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1 Introduction

Recently, many reports have concentrated on the development of lead-free piezoelectric materials due to the environmental concern about serious lead pollution caused by the high toxicity of lead oxide and its high volatility during processing. Many lead-free ceramics systems such as $BaTiO_3$ (BT), $Bi_{0.5}Na_{0.5}TiO_3$ (BNT), ($Bi_{0.5}K_{0.5}$)TiO₃ (BKT), and $K_{0.5}Na_{0.5}NbO_3$ (KNN) based materials, have been widely investigated due to their good electrical properties.^{1,2} Among these lead-free systems, the solid

Enhanced electrical properties of BNKT–BMN leadfree ceramics by CaSnO₃ doping and their bioactive properties†

Lalita Tawee,^a Narongdetch Boothrawong,^a Suwanan Thammarong,^b Phanrawee Sriprapha,^a Waraporn Boontakam,^a Denis Russell Sweatman,^{acd} Nopakarn Chandet,^e Chamnan Randorn^{de} and Gobwute Rujijanagul ^b*^{acdf}

There is an increasing interest in using piezoelectric materials, including lead-free piezoceramics for medical applications. As a result, more attention has been placed on investigating the biological properties of these materials. In this research experiment, electrical, mechanical, and biological properties of lead-free 0.99Bi_{0.5}(Na_{0.8}K_{0.2})_{0.5}TiO₃-0.01Bi(Mg_{2/3}Nb_{1/3})O₃ or 0.99BNKT-0.01BMN doped with CaSnO₃ (CSO) were investigated. The samples were synthesized by a modified solid-state reaction technique. X-ray diffraction (XRD) analysis showed that the samples presented a single perovskite phase. After adding CSO, electrical properties such as energy storage density (maximum $W_{rec} = 781$ mJ cm⁻³) and electro-strain properties (maximum $S_{max} = 0.3\%$) were improved at room temperature. Mechanical properties were also enhanced for the modified samples with maximum values for Vickers hardness (H_v) and elastic modulus (E_m) of 6.11 GPa and 135 GPa, respectively. Biological assays for cytotoxicity, indicated that the samples had high cell viability, while the simulated body fluid (SBF) test revealed a moderate apatite forming ability. The samples were then coated with hydroxyapatite to improve their apatite-forming ability. The SBF testing for the coated samples showed that the coated samples had high apatite-forming ability. The obtained results pointed to the possibility of ceramics being used for multifunction electrical devices at room temperature and biomaterial applications.

solution between BNT and BKT (BNKT) is of interest since it presents large spontaneous polarization, high permittivity, low dielectric loss, and a wide working temperature range.³ Normally, the BNKT solid solutions exhibited higher dielectric and piezoelectric properties near the morphotropic phase boundary (MPB) of the rhombohedral and tetragonal phases, at compositions ranging from x = 0.16 to 0.20 (for $[Bi_{0.5}(Na_{1-x}K_x)_{0.5}]$ TiO₃).⁴ Furthermore, the BNKT ceramics have a lower coercive field (E_c) as compared with the pure BNT which favors a poling process.^{2,5} The BNKT-based materials thus have a potential for many electronic device applications.^{1,2,6}

To achieve the high performance of BNK-based ceramics, many modified BNKT-based materials or their complex solid solutions have been synthesized and scrutinized. For example, the BNKT solid solution can be modified by modifier substances such as $Sr(Hf_{0.5}Zr_{0.5})O_3$ (improved the electrostrain),⁷ Bi(Mg_{0.5}Ti_{0.5})O_3 (enhanced normalized strain),⁸ Bi(Zn_{0.5}Ti_{0.5})O_3 (bettered both electro-strain and normalized strain),⁹ Ba(Zr_{0.04}Ti_{0.96})O_3 (enhanced piezoelectric properties),¹⁰ BiAlO₃ (improved electro-strain),¹¹ (Bi_{0.5}La_{0.5})AlO₃ (enhanced the electro-strain),¹² BiFeO₃ (bettered piezoelectric properties),^{13,14} Ag₂O (improved dielectric and piezoelectric properties),¹⁵ SrTiO₃ (improved the electro-strain),¹⁶ K_{0.5}Na_{0.5}NbO₃ (bettered electro-strain and normalized strain),¹⁷ and Bi(Mg_{2/})

[&]quot;Department of Physics and Materials Science, Faculty of Science, Chiang Mai University, Chiang Mai 50200, Thailand. E-mail: rujijanagul@yahoo.com

^bState Key Laboratory of Structure Analysis for Industrial Equipment, Department of Engineering Mechanics, Dalian University of Technology, Dalian 116024, China

^cOffice of Research Administration, Chiang Mai University, Chiang Mai 50200, Thailand

^dMaterials Science Research Center, Faculty of Science, Chiang Mai University, Chiang Mai 50200, Thailand

^eDepartment of Chemistry, Faculty of Science, Chiang Mai University, Chiang Mai 50200, Thailand

^fMultidisciplinary and Interdisciplinary School, Chiang Mai University, Chiang Mai 50200, Thailand

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 $_3Nb_{1/3})O_3$ (BMN) (exhibited good piezoelectric and electromechanical properties).^3

