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1. Introduction

Ferro-/piezo-electric ceramics have been widely used in many fields, owing to their applications as sensors, actuators, ultrasonic transducers, and so on.^{1,2} In particular, Pb-based perovskite ferroelectric ceramics have been under intense investigation for many years because of their much more excellent ferroelectric and piezoelectric properties than those of most Pb-free ceramics.3-7 However, issues such as the environmental restrictions on the usage of lead, the depoling and aging of PbZrO₃-PbTiO₃ (PZT) piezoelectric ceramics above 200 °C have attracted a great deal of attention to search for high Curie temperature lead-reduced materials.8,9 Therefore, BiMeO3-PbTiO₃ (Me represents cations of rare earth metal or transition metal, such as Sc, Fe, In, $Mg_{1/2}Ti_{1/2}$, and so on) binary system as a class of lead-reduced materials has been explored and developed, because most of BiMeO₃-PbTiO₃ family were found to display high Curie temperatures and fairish piezoelectric coefficients.¹⁰⁻¹⁵ For example, BiScO₃-PbTiO₃ (BS-PT) composition in the MPB region exhibits excellent piezoelectric properties d_{33} = 460 pC N⁻¹ and $T_{\rm C}$ = 450 °C, which is comparable with many Pb-based ceramics.¹⁶ The BiFeO₃-PbTiO₃ system displays high ferroelectric Curie temperature ($T_{\rm C} \sim 640~^\circ{\rm C}$) in the MPB region and weak-ferromagnetic ordering at high temperature ($T_{\rm N} \sim$ 104 °C), becoming a multiferroic material which was investigated widely and intensively.¹⁷ $(1 - x)Bi(Mg_{1/2}Ti_{1/2})O_3-xPbTiO_3$

Ferroelectric, piezoelectric properties and thermal expansion of new $Bi(Ni_{3/4}W_{1/4})O_3 - PbTiO_3$ solid solutions

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Ceramics of the Bi(Ni_{3/4}W_{1/4})O₃-PbTiO₃ (BNW-PT) ferroelectric system were synthesized using a conventional solid-state sintering process. A morphotropic phase boundary (MPB) region separating tetragonal and rhombohedral phases has been determined based on X-ray diffraction (XRD) analysis. The composition-dependent phase transition behavior, thermal expansion, di/ferro/piezoelectric properties have been systemically investigated. The volume TECs (thermal expansion coefficients) for 0.1BNW-0.9PT and 0.2BNW-0.8PT are -1.01×10^{-5} per °C and -0.94×10^{-5} per °C, respectively, indicating the negative thermal expansion of some compositions for the binary system. The BNW-PT ceramics have Curie temperatures, T_{c} , ranging from 460 °C to ~152 °C with the variation of BNW constituent. The tetragonal-rich composition 0.20BNW-0.80PT is found to have the largest remnant polarization, $P_r \sim$ 23.4 μ C cm⁻². The highest piezoelectric coefficient $d_{33} \sim 145$ pC N⁻¹ is achieved at MPB composition 0.32BNW-0.68PT. The maximum strain value ~0.194% is also obtained in the 0.32BNW-0.68PT ceramic.

> (x = 0.37) possesses high Curie temperature $T_{
> m C} \sim 450~^\circ{
> m C}$ and piezoelectric coefficient $d_{33} \sim 225 \text{ pC N}^{-1}$.¹⁸ The morphotropic phase boundary (MPB) composition x = 0.62 of $(1 - x)Bi(Ni_{1/2}Hf_{1/2})$ O₃-xPbTiO₃ (BNH-PT) shows a high piezoelectric coefficient of $d_{33} \sim 446 \text{ pC N}^{-1}$ and Curie temperature $T_{\rm C} \sim 290 \, {}^{\circ}\text{C.}^{19}$ Given these examples, there is no doubt that many BiMeO₃-PbTiO₃ systems may be useful for high temperature piezoelectric applications. In addition, many PbTiO₃-based compounds were found to display negative or near-zero thermal expansion, which is of great importance for designing some precision devices.19-22 Especially, it is meaningful to get negative or nearzero thermal expansion for BiMeO₃-PbTiO₃, because the Bi substitution plays an important role in enhancing the tetragonality c/a (high tetragonality c/a corresponds to negative thermal expansion) and Curie temperature $T_{\rm C}$ due to the strong coupling between the A-site Pb/Bi cations and the B-site ferroelectric cations.^{23,24} Therefore, in order to get a material which shows good ferro/piezoelectric properties, high Curie temperature and negative thermal expansive, it is worth to explore new BiMeO₃-PbTiO₃ system. Here, Bi(Ni_{3/4}W_{1/4})O₃-PbTiO₃ (BNW-PT) were chosen owing to the possibility of stable solid solution formed, according to the calculated tolerance factor $t \approx 0.89$ of BNW by the following formula (1):

