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Composition and Temperature Dependence of Structure and Piezoelectricity in $(1-x)(K_{1-y}Na_y)NbO_3$ - $x(Bi_{1/2}Na_{1/2})ZrO_3$ Lead-Free Ceramics

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Lead-free piezoceramics with the composition $(1-x)(K_{1-y}Na_y)$ NbO₃-x(Bi_{1/2}Na_{1/2})ZrO₃ (KNyN-xBNZ) were prepared using a conventional solid-state route. X-ray diffraction, Raman spectroscopy, and dielectric measurements as a function of temperature indicated the coexistence of rhombohedral (R) and tetragonal (T) phase, typical of a morphotropic phase boundary (MPB) as the BNZ concentration increased and by adjusting the K/Na ratio. High remnant polarization ($P_r = 24 \mu \text{C/cm}^2$), piezoelectric coefficient ($d_{33} = 320 \text{ pC/N}$), effective piezocoefficient ($d_{33}^* = 420 \text{ pm/V}$), coupling coefficient ($k_p = 48\%$), and high strain (S = 0.168%) were obtained at room temperature, but significant deterioration of P_r , d_{33}^* , and k_p were observed by increasing from room temperature to 160°C (17.5 μ C/cm², 338 pm/V, and 32%, respectively) associated with a transition to a purely T phase. Despite these compositions showing promise for room-temperature applications, the deterioration in properties as a function of increasing temperature poses challenges for device design and remains to be resolved.

Keywords: lead-free ceramics; piezoelectric materials/properties; potassium-sodium niobate/KNN

I. Introduction

P IEZOELECTRIC materials have being extensively investigated because of their widespread applications ranging from medical, environmental, and industrial process monitoring to robotics, energy harvesting, and high-frequency communication systems. ^{1,2} Most commonly used materials are based on lead zirconate titanate Pb(Zr,Ti)O₃ (PZT) due to their high piezoelectric performance. ^{3–5} However, regulations against hazardous substances such as lead in electric and electronic equipment has stimulated research in lead-free piezoelectrics over the last two decades. ^{6–14}

Potassium-sodium niobate (K,Na)NbO₃ (KNN) is one of the leading candidates to replace PZT and has been extensively investigated due to its moderately large piezoelectric coefficient (d_{33}) and high Curie temperature ($T_{\rm C}$), ^{15,16} since its discovery in the 1950s. ¹⁷ In particular, research on KNN accelerated when Saito et al. (2004) reported giant $d_{33} \sim 416$ pC/N, comparable to those of PZTs, in textured (Li,Ta,

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Sb)-modified KNN, ^{18,19} However, disadvantages such as attaining well-densified ceramics due to the high volatility of the alkaline components and low piezoelectric properties in polycrystalline ceramics have prevented KNN being commercialized. ^{20–24} In general, dopants in KNN enhance piezoelectricity ($d_{33} > 200 \text{ pC/N}$) by pushing the orthorhombic (O) to tetragonal (T) transition boundaries closer to room temperature. ^{25–44} This effect is well documented for dopants such as LiSbO₃, LiTaO₃, Bi_{0.5}Na_{0.5}TiO₃, and BaTiO₃. ^{25–32} But coexistence of rhombohedral (R) and O phases has been reported for dopants with the general formula AZrO₃ (A=Ba, Sr, or Ca) or BiMO₃ (M=Fe, Sc, Co). ^{33–38} Compositions which rely on R-O coexistence generally, however, exhibit $d_{33} \le 230 \text{ pC/N}$ that are significantly lower the compositions which utilize the O-T phase boundary to optimize d_{33} .