However, it should be noted that BMN can be used as a modifier for several Bi-based piezoelectric ceramic systems such as BNKT-BMT (improved electro-strain),¹⁸ BT-BMN (exhibited good dielectric/temperature/frequency dependent stability and good energy storage density),¹⁹ BNT-BMN (presented good energy storage properties),²⁰ Bi_{1.05}FeO₃-BT-BMN (showed large strain with low hysteresis strain),²¹ and Ba_{0.55}Sr_{0.45}TiO₃-BMN (improved breakdown strength and energy storage behavior).²²

Lately, CaSnO₃ (CSO) has attracted a growing interest because of its various applications in objects, such as gas sensors,²³ capacitor components,²⁴ anode materials for lithiumion batteries,²⁵ and catalysts.^{26–28} Moreover, CSO has been used as a modifier for some lead-free piezoelectric ceramics such as BT–CSO (improved dielectric and piezoelectric behaviours),²⁹ NaNbO₃–CSO (improved antiferroelectric behaviour),³⁰ and BiFeO₃–CSO (enhanced ferroelectric and piezoelectric properties).³¹ Since BNKT ceramics present many good properties and BMN as well as CSO modifiers, have positive effects on many properties of lead-free piezoelectric materials, it is interesting to synthesize the BNKT-based ceramics modified with BMN and CSO.

As of lately, there has been a higher appeal in the investigation of using piezoelectric materials in medical applications.^{32,33} Some lead-free piezoceramics have been proposed to combine with some bio-ceramics to search for a potential biocompatible material and their applications.^{34,35} Therefore, the property investigation of the biological properties of leadfree piezoceramics is essential, especially for medical purposes. In the present work, BNKT–BMN ceramics modified with CSO were synthesized and their mechanical and electrical properties were investigated. Furthermore, bio-properties of the researched ceramics were also explored to determine their potential for use in biomedical applications.

2 Experimental

A solid solution (1 - x)[0.99BNKT-0.01BMN]-xCSO ceramics (with x = 0.00, 0.02 and 0.04) were fabricated via a modified conventional solid-state reaction method, using high-purity raw materials of Bi2O3, Na2CO3, K2CO3, TiO2, MgO, Nb2O5, CaCO3 and SnO2. The BNKT, BMN, and CSO were calcined separately by stoichiometrically weighing the metal oxides for their composition and then mixed by a ball-milled technique for 24 h in an ethanol solution. The obtained products were dried at 120 °C in an oven overnight. The dried (and mixed) raw powders for BNKT, BMN, and CSO were separately calcined at 900 °C, 700 °C and 1200 °C for 2 h, respectively. After that, the BNKT, BMN and CaSnO₃ calcined powders were then weighed, mixed, and dried again to produce the mixed powders of (1 - x)[0.99BNKT-0.01BMN]-xCSO (x = 0.00, 0.02 and 0.04). The calcined powders were crushed and sieved to which polyvinyl alcohol (PVA) binder was afterward added. Next, the powders were uniaxially pressed into discs 10 mm in diameter. The binder removal process was carried out by heating the green pellets to 500 °C for 1 h and then sintering them at 1125 °C for

2 h dwell time with a heating/cooling rate of 5 $^{\circ}$ C min⁻¹ in closed alumina crucibles. To minimize the loss of the volatile elements, the green pellets were embedded in the powder of the same composition. The bulk density and apparent porosity of sintered pellets were measured using the Archimedes' method.

The crystal structure and phase formation of powders and ceramics were investigated by X-ray diffraction technique (XRD, PANalytical, X'Pert Pro MPD). A Raman scattering technique (T6400 JY, Horiba Jobin Yvon) was utilized to confirm the XRD result. A scanning electron microscope (FE-SEM, JEOL JSM-6335F) was used to determine the microstructure of the samples. The grain size of the ceramics was measured by using the linear intercept method. For electrical property measurements, the silver paste was applied on both sides of sintered pellets and then fired at 600 °C for 20 min. Dielectric constant and dielectric loss were measured in the temperature range 30-500 °C at different frequencies from 1 kHz to 1 MHz using a 4192A LCR meter connected to a high temperature furnace. The ferroelectric properties were carried out using a ferroelectric tester (Radiant Technologies, Inc. with Precision 10 kV HVI-SC). Remnant polarization (P_r) , maximum polarization (P_{max}) , and coercive field (E_{c}) values were determined from the hysteresis loops. The mechanical properties, including hardness, elasticity modulus $(E_{\rm m})$, and fracture toughness $(K_{\rm IC})$ of the material were measured using a hardness tester (SMV-1000).

To check the cell viability, the ceramics powder was tested by a MTT assay by using a model of the primary dermal fibroblast PCS-201-012 cell line (Manose Health and Beauty Research Center). Dulbecco's Modified Eagle Medium (DMEM) with 10% fetal bovine serum (FBS) and 1% penicillin/streptomycin was used to culture the suspended PCS-201-012 cell line in a 96-well plate (5000 cells per well). All cells well plates were retained at $37\pm0.1~^{\rm o}{\rm C}$ in a CO $_2$ incubator with 5% CO $_2$, and 100% relative humidity. The cultures and mediums were switched every week and twice a week, respectively. The cells were treated with fine sterilized ceramics powder with 190 ml of 0.1, 1, 10, 100, and 1000 μ g ml⁻¹ of concentrations for 24 h. The positive control was served by sodium lauryl sulfate (SLS). The optical density (OD) of the MTT assay was calculated by a microplate reader at the wavelength of 510 nm, and the average of three readings was counted. The bioactivity of a sample was determined utilizing a simulated body fluid (SBF) test. The samples were ultrasonically cleaned in ethyl alcohol for 30 minutes before drying overnight. The samples were immersed in a pH 7.4 SBF solution and kept in a temperature-controlled chamber (37 \pm 0.5 °C). The apatite layers that developed on the material were investigated using an SEM.