$$t = \frac{R_{\rm A} + R_{\rm O}}{\sqrt{2}(R_{\rm B} + R_{\rm O})} \tag{1}$$

(R_A , R_B , R_O are the effective ionic radius of A, B cations and O anion in ABO₃ compounds). When *t* is located in the range from 0.78 to 1.05, the compound may be formed in a stable perovskite structure.

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In this work, BNW–PT ceramics were prepared by conventional solid-state reaction. The phase structures, thermal expansion, dielectric, ferroelectric and piezoelectric properties of BNW–PT ceramics were studied and analyzed systematically.

2. Experimental procedure

The ceramic samples of *x*BNW–(1 - x)PT with compositions of x = 0.10, 0.20, 0.3, 0.32, 0.34, 0.36 and 0.40 were prepared by a conventional solid-state reaction process. Ceramic samples were fabricated using metal oxide powders with 99.9% purity,

sample preparation details are as follows: NiO and TiO₂ powders were weighed according to the stoichiometric amount, while PbO and Bi_2O_3 were weighed with 1–2 mol% excess to compensate for volatilization during sintering and WO₃ with 3 mol% excess to compensate for sublimation. At first, these metal oxide powders were mixed homogeneously and milled using ethanol for 2 h. After milling, the powders were pressed into pellets with 25 mm diameter at 4 MPa. The pressed powders were calcined at 900 °C for 5 h in air. After calcination, the pieces were remilled using ethanol for 2 h and mixed with 5 wt% polyvinyl alcohol (PVA) as a binder. The powders were



Fig. 1 Powder X-ray diffraction patterns of xBNW-(1-x)PT ceramics with x = 0.1-0.4 (right); the enlarged XRD profiles of (200) reflections for MPB compositions (left).



Fig. 2 Pseudocubic (200) reflections of the BNW-PT ceramics: (a) 0.30 BNW-0.70PT; (b) 0.32 BNW-0.68PT; (c) 0.34 BNW-0.66PT.

pressed into pellets with 5–6 mm thickness at 7 MPa, and then heated up to 550 °C and kept for 2 h to eliminate the binder. Finally, the discs were sintered at 930 °C to 960 °C for 3 h in a sealed Al_2O_3 crucible to form a series of desired ceramic samples.

The powder XRD patterns of sintered ceramics were collected by XRD equipment with Cu K α radiation (Rigaku, MinFlex 600, Japan). The microstructure of the fracture surface of sintered ceramic pieces was observed by scanning electron microscope (SEM, SU-8010, Tokyo, Japan). Two faces of pellets with 0.5 mm in thickness were polished and painted with silver paste for electrodes. Dielectric and piezoelectric measurements were carried out after ferroelectric measurements. The piezoelectric constant d_{33} was measured by a piezo- d_{33} quasi static meter (ZJ-3AN; Institute of Acoustics, Beijing, China). The temperature dependence of dielectric properties was collected on a TH2816A LCR (Changzhou, China) analyzer with a heating rate of 3 °C min⁻¹. The ferroelectric and strain measurements

were carried out by a ferroelectric test system (TF Analyzer 2000E; axiACCT, Aachen, Germany). The high temperature XRD data was collected on a power diffractometer (Rigaku, Ultima-IV, Japan). The macroscopical thermal expansion coefficients of ceramics were determined from dilatometric measurements using a thermal expansion analyzer (DIL402PC, Germany) with clubbed samples (diameter 4–6 mm, length 11 mm) at a heating speed of 8 $^{\circ}$ C min⁻¹.

