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High discharge energy density in novel $K_{1/2}Bi_{1/2}TiO_3$ - $BiFeO_3$ based relaxor ferroelectrics

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ABSTRACT

An increasing number of new dielectrics are being reported for the development of next-generation ceramic capacitors for power electronics used in clean energy technologies. Here, high discharge energy density (W_{dis}) $\sim 6.1 \text{ J cm}^{-3}$ with efficiency (η) $\sim 72\%$ under a pulsed field (E_{max}) of 410 kV cm^{-1} is reported along with temperature stability up to $150 \text{ }^\circ\text{C}$ ($E_{max} = 200 \text{ kV cm}^{-1}$) for $0.5 K_{0.5}Bi_{0.5}TiO_3$ - $0.42BiFeO_3$ - $0.08Sm(Mg_{2/3}Nb_{1/3})O_3$ (KBT-BF-SMN) bulk ceramics. The optimised composition is chemically heterogeneous but electrically homogeneous, similar to several $BiFeO_3$ -based dielectrics reported previously and adding to the growing body of evidence that electrical (measured at weak-field) not chemical homogeneity is the best guide to increased E_{max} and enhanced energy density. KBT-BF-SMN ceramics are therefore considered as promising candidates for pulsed power and power electronics applications.

1. Introduction

Given the importance of clean technologies and electrification in the coming decades, the development of power electronics and devices is recognised as a core future technology. Dielectric ceramic capacitors are passive components which provide superior power density due to their exceptionally fast charge-discharge rate ($\sim \mu\text{s}$) but are typically inferior in energy density with respect to supercapacitors and batteries [1–5]. As the electric vehicle (EV) market grows, not only has the demand for batteries and supercapacitors increased but also for multilayer ceramic capacitors (MLCCs) used in electronics and subsystems [6–10]. In particular, new dielectric materials are needed to facilitate near-engine power electronics to enable ultra-fast charging solutions that facilitate durable EV performance.

Energy storage performance for dielectrics is evaluated from integration of the polarisation-electric field (P-E) loop at the maximum pulsed electric field (E_{pulse}) [1,11]. The E_{pulse} here is the maximum

triangular AC electric field applied during P-E measurement. It differs from breakdown strength which is measured commercially using a continuous dc bias. The total and discharging energy density are obtained using $W_{total} = \int E_{pulse} dP$ and $W_{dis} = \int_{P_r}^{P_{max}} E_{pulse} dP$, where P_{max} and P_r are maximum polarisation and remnant polarisation, respectively [1]. Thus, both large ΔP ($P_{max}-P_r$) and E_{pulse} are desirable for realising high W_{dis} . New electroceramics for high energy-density MLCCs need to demonstrate optimised $E_{pulse} > 300 \text{ kV cm}^{-1}$, $W_{dis} > 4 \text{ J cm}^{-3}$, good temperature stability of permittivity (at least $25\text{--}150 \text{ }^\circ\text{C}$) as well as compatibility with low-cost internal electrodes (Ag, Ni and Ag-10%Pd) to facilitate mass market delivery.

In recent years, perovskite-type dielectrics (traditional class-II type capacitor candidates) have been reported with optimised energy storage performance based either on relaxor-ferroelectrics formed with end members that include $BaTiO_3$ (BT) [12–19], $Na_{0.5}Bi_{0.5}TiO_3$ (NBT) [20–24], $BiFeO_3$ (BF) [25–36] and antiferroelectrics such as $NaNbO_3$ (NN) [37–41] and $AgNbO_3$ (AN) [42–52]. The highest properties to date