Recently, PZT-like morphotropic phase boundary (MPBs) (R-T) have been reported in KNN-based ceramics by integrating compositions which were previously used to optimize the O-T and R-O phase boundaries.^{39–50} In 2011, Zuo et al. reported R-T phase coexistence in (Li,Ta,Sb) and BaZrO₃ multimodified KNN lead-free ceramics and obtained a high $d_{33} = 365 \text{ pC/N}.^{39-41}$ For similar compositions, Wu et al. obtained $d_{33} \sim 425 \text{ pC/N}.^{42}$ But more recently, Sb together with (Bi,M)NO₃ (M=Na, K, Li, Ag, N=Zr, Hf, Sn) have been used to enhance d_{33} and electromechanical strain (S) with d_{33} and S improved to >400 pC/N and 0.46% at 3 kV/cm, respectively.^{43–49} Despite promising d_{33} and S values there are still a number of concerns. For Sb doping, $T_{\rm C}$ is reported to decrease alarmingly and de-poling may jeopardize some applications. More importantly, however, to our knowledge, the temperature dependence of the piezoelectric properties is rarely reported. It is likely that the supposed MPB in many KNN-based compositions is not temperature independent and therefore phase coexistence and high d_{33}/S are not maintained during high-temperature applications or at high field cyclic loading in which temperature is likely to increase. Given the above concerns, this contribution presents the composition and temperature dependence of the piezoelectric properties $(1-x)(K_{1-y}Na_y)NbO_3-x(Bi_{1/2}Na_{1/2})$ ZrO₃ (KNyN-xBNZ) lead-free ceramics in which LiSbO₃ dopants have been excluded to maintain a high $T_{\rm C}$.

II. Experimental Procedure

The KNyN-xBNZ ceramics with compositions of (1-x) ($K_{1-y}Na_y$)NbO₃-x(Bi_{1/2}Na_{1/2})ZrO₃ (x = 0, 0.02, 0.03, 0.04, 0.045, 0.05, 0.06, 0.08, and y = 0.6) and (x = 0.045, and y = 0.4, 0.45, 0.49, 0.53, 0.56, 0.6, 0.65, and 0.7) were prepared using a conventional solid-state reaction. Raw materials, including Na₂CO₃ (99.9%, Fisher Scientific), K_2 CO₃

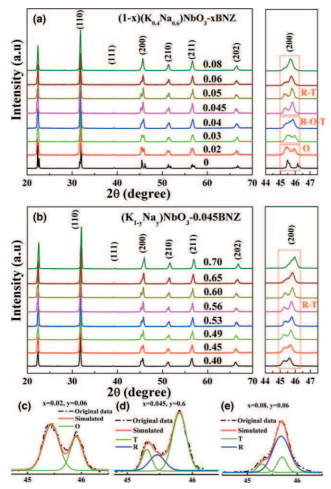


Fig. 1. Room temperature XRD patterns of KNyN-xBNZ (a) KN0.6N-xBNZ and (b) KNyN-0.045BNZ; amplified XRD patterns simulated by Gaussian for (c) KN0.06N-0.02BNZ (d) KN0.06N-0.045BNZ (e) KN0.06N-0.08BNZ.

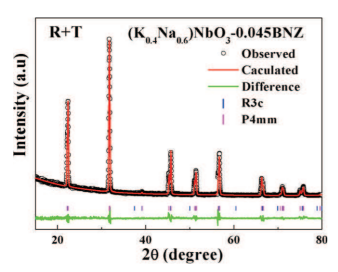


Fig. 2. Rietveld refinement analysis of KN0.6N-0.045BNZ using the GSAS+EXPGUI package. Two phases $R3c + P4 \ mm$ were used. $R_p = 6.2\%$, $R_{\rm wp} = 9.7\%$ and $\chi^2 = 1.88$.