3. Results and discussion

3.1 Structural characterizations

Fig. 1 shows the powder XRD patterns of the *x*BNW–(1 - x)PT ceramics with x = 0.10, 0.20, 0.3, 0.32, 0.34, 0.36 and 0.40. As shown at the bottom of the figure, 0.10BNT–0.90PT ceramics have crystallized in the tetragonal perovskite structure, resulting from more tetragonal symmetry PT content in the composition. The XRD profiles of (200) reflection splits into two peaks



Fig. 3 (a) The unit cell parameters a, c for all compositions; (b) the tetragonality c/a for all compositions.



Fig. 4 SEM micrographs of fracture surfaces for xBNW-(1 - x)PT ceramics: (a) 0.10BNW-0.90PT; (b) 0.20BNW-0.80PT; (c) 0.30BNW-0.70PT; (d) 0.40BNW-0.60PT.

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for the tetragonal (T) phase from x = 0.10 to x = 0.32, with an intensity ratio of the T(002) and T(200)/(020) peaks of 1:2. However, with increasing BNW content, the XRD profile of (200) reflection shows only a single peak R(200) for the rhombohedral (R) phase due to all the planes of (200) sharing the same lattice parameters from x = 0.34 to x = 0.40, resulting from BNW having rhombohedral symmetry at room temperature. Therefore, based on the (200) reflections (marked by black box around $2\theta = 45^{\circ}$), the MPB region from rhombohedral phase to tetragonal phase is determined at about 30-34 mol% BNW. The pseudocubic (200) reflections of xBNW-(1 - x)PT ceramics (x = 0.30-0.34) are shown in Fig. 2. The (200) reflection is composed of different numbers of peaks. The dominant peaks 1 and 2 represent the tetragonal phase, and the peak 3 corresponds to the rhombohedral phase. Thus, there is the coexistence of rhombohedral and tetragonal phases for 0.32BNW-0.68PT composition. The ratio of the rhombohedral phase and tetragonal phase for 0.32BNW-0.68PT composition in MPB region

has been calculated based on the formula $\frac{R}{T}\!=\!$ $I_{R(200)}$ $\overline{I_{\rm T(200)} + I_{\rm T(002)}}$ and the ratio value is 0.495. In addition, it can be seen that the phase is becoming impure with the increase of BNW content, the formation of impurity can be seen in position of $2\theta = 27.69^{\circ}$ for the compositions x = 0.36 and 0.40, indicating that it is difficult to form the stable perovskite structure for BNW-PT solid solution when $x_{(BNW)} > 0.36$. Based on the analysis of the impure peak in 27.69°, we estimate that the impure phase should be Bi3.84W0.16O6.24, the substance is formed easily in some solid-state reaction when Bi2O3 and WO3 are used as reaction raw materials. Yao et al. also observed the formation of Bi_{3.84}W_{0.16}O_{6.24} when they synthesized Bi₂WO₆ by solid-state method.²⁵ Fig. 3 presents that the compositional dependences of lattice parameter a(c) and the tetragonality c/a, it should be noted that there is an increase in *a* lattice parameter with BNW raising, but a decrease in c lattice parameter, which leads to a decrease of the tetragonality c/a with the increase of BNW.

Composition	Grain size (μm)	Theoretical density $(g \text{ cm}^{-3})$	Actual density $(g \text{ cm}^{-1})$	Relative density (%)
0.10BNW-0.90PT	5	8.5026	8.0293	94.43
0.20BNW-0.80PT	5	8.5914	8.2402	95.91
0.30BNW-0.70PT	0.6	8.7083	8.4822	97.40
0.40BNW-0.60PT	0.4	8.7755	8.6115	98.13



Fig. 5 High temperature X-ray diffraction patterns and the variation of the unit cell parameters a(c) with respect to the temperature for 0.10BNW-0.90PT (a and c) and 0.20BNW-0.80PT (b and d).