3 Results and discussion

3.1 Phase formation

The X-ray diffraction patterns of the (1 - x)[0.99BNKT-0.01BMN]-xCSO are shown in Fig. 1(a). All ceramics exhibited a single phase of perovskite structure without secondary phase, within the resolution limit of the XRD technique. This suggests the formation of the solid solution occurred after the

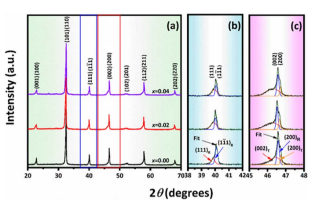


Fig. 1 X-ray diffraction patterns of (1 - x)[0.99BNKT-0.01BMN] - xCSO ceramics: (a) $2\theta = 20^{\circ}-70^{\circ}$, (b) $2\theta = 38^{\circ}-42^{\circ}$, and (c) $2\theta = 45^{\circ}-48^{\circ}$.

processing. The tolerance factor (t) of the ceramics system was calculated, as presented in Fig. S1 (in the ESI⁺) and the obtained values were in the range 0.9742-0.9762. Normally, the perovskite structure is stable within the range of 0.880 < t < 1.090.³⁶ Therefore, the *t*-values for the studied samples lie well within the *t*-value limit, indicating the materials belong to a stable perovskite structure.37 It should be noted that the positions of all peaks tend to shift toward lower 2θ angles when compared to the x = 0.00 samples. For example, the (110) peak exhibited a shift to a lower angle when the CSO content was increased. This corresponds with the change in the *t* value associated with the CSO content. Therefore, the lattice distortion in the modified samples occurred, which may be due to the diffusion of Ca²⁺ and Sn⁴⁺ ions into the lattices of the base ceramics. Similar evidence for peak shifting behaviour was also observed in a previous work.³⁸ For the x = 0.0 samples, there was a peak splitting at $2\theta \sim 40^{\circ}$ ((111)/(111) peaks) and broad ambiguity at 46° ((002)/(200) peaks), which indicated a mixture phase of rhombohedral and tetragonal phases. It should be noted that, a slight splitting of (002)/(200) tetragonal peaks was observed with increasing CSO content.³⁹⁻⁴³ Furthermore, the peak splitting of $(111)/(1\overline{1}1)$ peaks slightly merged into a single peak (Fig. 1(b) and (c)). This behaviour matches the trend of calculated tetragonality (c/a) of the samples slightly increasing with increasing CSO content, which indicated more tetragonality (Fig. S1(a) in the ESI[†]) with increasing CSO content. This result agrees with the previous studies for the phase transformation.44

The Raman spectra of (1 - x)[0.99BNKT-0.01BMN]-xCSO ceramics measured at room temperature (RT) over the range of 100–1000 cm⁻¹ are shown in Fig. 2. There are four main regions of Raman spectra identified by: (1) a vibration mode at ~120 cm⁻¹, (2) ~270 cm⁻¹, (3) ~450–650 cm⁻¹, and (4) peak at >700 cm⁻¹, respectively. Based on the reports from previous studies, the first mode observed between 100 and 150 cm⁻¹ is related to the vibration of A-site cations in the perovskite structure, influenced by A-site doping.^{45,46} The peak around 270 cm⁻¹ is associated with Ti–O vibrations at the B-site.⁴⁶ The peak at 450–620 cm⁻¹ can be related to the vibration of TiO₆ octahedra mode.^{47,48} It has been suggested from previous experiments for the work done on BNKT-based ceramics that

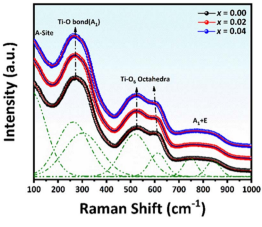


Fig. 2 Raman spectra of the (1 - x)[0.99BNKT-0.01BMN]-xCSO ceramics at RT.

the splitting peak in the vibration mode of Ti–O and TiO₆ octahedra mode indicates a coexistence of rhombohedral (*R3c*) and tetragonal (*P4bm*) phases.^{48,49} In this work, there was a shoulder for the peak around 270 cm⁻¹ and a splitting peak TiO₆ octahedra mode, which indicated the coexistence of the two phases. Finally, the broad peak observed in the fourth region was assigned to be the A₁ (longitudinal optical) and E (longitudinal optical) overlapping bands.⁴⁸ However, the presence of these broader peaks was related to the presence of oxygen vacancy.⁵⁰ Thus, the obtained Raman data in the current study is consistent with the XRD result, which confirms the coexistence of the phases for all compositions.