(99.7%, Fisher Scientific), Nb₂O₅ (99.5%, Sigma-Aldrich), ZrO₂ (99%, Sigma-Aldrich), and Bi₂O₃ (99.9%, Sigma-Aldrich) were batched stoichiometrically according to the nominal compositions and ball-milled in isopropanol for 24 h. The dried mixed powders were calcined at 850°C for 6 h to synthesize the compound and then ball milled in isopropanol

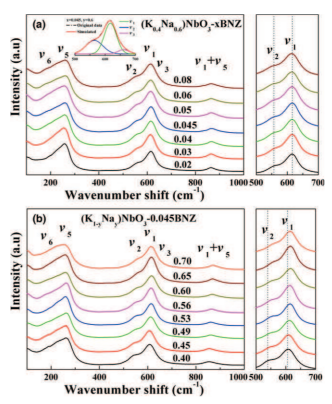


Fig. 3. Room temperature Raman spectra of KNyN-xBNZ (a) KN0.6N-xBNZ, inset shows the Raman amplified peaks around $600~{\rm cm}^{-1}$ simulated by Gaussian and (b) KNyN-0.045BNZ.

for 12 h. The calcined powders were mixed with a polyvinyl alcohol (PVA) binder solution, granulated and pressed into pellets with 10 mm in diameter. Following binder burnout at 550°C, the pellets were sintered in a sealed crucible with sacrificial powders at 1150°C–1230°C for 3–6 h.

The density of the sintered samples was measured by the Archimedes method. The relative density of all studied samples was higher than 95%. The phase structure of the sintered samples was studied using a Bruker D2 Phaser Xray powder diffraction (XRD) and in situ XRD performed for selected samples using a Siemens D5000 HTXRD in the temperature range 30°C-350°C. Ceramic morphology and microstructure was examined using an FEI Inspect F scanning electron microscope (SEM). Raman spectra were acquired from -150°C to 350°C using a Renishaw inVia Raman microscope. For electrical tests, sintered samples were electroded using fire-on silver paste, followed by the samples being poled in silicon oil at room temperature with an applied electric field of 30-40 kV/cm. Piezoelectric coefficient (d_{33}) was measured using a Piezotest PM300 d_{33} meter. Polarization hysteresis and strain-electric field behavior were determined using an aixACCT TF 2000 ferroelectric tester at a frequency of 1 Hz from room temperature to 160°C. The displacement data were synchronously captured by a laser interferometer. The temperature-dependent dielectric permittivity from 25°C to 600°C was measured using an Agilent 4184A multifrequency precision LCR meter. The planar electromechanical coupling factor (k_p) was determined from the resonance and antiresonance frequencies, which were measured using an Agilent 4294A Impedance/Gain-phase analyzer according to IEEE standards on piezoelectricity.

III. Results and Discussion

(1) Compositional Evolution of Structure–Property Relations

The room-temperature XRD patterns of KNyN-xBNZ in the 2θ range of $20^{\circ}-70^{\circ}$ are shown in Fig. 1. All peaks

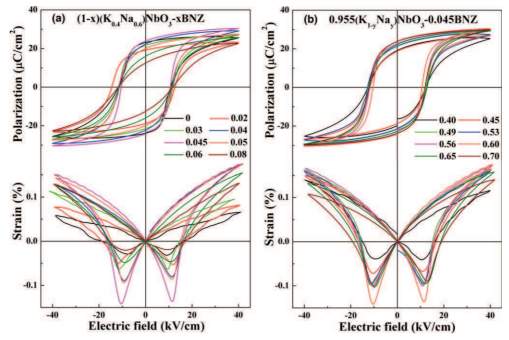


Fig. 4. High electric field bipolar polarization hysteresis and strain loops for KNyN-xBNZ (a) KN0.6N-xBNZ and (b) KNyN-0.045BNZ.