The SEM micrographs of a fractured surface for selected xBNW-(1 - x)PT ceramics with x = 0.10, 0.2, 0.3 and 0.4 are presented in Fig. 4. It can be seen clearly that well sintered ceramics were obtained for all ceramics. The grain size is gradually decreasing with increasing the BNW content. The actual density was measured using Archimedes method, and the theoretical density was calculated based on XRD. The theoretical and actual density reaches up to 94.4–98.1%, indicating high density.

3.2 Thermal expansion

The high temperature X-ray powder diffraction patterns (HTXRD) and unit cell parameters *a*, *c* of the *x*BNW–(1 - x)PT ceramics with x = 0.1 and 0.2 are shown in Fig. 5. The Curie temperatures $T_{\rm C}$ of the two compositions by HTXRD are in good agreement with those of dielectric spectra (the Section 3.3). The Curie temperatures $T_{\rm C}$ are in the range of 420–460 °C for 0.1BNW–0.9PT and 320–360 °C for 0.2BNW–0.8PT, based on (100) peaks. The Fig. 6(a) presents the tetragonality *c/a* as a function of temperature. It is observed that the tetragonality *c/a* as a larger decrease of the tetragonality *c/a* for 0.1BNW–0.9PT



Fig. 6 (a) The tetragonality c/a as functions of temperature for xBNW–(1 – x)PT ceramics with x = 0.10-0.30; (b) the unit cell volume as functions of temperature for xBNW–(1 – x)PT ceramics with x = 0.10-0.30.

ceramic, which leads to enhanced NTE.26 Therefore, as shown in Fig. 6(b), both 0.1BNW-0.9PT and 0.2BNW-0.8PT display reduced cell volumes, but 0.3BNW-0.7PT shows increased cell volume. The volume TECs for these ceramics are listed in Table 2. The measurements of dilatometric thermal expansion on clubbed BNT-PT ceramics were taken to characterize the materials' apparent thermal expansion, and the results are shown in Fig. 7. The linear TECs are also listed in Table 2. It can be seen that the TECs for 0.1BNW-0.9PT (-1.01×10^{-5} per °C) and 0.2BNW-0.8PT (-0.94×10^{-5} per °C) are very small, compared to BiFeO₃-PbTiO₃ (-3.92×10^{-5} per °C) and Pb_{1-r}- Cd_rTiO_3 (-2.40 × 10⁻⁵ per °C), indicating that the variation of temperature will have less impact on the volumes of the materials.^{21,22} In addition, the ceramic sintering process is much affected by their discrepant NTEs, it is difficult to sinter dense ceramics for the strong NTE materials, this is why the relative density is decreasing with the content of BNW decreasing (as shown in Table 2).26

3.3 Dielectric properties

The temperature dependences of the real permittivity (ε') and dielectric loss (tan δ) for the selected compositions *x*BNW-(1 – *x*)PT (*x* = 0.2, 0.3, 0.34, 0.36) measured at different frequencies are shown in Fig. 8. In general, the paraelectric-ferroelectric phase transition varies in the range from 460 to 152 °C, the Curie temperature *T*_C in MPB region is around 220 °C. It is

Table 2 Average TECs of xBNW- $(1 - x)$ PT ceramics ($x = 0.1-0.3$)							
			TEC $(10^{-5}$ per °C)				
Composition	$lpha_{ m V}$	α_{l}	Temperature range (°C)				
x = 0.1 x = 0.2 x = 0.3	$-1.01 \\ -0.92 \\ 1.98$	-0.16 0.27 0.49	30–450 30–350 30–240				



Fig. 7 Dilatometric curves of ceramic bars for xBNW-(1-x)PT for x = 0.10-0.30.