3.2 Structural analysis

Figs. 3(a)–(d) shows the SEM images of the surface samples. The SEM image illustrates that all ceramics were densely sintered

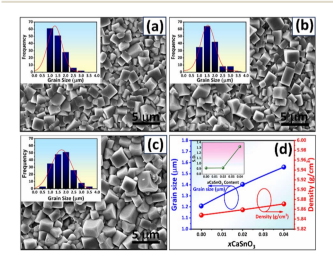


Fig. 3 SEM micrographs of (1 - x)(0.99BNKT-0.01BMN)-xCSO sintered ceramics: (a) x = 0.00, (b) x = 0.02, (c) x = 0.04, and (d) grain size with their corresponding density value as a function of CSO content. Insets (a)–(c): grain size distributions of the samples. Inset (d): S.D. vs. CSO content.

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and contained closely compacted grains with low surface porosity. This corresponds with the density value (5.85-5.87 g cm⁻³), where the density slightly increased with CSO content (Fig. 3(d)). Furthermore, all samples exhibited equiaxed grain shape and uniform grain morphology with well-defined grain boundaries. For the x = 0.00-0.02 samples, a cubic-like shape with a sharp edge was observed, while the x = 0.04samples presented a rounder edge grain. The CSO additive produced a slight increase in the average grain size, *i.e.*, the average grain size increased from 1.21 μ m for the x = 0.00samples to 1.56 μ m for the *x* = 0.04 samples (Fig. 3(d)). To check the degree of uniformity of the grain size, grain size distributions of the samples were plotted and a standard deviation (S.D., μ m) of the grain size was determined as presented in the insets of Fig. 3(a)-(d). All grain size distributions presented a monomodal normal distribution, and the S.D. value slightly increased with CSO content (inset of Fig. 3(d)). This confirms that all samples had a uniform grain size, which revealed a good sintering processing.

3.3 Dielectric properties

The temperature dependence of dielectric constant (ε_r) and dielectric loss (tan δ) measured at different frequencies of poled samples are shown in Fig. 4(a)-(c). The corresponding dielectric properties are also given in Table S1 (in the ESI⁺). Two dielectric anomaly peaks were observed for the dielectric curves. The first anomaly peak (near 100 °C) indicates the presence of a relaxor ferroelectric characteristic, which is typically caused by thermal evolutions of discrete polar nanoregions.51 The second, denoted as $T_{\rm m}$ and developing at approximately 300 °C, is associated with the maximum dielectric constant ($\varepsilon_{r,max}$). As suggested by many previous studies, the temperature at the peak of tan δ -T curve of the poled sample at a temperature less than 100 °C can be defined as the ferroelectric to relaxor phase transition temperature $(T_{\rm F-R})$.^{52,53} For BNKT-based ceramics, the $T_{\rm F-R}$ has been proposed to be related to an evolution of the initially coexisting R3c and P4bm PNRs, and $T_{\rm m}$ has a relevance to the

P4bm PNRs that entirely transform from the *R3c* PNRs with increasing the temperature.^{54,55} In this work, the $T_{\rm F-R}$ could not be observed, because there was no clear sharp peak in the tan δ -*T* curve. However, the broad or hump curve at the temperature around $T_{\rm F-R}$ of tan δ -*T* curve tended to shift to a lower temperature with increasing CSO content.

The Burns temperature $(T_{\rm B})$, which denotes the temperature from which the dielectric constant starts to deviate from the Curie-Weiss law, was also evaluated and the result is presented in Fig. S2 (in the ESI^{\dagger}). The trend of $T_{\rm B}$ value was similar to the trend of $T_{\rm m}$, *i.e.*, it decreased with CSO content. Yet, the $\Delta T (=T_{\rm B})$ $(-T_m)$ value (Table S1 in the ESI[†]) increased with the increasing CSO content, indicating the diffused phase transition behaviour was also enhanced.⁵⁶ In addition, the dielectric peak at $T_{\rm m}$ became broader, and the dielectric constant value shifted down. This was evident with the decrease in the degree of frequency dispersion of the dielectric constant, and the intensity of frequency dispersion $\Delta \varepsilon_r$ (= $\varepsilon_{r,1 \text{ kHz}} - \varepsilon_{r,1 \text{ MHz}}$) decreased with increasing CSO content (Table S1†).51,57 These results can affect electrical properties such as ferroelectric and piezoelectric properties (next sections). The observed decrease in transition temperatures as the CSO concentration increased revealed that these transition temperatures can be controlled by the CSO content. It should be noted that, the tan δ value slightly changed with CSO, where its values at RT and $T_{\rm m}$ were 0.0336–0.0415 and 0.0303-0.0398, respectively. This is beneficial for the development of the ceramics for capacitor applications (Table S1 in the ESI[†]). Thus, the dielectric properties of the studied system were controlled by the change in their composition.

3.4 Ferroelectric properties

Fig. 5 displays the *P*–*E* hysteresis loops of the samples at different CSO contents and applied electric fields. Plots of P_{max} , P_{r} , and E_{c} as a function of CSO content are also presented in Fig. 5(d). The corresponding ferroelectric properties are listed in

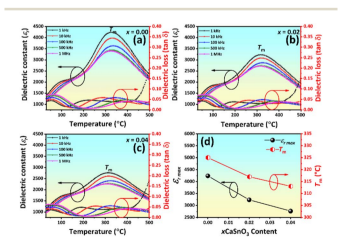


Fig. 4 (a)–(c) Temperature dependence of dielectric constant (ε_r) and dielectric loss (tan δ) of the ceramics as a function of CSO content and (d) plots of $\varepsilon_{r,max}$ and T_m as a function of CSO content.