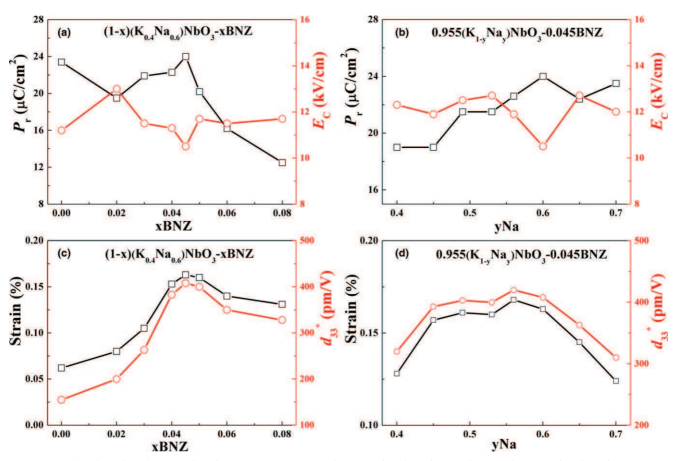


Fig. 5. Ferroelectric and strain properties of KNyN-xBNZ (a) P_r and E_C as a function of BNZ, (b) P_r and E_C as a function of Na, (c) S and d_{33}^* as a function of BNZ, (d) S and d_{33}^* as a function of Na.

could be attributed to perovskite phases, indicating that a stable solid solution was formed in the studied range. Diffraction peaks shifted to a lower and higher diffraction angle with increasing BNZ and Na concentration, respectively, consistent with differences in relative ionic radius of the dopant/substituent ions with the matrix composition.

Splitting of (200) peak at $\sim 2\theta = 45^{\circ}$ was used to determine the likely symmetry of compositions. In the expanded XRD patterns [Fig. 1(a)], O phase was dominant in KN0.6N-xBNZ with $x \le 0.02$, but R, O and T phases coexisted for compositions of $0.03 \le x \le 0.04$. As x increased further, ceramics with $0.045 \le x \le 0.05$ showed mixed R and T

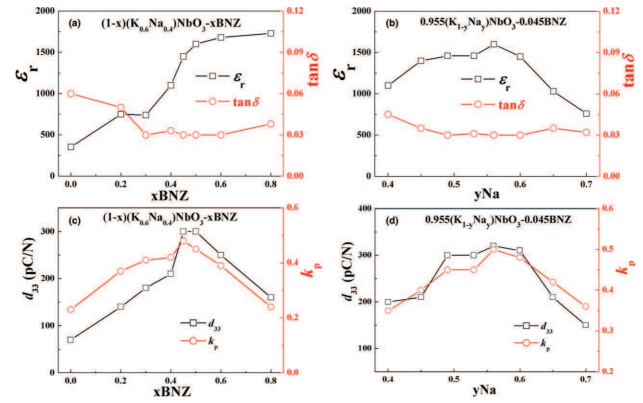


Fig. 6. Dielectric and piezoelectric properties of KNyN-xBNZ (a) ε_r and tanð as a function of BNZ, (b) ε_r and tanð as a function of Na, (c) d_{33} and k_p as a function of BNZ, (d) d_{33} and k_p as a function of Na.

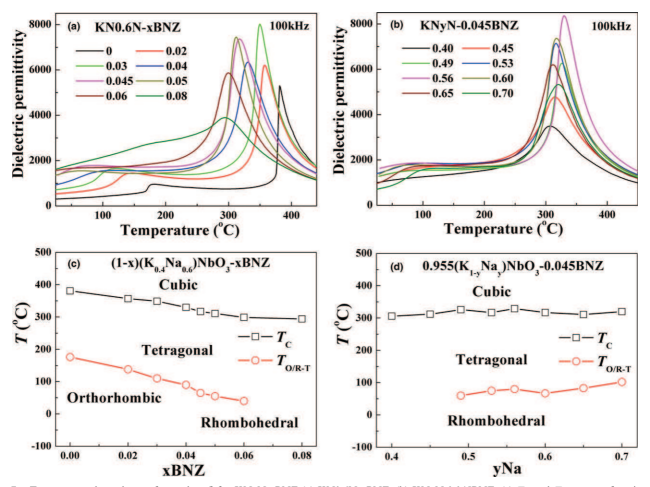


Fig. 7. Temperature dependence of ε_r and $\tan\delta$ for KNyN-xBNZ (a) KN0.6N-xBNZ, (b) KNyN-0.045BNZ, (c) T_C and $T_{O/R-T}$ as a function of BNZ, (d) (c) T_C and $T_{O/R-T}$ as a function of Na.