Fig. 8 Temperature dependence of dielectric permittivity ε' and dielectric loss tan δ for xBNW–(1 – x)PT ceramics: (a) 0.20BNW–0.80PT; (b) 0.30BNW–0.70PT; (c) 0.34BNW–0.64PT; (d) 0.36BNW–0.62PT.

common sense that the Curie temperature $T_{\rm C}$ of tetragonal ABO₃ perovskites is generally correlated to the tetragonality c/a. High tetragonality c/a means large distorted lattice symmetry, which requires higher temperature to eliminate the crystalline distortion.^{19,27} This is in excellent agreement with previous reports.^{28–30} Fig. 9 displays the compositional dependences of the Curie temperature $T_{\rm C}$ and room-temperature dielectric permittivity $\varepsilon'_{\rm RT}$, it is found that the 0.60BNW–0.40PT has the lowest Curie temperature, which is about 152 °C and increasing to the highest Curie temperature 460 °C for 0.10BNW–0.90PT. The compositional dependence of the Curie temperature is



Fig. 9 Curie temperature $T_{\rm C}$ and the real permittivity $\varepsilon'_{\rm RT}$ as functions of BNW content for xBNW-(1 - x)PT ceramics.

almost linear with changing BNW content in the process from tetragonal phase to the rhombohedral region. The result is analogous to some ferroelectric systems such as Pb(Yb_{1/2}Nb_{1/2}) O₃–PbTiO₃ ceramics and crystals.³¹ However, the room-temperature dielectric constant $\varepsilon'_{\rm RT}$ increases with the increase of BNW content. This is because ignoring the influence of the ionic radius difference, raising BNW content would lead to the increase of the sum of ionic polarizability, as a result, the dielectric constant would be enhanced.³²

3.4 Ferroelectric properties

Fig. 10(a) and (b) show the polarization-electrical field (P-E)hysteresis loops and switching currents of xBNW-(1 - x)PT ceramics with all compositions under bipolar electric fields of 60–90 kV cm⁻¹ at room temperature. Good saturated loops can be obtained, and the shape of hysteresis loops is developing from rectangular to slim with raising the content of BNW, as a result of the decrease in the c/a ratio and the smaller lattice distortion. Therefore, the coercive field $E_{\rm C}$ is monotonically decreasing from 57.15 to 13.63 kV cm⁻¹ with increasing the content of BNW. However, the remnant polarization P_r firstly increases and then decreases, and reaches a maximum value of \sim 23.4 μ C cm⁻² at 0.20BNW-0.80PT which is tetragonal phaserich composition near MPB region. The cause is that ferroelectric domains of compositions near the MPB are switched more easily, due to the coexistence of two phases giving a total of fourteen possible polarization directions (six tetragonal (001)and eight rhombohedral (111).³³ The P_r value gets largest in the



Fig. 10 (a) Polarization–electrical field (P-E) hysteresis loops for xBNW–(1 – x)PT ceramics with different x values at bipolar electric fields of about 60–100 kV cm⁻¹ at 5 Hz; (b) switching current; (c) bipolar electric-field-induced strain curves; (d) unipolar electric-field-induced strain curves.

composition 0.2BNW–0.8PT because the composition is close to the MPB region, and there is the coexistence of most tetragonal phase and a small amount rhombohedral phase. Therefore, the remnant polarization $P_{\rm r}$ may be optimal for 0.20BNW–0.80PT composition (Table 3).

Fig. 10(c) and (d) show the bipole and unipole electric-fieldinduced strain curves of *x*BNW–(1 - x)PT ceramics with different *x* measured at 5 Hz at room temperature. It is found that the compositions (x = 0.2-0.34) display more outstanding strain hysteresis under bipolar cycling, that is symmetric butterfly-shaped strain hysteresis with relatively large negative strain originating from domain back switching, whereas the rhombohedral compositions with x = 0.36 and 0.4 show less obvious strain hysteresis, this may be because domain switching is hindered by point defects resulted from oxygen vacancy with BNW increasing. Therefore, the strain reached the maximum value ~0.194% at the composition of x = 0.32 which lies in MPB region, as presented in Fig. 11(b). The Fig. 12(a) presents the XPS spectrum for some compositions, the peaks of O 1s are shown in Fig. 12(b). It can be seen that the peaks of O 1s are composed of oxygen in BNW-PT lattice ((I) in 529.5 eV) and oxygen vacancies/OH⁻ groups ((II) in 531 eV), which is in good agreement with many references.^{34,35} The XPS results show the existence of many oxygen vacancies for 0.36BNW-0.64PT and 0.40BNW-0.60PT ceramics. In order to make the results convincing, we further calculated the ratio of the oxygen atoms