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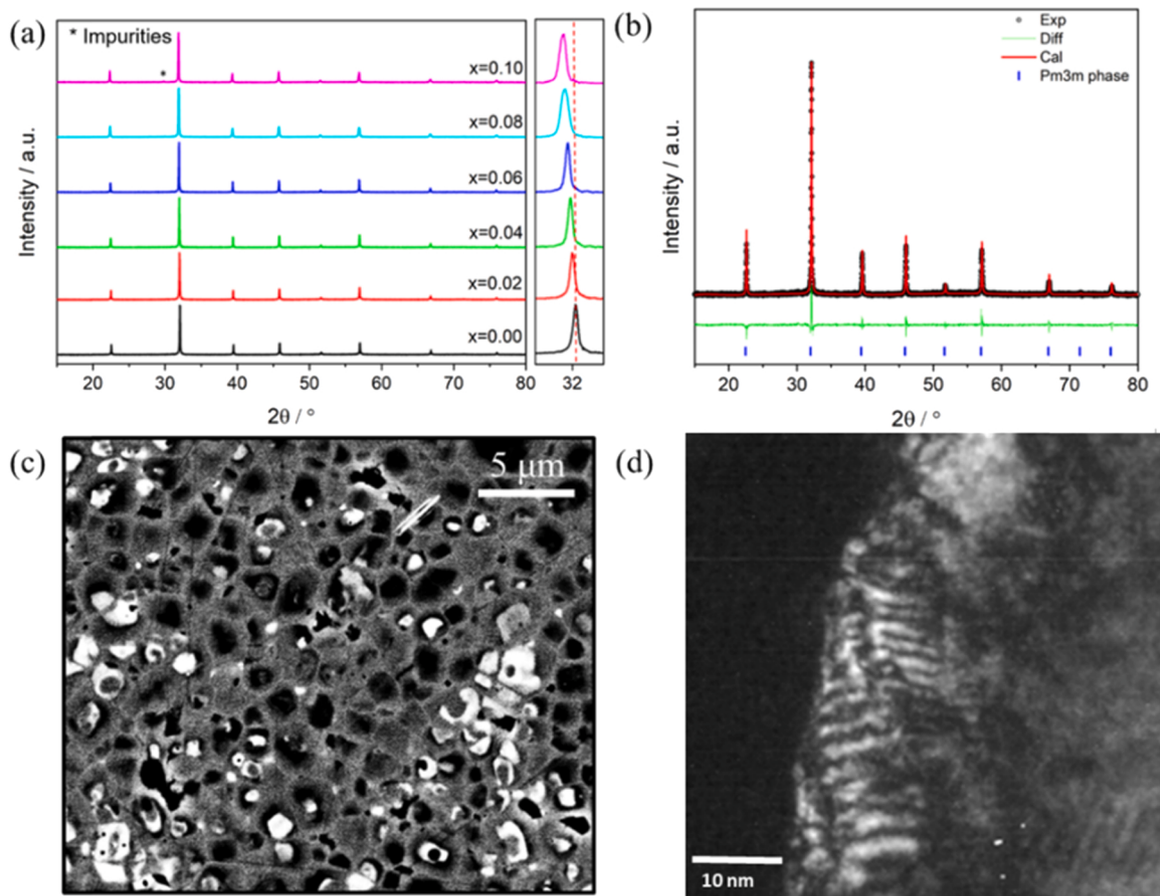


Fig. 1. (a) XRD patterns of KBT-BF-xSMN ($x = 0.00$ – 0.10) ceramics with expanded $\{110\}_p$ peaks. (b) Full-pattern Rietveld refinement using the space group $Pm\bar{3}m$ for $x = 0.08$. (c) BSE image of a polished ceramic surface and (d) TEM image of a grain with lamellar nanodomains for $x = 0.08$.

are reported for relaxor-ferroelectric compositions: $W_{dis} \sim 12.2 \text{ J cm}^{-3}$ under E_{pulse} of 680 kV cm^{-1} was reported by Zuo *et al.* in $0.76\text{NN}-0.24\text{NBT}$ bulk ceramics [41] and Reaney and co-workers succeeded in tailoring the structure and electrical microstructure in BF and NBT-based systems [23,27,28,30] to give W_{dis} up to 7.5 J cm^{-3} and 18 J cm^{-3} in bulk and MLCCs, respectively. The following briefly discusses some further key findings reported for BT, AN and BF-based compositions.

Bi- and Pb-free BT compositions dominate commercial MLCCs market and are compatible with low-cost base metal electrodes but energy densities and breakdown strengths are lower than required in pulsed power and power electronic applications. Modifications using Bi based end members, e.g. $(\text{Bi}_{1/2}\text{Li}_{1/2})\text{TaO}_3$ and $\text{Bi}(\text{Mg}_{2/3}\text{Nb}_{1/3})\text{O}_3$, improve breakdown strength and energy density but necessitate the use Ag-Pd or Pt internal electrodes [12–14,18].