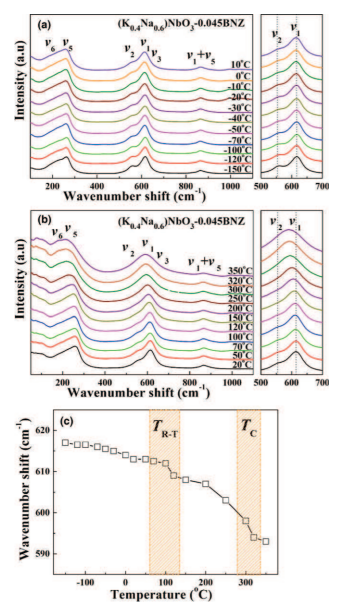


Fig. 8. In situ temperature dependence of Raman spectra for KN0.6N-0.045BNZ (a) $-150^{\circ}\text{C}-10^{\circ}\text{C}$, (b) $20^{\circ}\text{C}-350^{\circ}\text{C}$, (c) ν_1 peak shift as a function of temperature.

phases. For x > 0.05, a single peak emerged, suggesting that R phase dominated. For compositions in which x was maintained at 0.045 (KNyN-0.045BNZ), the splitting of (200)/(002) peaks gradually disappeared as y increased, suggesting the coexistence of R and T for a wide variation in Na:K ratio, Fig. 1(b). To confirm the phase evolution in KNyN-xBNZ as a function of composition, simulations were performed, Fig. 1(c-e), which are consistent with experimental data and also the interpretation by Wang et al. 43-50 Rietveld refinement analysis of the MPB composition KN0.6N-0.045BNZ was also performed using the GSAS+EXPGUI package, 53,54 where a two phases refinement (R3c + P4 mm) was used, as shown in Fig. 2. A good agreement between the observed and calculated patterns was obtained with $R_p = 6.2\%$, $R_{wp} = 9.7\%$ and chi-squared $(\chi^2) = 1.88$, indicating the coexistence of R (R3c, 32.5%) and T (P4 mm, 67.5%) phases in the MPB composition.

To further confirm phase evolution as a function of composition, room-temperature Raman spectra of KNyN-xBNZ ceramics were obtained as shown in Fig. 3. Among the full Raman active mode of KNN, v_1 , v_2 , and v_3 are stretching modes, v_5 and v_6 are bending modes of the NbO₆

octahedra.55 The strongest peak around 600 cm⁻¹ is related to the v₁ mode; whereas the left and right weaker peaks are assigned to the v_2 and v_3 mode, respectively, as indicated by the simulated data shown in the inset of Fig. 3(a). With increasing BNZ content, the v_1 and v_2 peaks were found to shift to lower wave number as shown in Fig. 3(a). On the other hand, the v_1 and v_2 peaks shift to higher wave number with an increase in Na content as shown in Fig. 3(b). The peak shift to a lower/higher frequency is due to an increase/ decrease in binding strength caused by the expansion/shrinkage of the distance between Nb5+ and its coordinated oxygen, resulting in the corresponding variation in crystal cell volume. 55,56 However, the clear but gradual symmetry changes observed by XRD were not apparent in the Raman spectra, as evidenced by the absence of the appearance of new symmetry related modes. Raman, however, is not only sensitive to the macroscopic symmetry but is influenced strongly by local distortions to the lattice. It is postulated that within the KNyN-xBNZ system there are compositional regions dominated by "pseudosymmetry" in a manner discussed for La-doped BiFeO₃ compositions by Khesro et al.⁵ and $Bi_{1/2}(K_{1-x}Na_x)_{1/2}TiO_3$ by Levin and co-workers.^{58,59} In these compositions, there are local perturbations in the average macroscopic symmetry more commonly associated with adjacent phases within the phase diagram, for example, short-range antipolar order in the paraelectric phase for Ladoped BiFeO₃. We propose a similar model for KNyNxBNZ based on the absence of new symmetry modes in Raman and the gradual changes in peak splitting in XRD but extensive transmission electron microscopy (TEM) work is required before this can be proved conclusively.

