increases and then decreases with the increase of BF content, reaching the maximum values (MWS: ~425 pC/N, CS: ~360 pC/N) at $x = 0.004$. This can be attributed to the R-T phase boundaries near room temperature in the ceramic with $x = 0.004$. As is known, multiphase coexistence

| $x$   | Sintering Method | Density (g/cm$^3$) | Sintering Method | Density (g/cm$^3$) |
|------|----------------|--------------------|----------------|--------------------|
| 0.002 | CS             | 5.8005             | MW             | 5.8364             |
| 0.004 |                | 5.8285             |                | 5.9100             |
| 0.006 |                | 5.7959             |                | 5.8653             |
| 0.008 |                | 5.7644             |                | 5.8435             |
| 0.010 |                | 5.7721             |                | 5.8195             |
| 0.012 |                | 5.8059             |                | 5.8065             |

Figure 4. Surface morphology images of the BTS-BCT-xBF ceramics sintered by (a–d) MWS and (e–h) CS, (i) Grain sizes of MWS and CS BTS-BCT-xBF ceramics.
provides more possible polarization directions, resulting in easy polarization rotation, which can greatly facilitate the polarization switching [30]. Moreover, the bimodal grain sizes distribution can also enhance the $d_{33}$ value in MWS ceramics [25].

![Figure 5](image_url)  
**Figure 5.** $P$-$E$ and $S$-$E$ loops of (a) MWS and (b) CS sintered BTS-BCT-xBF ceramics.

![Figure 6](image_url)  
**Figure 6.** (a) $P_r$, (b) $E_c$, (c) $S_{pos}$, (d) $e_r$, (e) $\tan \delta$, and (f) $d_{33}$ of BTS-BCT-xBF ceramics sintered with MWS and CS.

### 3.4. Energy Storage Performance and Electrostriction

Figure 7a shows the $P$-$E$ loops of the MWS and CS ceramics with $x = 0.012$. It is reported that the thinner $P$-$E$ loop with high $P_{max}$ and low $P_r$ value is beneficial to energy storage due to the low energy loss in the materials. Compared with the $P$-$E$ loops of CS ceramics, a thinner $P$-$E$ loop is obtained in MWS ceramics, indicating that it has better energy storage performance than CS ceramics. To further make a quantitative analysis on the energy storage performance of MWS and CS ceramics, the calculated energy storage performance of MWS and CS ceramics with $x = 0.012$ are plotted in Figure 7b,c, measured at 30 kV/cm. As is known, the discharge storage density $W_1$ can be obtained from the
integration of the discharge curve, and the area between the charge curve and the discharge curve is the energy loss $W_2$, $\eta = W_1/(W_1 + W_2)$ is used to express energy storage efficiency [31–33]. Although the $P_{\text{max}}$ of MWS ceramics is slightly lower than that of CS, the MWS ceramics have better energy storage density and energy storage efficiency (MWS: $W_1 = 0.1019 \text{ J/cm}^3$, $\eta = 90.10\%$; CS: $W_1 = 0.0883 \text{ J/cm}^3$, $\eta = 73.46\%$) due to the slender hysteresis loop resulting from the smaller grain and high cubic phase. Therefore, MWS may be beneficial to the enhanced energy storage performance. Polarization reveals that the strain is mainly contributed by electrostriction in the ceramics, and the $S$-$P$ curves of some components deviate from the quadratic relationship caused by the irreversible domain switching [34–39]. The $Q_{33}$ of the ceramics shows a similar trend, that is, $Q_{33}$ increases first and then decreases, and reaches the maximum value at $x = 0.004$ (MWS: $\sim 0.049 \text{ m}^4/\text{C}^2$, CS: $\sim 0.045 \text{ m}^4/\text{C}^2$), as shown in Figure 8. Moreover, the MWS ceramics have higher $Q_{33}$ than CS.

![Figure 7](image_url) 

**Figure 7.** (a) P-E loops, energy storage performance of (b) MWS and (c) CS of BTS-BCT-$x$BF ceramics with $x = 0.012$.

![Figure 8](image_url) 

**Figure 8.** S-P loops of BST-BCT-$x$BF ceramics sintered with (a) MWS and (b) CS.

4. Conclusions

In this work, $0.82\text{Ba(Ti}_{0.82}\text{Sn}_{0.11})\text{O}_3-(0.18-x)(\text{Ba}_{0.7}\text{Ca}_{0.3})\text{TiO}_3-x\text{BiFeO}_3$ (BTS-BCT-$x$BF) piezoelectric ceramics were prepared by microwave sintering method and conventional sintering method. The phase structure of two group samples presents a similar evolution, and the more refined grain size was observed in MWS ceramics, resulting in the denser structure than CS ceramics. Owing to the refined grain size and denser microstructure in MWS ceramics, the electrical properties (e.g., $t_1 \sim 8273$, $d_{33} \sim 425 \text{ pC/N}$, tan $\delta \sim 0.022$, Strain $\sim 0.102\%$, $Q_{33} \sim 0.049 \text{ m}^4/\text{C}^2$) were better than the performance of CS ceramics (e.g.,
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