AN based antiferroelectrics have also been studied with a view to optimising energy density. Most studies have focused on optimising E_{pulse} and W_{dis} by stabilising the antiferroelectric with respect to the induced ferroelectric phase, thereby increasing the field required to switch to the polar rather than antipolar state [48]. To date, the highest W_{dis} for AN based bulk ceramics and MLCCs reaches approximately 7 J cm^{-3} (under E_{pulse} of 476 kV cm^{-1}) and 7.9 J cm^{-3} (under E_{pulse} of 1020 kV cm^{-1}), respectively. Most antiferroelectric type dielectrics have promising ΔP values but their energy density is restricted by reaching saturated polarisation in the induced ferroelectric phase. The associated double hysteresis loops generally show conversion efficiency, $< 80\%$, because energy is lost through realignment of dipoles from antipolar to polar. They also suffer from large electric field-induced strain as a consequence of the field induced transition. [52] Some

relaxors type dielectrics however, are reported with high E_{pulse} and energy conversion efficiency (90%). The W_{dis} also benefits from an unsaturated polarisation at higher electric field whose value increases until breakdown. Temperature stability ($>200 \text{ }^\circ\text{C}$) still remains a challenge for both antiferroelectric and relaxors type dielectrics.

BF based dielectrics, mostly alloyed with BT and SrTiO_3 (ST) demonstrate excellent performance not only as bulk ceramics (W_{dis} up to 7.5 J cm^{-3}) but also as MLCCs (W_{dis} up to 15 J cm^{-3}) and thin films (W_{dis} up to 150 J cm^{-3}) [27,28,53]. Their conduction mechanisms (p- or n-type) and electrical microstructures (electrical homogeneity) that underpin superior E_{pulse} were investigated by Wang, Reaney and co-workers who proposed several key factors to enhance the energy storage performance [27,28], such as: reduction in conductivity (controlling defect chemistry); a homogeneous electrical microstructure and a reduction in polar coupling whilst retaining ions with high average ionic polarisability. Several workers have also noted “core-shell” microstructures (bulk ceramics and MLCCs), particularly in BF-BT, but it remains to be elucidated whether these are critical to enhancing energy density [23,27,28,53].

Despite promising energy densities in many current ceramic systems, most experience a widening of the P-E loops at $> 100 \text{ }^\circ\text{C}$ associated with the onset of the movement of V_O under applied field [30]. In 2013, BF- $\text{K}_{0.5}\text{Bi}_{0.5}\text{TiO}_3$ (BF-KBT) based ceramics were reported as high-temperature, piezoelectrics and multiferroics [54–57] with for example, BF-KBT- PbTiO_3 exhibiting good piezoelectric properties up to $450 \text{ }^\circ\text{C}$ [54]. Temperature stable and fatigue-resistant ceramics, KBT-BF- NdFeO_3 were also reported by Khesro *et al.* to exhibit strain variation $< 10\%$ up to $175 \text{ }^\circ\text{C}$ for bulk ceramics and $< 15\%$ for MLCCs up to $300 \text{ }^\circ\text{C}$ [58]. However, KBT is difficult to densify via conventional