The SEM images for the surface of KNyN-xBNZ ceramics (not shown) revealed that the grain size of KN0.6N-xBNZ decreased with increasing BNZ content, from \sim 5 µm for x = 0 and to \sim 1 µm for x = 0.08 but the grain size of KNyN-0.045BNZ was insensitive to the K/Na ratio.

The high electric field bipolar polarization hysteresis and strain loops for KNyN-xBNZ as a function of BNZ and Na content are shown in Fig. 4, from which the remnant polarization (P_r), coercive field (E_C) and average electric field induced strain (S) as a function of BNZ and Na content can be obtained, Fig. 5. The normalized strain coefficient d_{33}^* , representing the average strain per unit of electric field, is calculated by, $d_{33}^* = S_{\text{max}}/E_{\text{max}}$, where E_{max} is the maximum electric field value, and $S_{\rm max}$ is the average value of the corresponding maximum strain. As shown in Fig. 5(a and b), with increasing BNZ and Na concentration in KNyN-xBNZ, $P_{\rm r}$ increased and then decreased after reaching a peak value, at which point the lowest $E_{\rm C}$ was obtained. The highest value of $P_r = 24 \,\mu\text{C/cm}^2$ and lowest value of $E_C = 10.5 \,\text{kV/cm}$ was achieved for x = 0.045, y = 0.6 at the MPB.^{60,61} The S and d_{33}^* values increased with increasing BNZ and Na content, reaching maximum values of 0.163%, 408 pm/V at x = 0.045, y = 0.6 and 0.168%, 420 pm/V at x = 0.045, y = 0.56, respectively, above which they reduced, Fig. 5(c and d). d_{33}^* is dominated by extrinsic effects, mainly domain wall motion, which the coexistence of R and T phases encourages at the MPB. 61-63

The dielectric and piezoelectric properties of KNyNxBNZ as a function of BNZ and Na content are shown in Fig. 6. The room-temperature permittivity (ε_r) of KN0.6NxBNZ increases with BNZ concentration, whereas it initially increases and then decreases as a function Na/K ratio, Fig. 6(a and b). tanð initially decreased with BNZ and Na concentration but then became stable for a broad range of compositions, Fig. 6(a and b). d_{33} and k_p both increased significantly with increasing BNZ and Na concentration as shown in Fig. 6(c and d), reaching a maxima of 310 pC/N and 48% for x = 0.045, y = 0.06, and 320 pC/N and 50% for x = 0.045, y = 0.056, respectively, and then decreasing for BNZ > 0.05 and Na > 0.6. As with d_{33}^* , d_{33} is optimized at the MPB largely through extrinsic