Table 3 The remanent polarization P_r , coercive field E_c , Curie temperature T_c , real permittivity (ϵ '), dielectric loss (tan δ) and piezoelectric coefficient d_{33} for xBNW-(1 - x)PT ceramics

Composition	$P_{\rm r}$ (µC cm ⁻²)	$E_{\rm C}$ (kV cm ⁻¹)	S (%)	$T_{\rm C}$ (°C)	ε' (@RT)	$\tan \delta$ (@RT)	$d_{33} ({ m pC}\;{ m N}^{-1})$
0.10BNW-0.90PT	12.35	57.15	0.04	460	177	0.095	51
0.20BNW-0.80PT	23.40	44.65	0.132	360	369	0.096	89
0.30BNW-0.70PT	22.86	31.25	0.162	234	836	0.040	105
0.32BNW-0.68PT	17.01	23.91	0.194	218	1284	0.038	145
0.34BNW-0.66PT	8.84	17.33	0.166	192	1469	0.058	59
0.36BNW-0.64PT	4.97	13.64	0.093	179	1337	0.066	34
0.40BNW-0.60PT	4.72	13.63	0.074	152	1568	0.068	20



Fig. 11 Coercive field E_{C} , remnant polarization P_r , strain S% and piezoelectric constant d_{33} as functions of BNW content for xBNW-(1 - x)PT ceramics.



Fig. 12 (a) XPS spectra of xBNW-(1 - x)PT ceramics (x = 0.32, 0.34, 0.36 and 0.40), (b) high-resolution XPS spectra of the O 1s.

in lattice, the theory ratio for the two compositions should be 60% but the calculated ratios based on the XPS spectra are 56.60% for 0.40BNW–0.60PT ceramic and 57.13% for 0.36BNW–0.64PT ceramic, respectively.

3.5 Piezoelectric properties

The compositional dependence of piezoelectric coefficient d_{33} is presented in Fig. 11. It can be seen that piezoelectric coefficient

 d_{33} varies in the range 20–145 pC N⁻¹ with the variation of BNW content, which initially increases and reaches the maximum, and then decreases. The maximum of piezoelectric coefficient $d_{33} \sim 145$ pC N⁻¹ is obtained at 0.32BNW–0.68PT composition, which is located in the transition region from rhombohedral phase to tetragonal phase based on the XRD analysis. This is because fourteen polarization directions in MPB region enables optimized crystallographic orientations to be established easily in the poling process, which results in anomalously high piezoelectric properties in MPB region.³³

4. Conclusions

In summary, BNW-PT ferroelectric ceramics were fabricated by a conventional solid-state sintering process. The composition and temperature-dependent phase transition, thermal expansion behavior, electric properties have been systemically investigated. The results indicate that the variation of BNW content significantly affects the structure and electric properties of the BNW-PT ceramics. Ceramics of 0.1BNW-0.9PT and 0.2BNW-0.8PT display reduced cell volumes of -1.01×10^{-5} per °C and -0.94×10^{-5} per °C, respectively, but 0.3BNW-0.7PT shows increased cell volume of 1.98 \times 10⁻⁵ per °C. The dielectric measurements show that the BNW-PT ceramics have high Curie temperature T_C which ranges from 460 °C to 152 °C with the variation of BNW content. The tetragonal-rich composition 0.20BNW-0.80PT possesses the largest remnant polarization Pr \sim 23.4 µC cm⁻². The highest piezoelectric coefficient $d_{33} \sim$ 145 pC N^{-1} is achieved at MPB composition 0.32BNW-0.68PT. The maximum strain value ~0.194% is also obtained in the 0.32BNW-0.68PT ceramic.

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