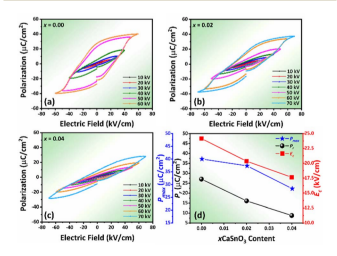


Fig. 5 Polarization-electric field (*P*–*E*) hysteresis loops of (1 - x) [0.99BNKT–0.01BMN]–xCSO ceramics at RT and different electric fields: (a) x = 0.00, (b) x = 0.02, (c) x = 0.04, and (d) plots of P_{max} , P_{r} , and E_{c} as a function of CSO content.

Table S2 (in the ESI[†]). For the x = 0.00 ceramics, the shape of the *P*–*E* loop indicated a ferroelectric order behaviour. However, the shape of the *P*–*E* loop transformed into a constriction loop that became slimmer with increasing CSO content. The *P*_{max}, *P*_r and *E*_c values also tended to decrease with CSO content. This indicates that the long-range ferroelectric order in the unmodified ceramics was disturbed by the CSO and transformed to be into the ergodic relaxor (ER) phase, where the ER phase can transform into a ferroelectric (FE) phase by an external electric field.⁴⁸ In the current experiment, the energy storage density (*W*_{rec}) and energy storage efficiency (η), were calculated from the *P*–*E* loop data by using the following equations:³⁸

$$W = \int_0^{P_{\text{max}}} E \mathrm{d}P,\tag{1}$$

$$W_{\rm rec} = \int_{P_{\rm r}}^{P_{\rm max}} E dP \tag{2}$$

$$\eta (\%) = \frac{W_{\text{rec}}}{W_{\text{rec}} + W_{\text{loss}}} \times 100$$
(3)

where W_{loss} , is energy loss density ($W_{\text{loss}} = W - W_{\text{rec}}$) and *E* is the external electric field. At RT, the unmodified ceramic showed $W_{\text{rec}} = 320 \text{ mJ cm}^{-3}$ and $\eta = 19.68\%$. The W_{rec} and η values increased with increasing CSO content and reached the maximum values of 781 mJ cm⁻³ and 42.99%, respectively for the x = 0.02 samples. This improvement matched the $\Delta P (=P_{\text{max}} - P_{\text{r}})$ which was the maximum for the x = 0.02 samples.

In addition, this improvement can be related to the formation of the constrictive *P*–*E* loop for this composition, which indicated the formation of the mixed FE and ER phases.⁴⁸ It should be remarked that the $W_{\rm rec}$ value for the 0.02 samples at RT is higher than many BNKT based piezoelectric ceramics (at RT, Table 1). To compare the ability of the $W_{\rm rec}$ value under a low working electric field at RT, a normalized energy density ($W_{\rm rec}/E$ = energy storage density/applied electric field) was determined. A comparison of $W_{\rm rec}/E$ value of the current work with other previous works for lead-free ceramics is presented in Table 1.^{59–68} The $W_{\rm rec}/E$ value for the 0.02 samples is considered high when compared with many previous studies.

The influence of the applied electric field on the $W_{\rm rec}$ and η values at RT is depicted in Fig. 6(a) and (b), respectively. The $W_{\rm rec}$ value tended to increase with the electric field, especially for the modified sample due to ΔP increasing with the electric field (Fig. S3 in the ESI[†]). This also consists with the work done by Dai et al. for BaSrTiO₃-Bi(Zn_{1/2}T_{1/2})O₃ ceramic system.⁶⁹ However, the η value decreased with the electric field because $W_{\rm loss}$ increased with applied electric field at a higher rate when compared to the W_{rec} value (see Fig. S2 in the ESI[†]). The *P*-*E* hysteresis loop at different temperatures of the samples was plotted and results are shown in Fig. S4 (in the ESI[†]). Plots of $W_{\rm rec}$ and η values as a function of temperature are shown in Fig. 6(c) and (d), respectively. The related values are also listed in Table S2 (see the ESI[†]). The trend of constrictive loop formation for each composition at high temperatures, confirmed the trend of $T_{\rm F-R}$ value that shifted down with increasing CSO content, *i.e.*, hump curve of $\tan \delta - T$ curve shifted down with CSO. In addition, the W and η values, as calculated from the *P*-*E* loop, increased with increasing temperature, and reached the

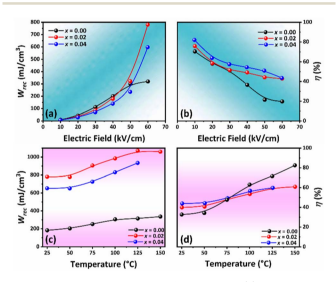


Fig. 6 Energy storage properties of the samples: (a) $W_{\rm rec}$ vs. electric field, (b) η vs. applied electric field, (c) $W_{\rm rec}$ vs. temperature, and (d) η vs. temperature.