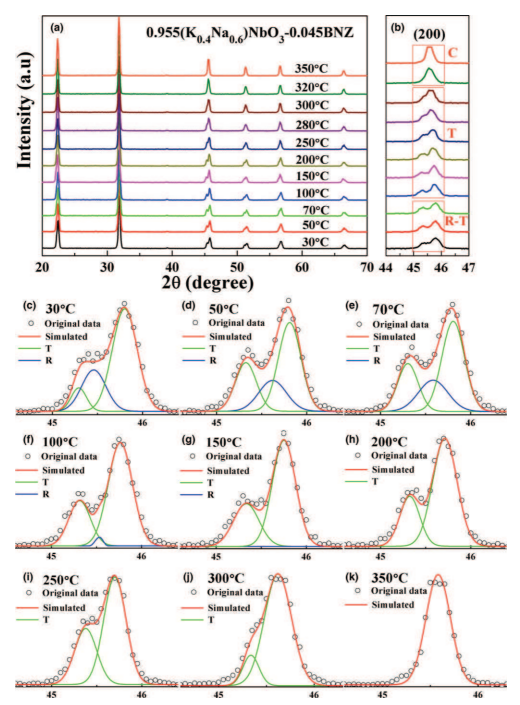


Fig. 9. In situ temperature dependence of XRD patterns for KN0.6N-0.045BNZ (a) $2\theta = 20^{\circ}-70^{\circ}$, (b) $2\theta = 44^{\circ}-47^{\circ}$; simulations of splitting of the {200} peaks with different temperatures (c) 30° C, (d) 50° C, (e) 70° C, (f) 100° C, (g) 150° C, (h) 200° C, (j) 300° C, (j) 300° C, and (k) 350° C.

contributions. The facile rotation of the polarization vector between states of similar free energy facilitated by phase coexistence, allows the domain walls to displace reversibly, resulting in a strong extrinsic contribution to the piezoelectric properties. ^{61,64}

(2) Evolution of Structure–Property Relations as a Function of Temperature

The temperature dependence of dielectric permittivity and loss for KNyN-xBNZ with different BNZ concentration and K/Na ratio is given in Fig. 7. It is well-known that KNN exhibits two dielectric peaks above room temperature, corresponding to the ferroelectric O-T phase transition at ~176°C ($T_{\text{O-T}}$) and the T-cubic (C) phase transition at ~381°C (Curie temperature, T_{C}). With increasing BNZ concentration, the

maximum $\varepsilon_{\rm r}$ at $T_{\rm C}$ ($\varepsilon_{\rm max}$) increased and $T_{\rm C}$ decreased continuously, meanwhile, $T_{\rm O-T}$ decreased and merged with the ferroelectric R-O phase transition ($T_{\rm R-O}$), resulting in coexistence of R and T phases at room temperature, Fig. 7(a,c), which is consistent with the XRD results (Fig. 1) and previous reports. 43–50 However, $T_{\rm C}$ remained at ~310°C and $T_{\rm R-T}$ increased with increasing Na concentration, suggesting the K/Na ratio had little effect on the ferroelectric-paraelectric phase transition of KNyN-xBNZ and that its major influence is on lower temperature transitions. We note also that broad relaxor-like peaks are observed in all compositions if the BNZ concentrations exceeds, x=0.08. Relaxor behavior, however, is not the focus of this study and we note only that KNyN-xBNZ follows trends typically observed in ferroelectric to relaxor compositional transitions with a transformation from a clearly defined macroscopic symmetry

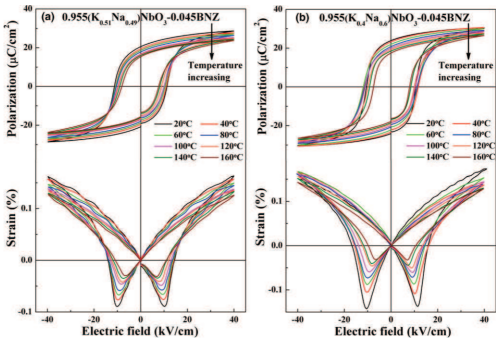


Fig. 10. In situ temperature dependence of high electric field bipolar polarization hysteresis and strain loops for two MPB compositions (a) KN0.49N-0.045BNZ and (b) KN0.6N-0.045BNZ.

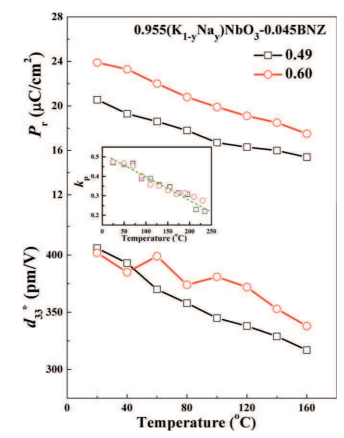


Fig. 11. $P_{\rm r}$ and d_{33}^* as a function of temperature for KN0.49N-0.045BNZ and KN0.6N-0.045BNZ, inset shows the temperature dependence of $k_{\rm p}$.