Table 1 Comparison of the energy storage performance at RT between the (1 - x)[0.99BNKT-0.01BMN]-xCSO (with x = 0.02) ceramics and other BNKT-based ceramics

Materials	W (mJ cm ⁻³)	E (kV mm ⁻¹)	W/E (µC mm ⁻²)	η (%)	Ref.
$(1 - x)[0.99BNKT-0.01BMN]-xCaSnO_3, x = 0.02$	781	6	0.13	43.00	This work
$Bi_{0.5}(Na_{0.82}K_{0.18})_{0.5}Ti_{1-x}(Mg_{1/3}Nb_{2/3})_xO_3, x = 0.02-0.045$	650	7	0.09	34.15	59
$(1 - x)Bi_{0.50}(Na_{0.80}K_{0.20})_{0.50}TiO_3 - xBa_{0.90}Ca_{0.10}Ti_{0.90}Zr_{0.10}O_3$	370	4	0.09	50.00	60
Bi _{0.5} (Na _{0.8} K _{0.2}) _{0.5} (Ti _{0.96} Sn _{0.04})O ₃	400	5	0.08	41.70	61
$Bi_{0.5}(Na_{0.80}K_{0.20})_{0.5}TiO_3$ - $SrZrO_3$ - Eu^{3+}	750	15	0.05	80.00	62
$(1 - x)(0.75Bi_{0.5}Na_{0.5}TiO_3 - 0.25Bi_{0.5}K_{0.5}TiO_3) - xBiAlO_3, x = 0.06$	710	7	0.10	71.30	63
$(1 - x)[Bi_{0.5}(Na_{0.80}K_{0.20})_{0.5}TiO_3 - 0.03(Ba_{0.70}Sr_{0.03})TiO_3] - xBa(Fe_{0.5}Ta_{0.5})O_3, x = 0.03$	490	5	0.10	60.70	64
$(1 - x)[Bi_{0.5}(Na_{0.8}K_{0.2})_{0.5}TiO_3] - x[Ba_{0.844}Ca_{0.156}(Zr_{0.096}Ti_{0.904})O_3], x = 0.075$	430	5	0.09	50.70	65
$(1 - x)[Bi_{0.5}(Na_{0.4}K_{0.1})TiO_3]-xBaTiO_3, x = 0.04$	300	4.5	0.07	24.6	66
Bi _{0.485} (Na _{0.388} K _{0.097})Ba _{0.021} Sr _{0.009} TiO ₃ , sintered at 1125 °C	270	4	0.07	28.10	67
(1 - x)[Bi _{0.5} (Na _{0.4} K _{0.1})TiO ₃]-xBi ₄ Ti ₃ O ₁₂ , $x = 0.15$	120	4.63	0.03	0.17	68

maximum value of 1071 mJ cm⁻³ and 59.60%, respectively for the *x* = 0.02 composition at 150 °C.

3.5 Electro-strain behaviour

Fig. 7 presents the electro-strain properties of the samples. The maximum strain values (S_{max}) and the negative strain (S_{neg}) , where S_{neg} is the difference between zero-field strain and the lowest strain, are displayed in Fig. 7(e), and the normalized strain coefficient $(d_{33}^* = S_{\text{max}}/E_{\text{max}})$.⁷⁰ are depicted Fig. 7(f). The related data are also presented in Table S3 (in the ESI[†]). With increasing CSO content, the S_{neg} value approached zero and the $S_{\rm max}$ reached its maximum value (0.3%) for the x = 0.02samples. The d_{33}^* value increased with CSO content and reached the maximum value for the x = 0.02 sample, then decreased for the x = 0.04 samples (Fig. 7(f)). In the present experiment, the significant enhancement in S_{max} can be described by the fact that the addition of CSO decreases the FE ordering and finally transformed into an ER state at an optimum between FE (nonergodic relaxor) and ER portion. However, upon the application of an electric field, the ER at zero field can easily be transformed into a long-range FE phase, and thus a large strain response can be achieved at around the critical phase composition.71-73 This result is consistent with the transformation of the P-E hysteresis-shaped loop from a normal ferroelectric loop for the x = 0.00 samples to a constriction loop for the x = 0.02 samples. However, the decrease of S_{max} for the x = 0.04 samples may be due to the higher degree of ergodicity for this composition.72

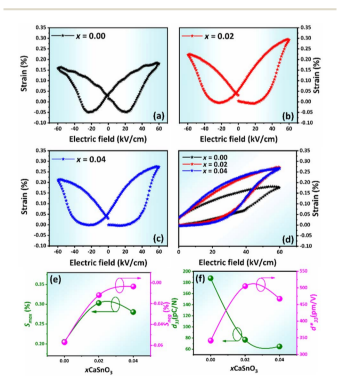


Fig. 7 Electro-strain and piezoelectric properties of the samples at different CSO contain and RT: (a–c) bipolar *S–E* loop *versus* CSO content for the x = 0.00-0.04 samples, (d) unipolar *S–E* loop *versus* CSO content, (e) *S*_{max} and *S*_{neg} *versus* CSO content, and (f) *d*₃₃ and *d*^{*}₃₃ value as a function of CSO content.

The low field d_{33} value as a function of CSO content is also shown in Fig. 7(f). The d_{33} value tended to decrease with CSO, as expected, due to the additive deteriorating the ferroelectric ordering and increasing the degree of ergodicity, which aligns with previous work.⁷⁴ This evidence can be roughly described by the thermodynamic theory of ferroelectrics, which follows the equation: $d_{33} = 2\varepsilon_{33}\varepsilon_0 Q_{11}P_r$, where ε_{33} is the dielectric constant, and Q_{11} is an electrostrictive coefficient (which is constant for perovskite materials).⁵

3.6 Mechanical properties

Normally, the practical uses of actuators or other applications such as biomedical applications are significantly influenced by their mechanical performance. In the current study, the mechanical properties of the samples, including Vickers (H_V) and Knoop (H_K) hardnesses, elastic modulus (E_m), and fracture toughness (K_{IC}) were determined. Plots of H_V and H_K values as a function of CSO content are shown in Fig. 8(a) and their corresponding values are listed in Table S4 (in the ESI†). In the current work, the H_V and H_K values were calculated *via* eqn (4) and (5).

$$H_{\rm V} = \frac{(1.854)P}{d^2} \tag{4}$$

$$H_{\rm K} = \frac{(14.23)P}{d^2}$$
(5)

where P = load and d = half-length of the long diagonal.⁷⁵⁻⁷⁷ The results indicated an increase in H_V and H_K values as the CSO content increased. Normally, the increases in H_V and H_K values can be attributed to many factors, such as density and grain size, *i.e.*, a higher density and lower grain size often produce a higher Vickers and Knoop's values. In the current work, there was a slight change in grain size. Therefore, the H_V and H_K values of the samples were predominantly affected by the increased density associated with the higher CSO content, a result stemming from the altered composition.

Plots of $E_{\rm m}$ and $K_{\rm IC}$ values as a function of CSO content are shown in Fig. 8(b). The $E_{\rm m}$ and $K_{\rm IC}$ values were also calculated using the following formulas (6) and (7):^{77,78}

$$E_{\rm m} = \frac{\alpha H_{\rm K}}{\left[\left(\frac{b}{a}\right) - \left(\frac{b'}{a'}\right)\right]} \tag{6}$$

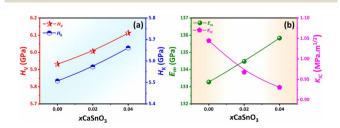


Fig. 8 Plots of (a) H_V and H_K as a function of x content and (b) E_m and K_{IC} values as a function of CSO content.

$$K_{\rm IC} = 0.016 \left(\frac{E}{H_{\rm V}}\right)^{1/2} \left(\frac{P}{c^{3/2}}\right)$$
 (7)

where α is the Marshall constant (~0.45), b/a is the length of the $H_{\rm K}$ tip mark with short diagonal/length of the $H_{\rm K}$ tip mark with long diagonal (0.14), b'/a' is the length of the $H_{\rm K}$ indentation with short diagonal/length of the $H_{\rm K}$ indentation with long diagonal, and c is the radial, or the length, from the center of the indentation imprint to the crack tip. The $E_{\rm m}$ value had a similar trend to that of $H_{\rm V}$ and $H_{\rm K}$, but the $K_{\rm IC}$ value had the opposite trend. According to the results, the $H_{\rm V}$, $H_{\rm K}$, and $E_{\rm m}$, values of the studied ceramics were enhanced due to the compositional tuning associated with varying concentrations of CSO addition.

3.7 Biological properties

Recently, there has been an increasing interest in using piezoelectric materials, including lead free piezo ceramics for medical applications.^{34,58} Thus, the details of bio-properties of piezoelectric materials have become more important. In this work, some bio-properties of the studied materials were investigated to search for potential uses in medical applications.

3.7.1 *In vitro* cell viability assay. Cell viability assays can be used to detect whether cells survive or die after exposure to a drug or chemical. In this experiment, the MTT test, which is a calorimetric technique, was employed for the cell viability assays. The MTT test was carried out on skin fibroblast cells, with sodium lauryl sulfate (SLS) as a positive control. The cell viability assay results are shown in Fig. 9. Furthermore, the percentage of cell viability was calculated from the ratio of absorbance of the cells incubated with the sample suspensions to that of the cells incubated with culture medium control using the following formula.⁷⁸

Cell viability (%) =
$$\frac{\text{absorbance}_{\text{ceramics}}}{\text{absorbance}_{\text{control}}} \times 100$$
 (8)

Comparative histogram plots between cell viability and concentration of positive control and treated-ceramics groups are shown in Fig. 10. The percentage cell viability for all samples was higher than 90% of all concentrations despite increasing of the x value.^{79,80} Thus, there were non-cytotoxic effects to cells compared with percentages shown by cells cultured in the SLS (positive control). These small differences of cell viability of all samples were not statistically significant. This indicates that ceramics were non-cytotoxic and beneficial for cell proliferation at an appropriate dosage. Comparing the obtained results with the standard sample result, the studied samples demonstrate high compatibility with skin fibroblast cells. This may be due to the composition of the samples, which contained elements and/ or components that were less or nontoxic to the cells. The result is also favourable when compared with previous studies.81,82 This suggests that the samples have potential roles in uses for biomedical applications.