(x = 0) to pseudosymmetry (x > 0.08) that is associated with loss of long range ferroelectric order through chemical disorder that induces local competition/frustration between phases of similar free energy.

To clarify the structural evolution of KNyN-xBNZ as a function of temperature, *in situ* temperature dependence of Raman and XRD measurements for the selected MPB composition KN0.6N-0.045BNZ were performed, the results from which are plotted in Figs. 8 and 9. As shown in Fig. 8, the general trend is that the v_1 peak shifted to lower wave number and broadens, accompanied by a gradual fading of the v_2 peak. More specifically, the v_1 peak [Fig. 8(c)] revealed a gradual shift in frequency (wave number) up to $70^{\circ}\text{C}-100^{\circ}\text{C}$ after which there was a sharp change in frequency, coincident with the broad peaks in ε_{r} at the same temperature range in Fig. 7.

In situ high-temperature XRD data for KN0.6N-0.045BNZ is shown in Fig. 9 which focused on evolution of splitting of the $\{200\}$ peaks. Phase coexistence was apparent at room temperature as shown by broad multiple peaks that gradually disappeared to become a single peak at high temperatures in the cubic phase. To confirm the phase evolution as a function of temperature, simulations were performed, Fig. 9(c–k). As temperature increased, the coexistence of R-T phases remained up to 70° C– 100° C, after which the phase structure total transformed to a T and then C phase above $T_{\rm C}$, consistent with experimental $\varepsilon_{\rm r}$ and Raman spectroscopy data as function of temperature.

The in situ temperature dependence of high electric field bipolar polarization hysteresis and strain loops for two MPB compositions (KN0.49N-0.045BNZ and KN0.6N-0.045BNZ) are shown in Fig. 10, from which the $P_{\rm r}$ and d_{33}^* as a function of temperature were obtained, Fig. 11. As temperature increased, P_r and d_{33}^* for both MPB compositions decreased linearly, coincident with a similar temperature dependence of $k_{\rm p}$, as shown in the inset of Fig. 11. The piezoelectric effect in ferroelectric ceramics is attributed to both intrinsic (lattice deformation) and extrinsic (domain wall motion) contributions. Normally, $P_{\rm r}$ reflects intrinsic contribution, and d_{33}^* and $k_{\rm p}$ contain both components. However, detailed knowledge of the domain morphology is required to fully appreciate how the extrinsic contributions is likely to vary as a function of temperature and composition and extensive transmission electron or piezoforce microscopy is required. Nonetheless, the data presented in this contribution clearly illustrates that KNN-BNZ suffers from extreme temperature

dependence which may inhibit its usage in high drive applications and high-temperature applications despite the attractive values of k_p and d_{33} at room temperature.

IV. Conclusion

In this work, KNN based lead-free ceramics with the composition $(1-x)(K_{1-y}Na_y)NbO_3-x(Bi_{1/2}Na_{1/2})ZrO_3$ were successfully prepared by a conventional solid-state route. With the increase in BNZ content, the phase structure of KNN gradually transformed from O to mixed R-T and then R phase, indicative of the construction of an R-T based MPB with $0.045 \le BNZ \le 0.05$ and $0.4 \le Na \le 0.7$. Optimum ferroelectric, piezoelectric, and strain properties were obtained for the MPB compositions with P_r , d_{33} , k_p , S, and d_{33}^* of 24 μ C/cm², 320 pC/N, 48%, 0.168%, and 420 pm/V, respectively. However, despite these attractive room-temperature values at the MPB, there is a dramatic decrease in properties with temperature indicating the MPB is not temperature independent unlike in PZT. Hence there are concerns about the viability of these compositions for high-temperature or high field applications.

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