3.7.2 Bone-like apatite forming. Growth of bone-like apatite forming from SBF is an important marker of bioactivity which can indicate an ability to bond the samples to the

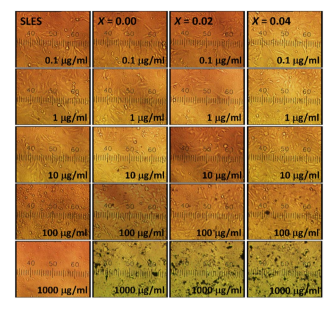


Fig. 9 Cell viability assays observed by an optical microscope at magnification of ×100, using fibroblast cells at different concentrations of SLES (as the positive control in the unit of μ g ml⁻¹) and x concentrations of the (1 – x)[0.99BNKT–0.01BMN]–xCSO ceramics.

surface surrounding tissue. This process is essential for developing bioactive materials with enhanced physical, chemical, and biological functions. In the current study, the bone-like apatite forming of samples was investigated by immersing the samples in the SBF. Fig. 11(a) and (b) present the surface of the x = 0.02 samples before being immersed in SBF and after the SBF test for 15 days, respectively. As compared with the nonimmersed SBF samples (Fig. 11(a)), a cluster of apatite layer was found that did not cover the entire area. EDS analysis for the samples after immersed in SBF is shown in Fig. S6 (in the ESI†). The EDS analysis indicated the formation of an apatite layer, as evidenced by the presence of Ca and P elements.

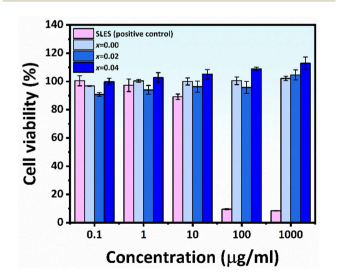


Fig. 10 Histogram plots of cell viability of samples at different CSO contents.

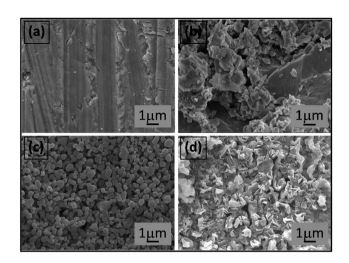


Fig. 11 SEM images of the 0.02 samples for the SBF test: (a) surface of the sample before SBF test, (b) apatite forming after the sample was immersion of in SBF for 30 days, (c) surface of sample coated with hydroxyapatite before SBF test, and (d) surface of the sample coated with hydroxyapatite after immersion in SBF for 15 days.

Notably, samples not immersed in SBF should not contain P, a constituent of the apatite layer.

To increase the ability of apatite formation, the x = 0.02 samples were coated with hydroxyapatite (HAp) by a dip-coating technique, where it is well known that the HAp is a bioactive material which has piezoelectric and pyroelectric properties³⁴ (see additional experimental procedures for the films dip coating in Fig. S5 in the ESI[†]).

The surface of a sample coated with HAp is illustrated in Fig. 11(c). The coated surface showed uniform microstructure and a round grain shape with an average grain size of 0.6 μ m. After the coating process, the samples were immersed in SBF for 15 days. Fig. 11(d) presents the apatite formation after the coated samples were immersed in the SBF. A clear continuous cluster of apatite-like layers was found, as seen in the SEM image in Fig. 11(d). The formation of apatite should be due to negative charge such as PO₄³⁻ anions from the sample surface attracting Ca²⁺ cations from the surrounding SBF, and subsequently attracting PO₄³⁻ to form nucleation sites for apatite development, resulting in the production of apatite layers.^{83,84} The presence of a clear appetite layer for the coated samples suggests the coated samples are a bioactive material.

4 Conclusions

The lead-free ceramics of (1 - x)[0.99BNKT-0.01BMN]-xCSO (x = 0.00-0.04) were synthesized *via* a modified solid state reaction technique. The effects of CSO on the electrical, mechanical, and bio-properties were investigated. The following is a summary of this work.

(1) All samples exhibited a single perovskite phase, with both density and grain size being increased by the CSO additive.

(2) The addition of CSO notably enhanced the electrical properties, where the x = 0.02 samples exhibited the highest energy storage density and electro-strain values at RT.

(3) The mechanical properties, including $H_{\rm V}$, $H_{\rm K}$, and $E_{\rm m}$, were improved by the CSO additive.

(4) The x = 0.02 samples were selected for bio-property tests. The cytotoxicity test indicated that the samples had high cell viability, while the SBF test revealed a moderate apatite forming ability. The SBF test for the samples that were coated with HAp demonstrated an effective apatite-forming ability.

The obtained results suggested that the studied ceramics have potential uses in multifunction electrical devices at RT. In addition, the samples have the potential for further development in biomedical applications in the future.

Data availability

The data supporting this article have been included as part of the ESI.[†]

Author contributions

Lalita Tawee: methodology-sample preparation and original draft preparation. Narongdetch Boothrawong: help the preparation. Suwanan Thammarong: help the preparation. Phanrawee Sriprapha: help the bio-methodology preparation part. Waraporn Boontakam: help the bio-methodology preparation part. Denis Russell Sweatman: electrical measurement, Nopakarn Chandete: improve the manuscript. Chamnan Randorn: improve the manuscript. Gobwute Rujijanagul: conceptualization writing-original draft preparation improve the manuscript, supervision and coresponding authors.

Conflicts of interest

There are no conflicts to declare.

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