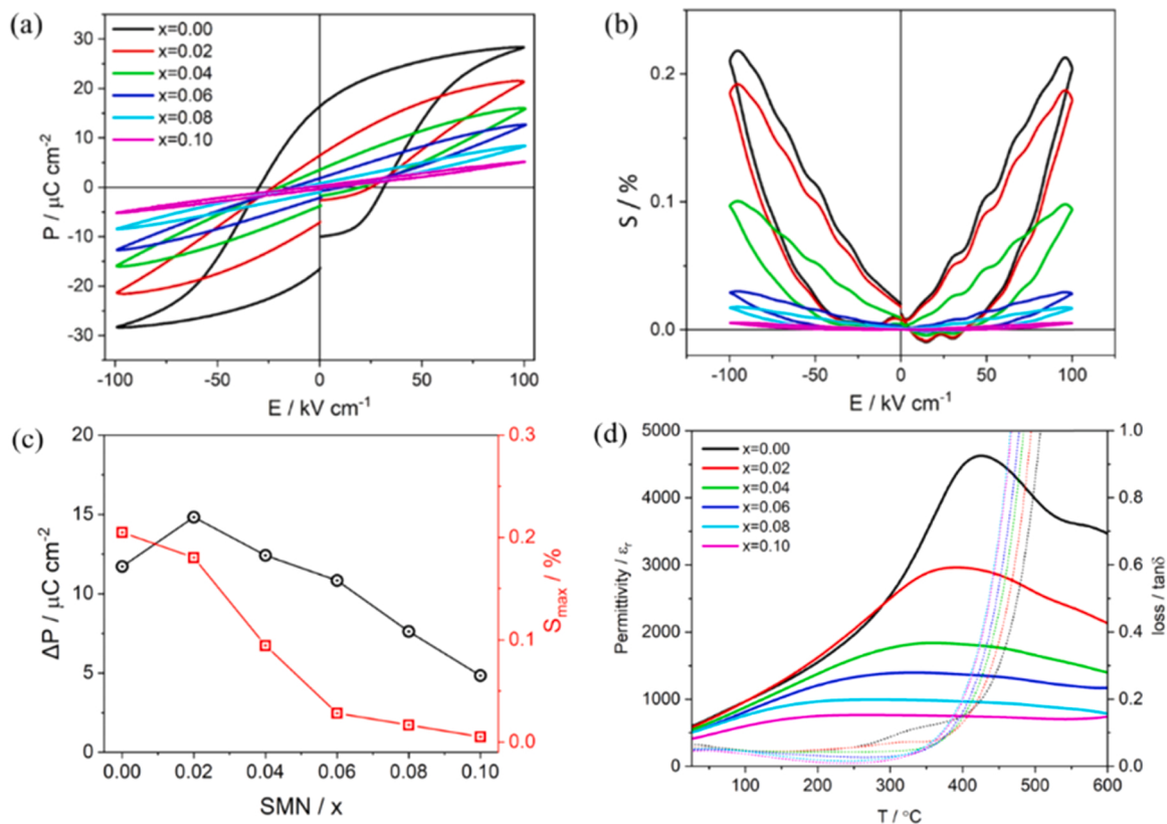


Fig. 2. (a) Bipolar P-E loops and (b) bipolar S-E loops for KBT-BF-xSMN ceramics ($x = 0.00\text{--}0.10$). (c) The changes of ΔP and S_{max} versus SMN concentration, x . (d) Temperature-dependent permittivity and dielectric loss data for KBT-BF-xSMN ($x = 0.00\text{--}0.10$) ceramics at 100 kHz.

solid-state reaction due to volatilisation of K. Therefore, few studies have investigated KBT based compositions as potential candidates for high energy density capacitors with hot pressing often employed to densify ceramics and only modest energy storage properties reported. [59–64].

Here, we report an electrically homogenous, highly-resistive relaxor, $0.5\text{K}_{0.5}\text{Bi}_{0.5}\text{TiO}_3\text{--}0.42\text{BiFeO}_3\text{--}0.08\text{Sm}(\text{Mg}_{2/3}\text{Nb}_{1/3})\text{O}_3$ (KBT-BF-0.08SMN), prepared using conventional solid-state synthesis that exhibits low dielectric loss ($\tan\delta < 0.04\%$ up to $375\text{ }^{\circ}\text{C}$), excellent energy storage performance ($W_{\text{rec}} \sim 6.1\text{ J cm}^{-3}$ at $E_{\text{pulse}} \sim 410\text{ kV cm}^{-1}$) and $< 15\%$ change in W_{dis} up to $150\text{ }^{\circ}\text{C}$ at $E_{\text{max}} \sim 200\text{ kV cm}^{-1}$.

2. Experimental procedure

$0.5\text{KBT}\text{--}(0.5\text{--}x)\text{BF}\text{--}x\text{SMN}$ ($x = 0.00\text{--}0.10$) ceramics were prepared via conventional solid-state reaction using Bi_2O_3 ($>99.9\%$, Sigma Aldrich), Fe_2O_3 ($>99\%$, Sigma Aldrich), K_2CO_3 ($>99\%$, Sigma Aldrich), TiO_2 ($>99.9\%$, Sigma Aldrich), Sm_2O_3 ($>99.9\%$, Sigma Aldrich), MgO ($>99.9\%$, Sigma Aldrich) and Nb_2O_5 ($>99.5\%$, Sigma Aldrich). All reagents were weighed, ball milled for 8 h in isopropanol and calcined for 2 h at $850\text{ }^{\circ}\text{C}$. Calcined powders were then ball-milled again for 8 h and the dried powders uniaxially pressed into 10 mm diameter pellets. The pressed pellets were sintered at $1050\text{--}1110\text{ }^{\circ}\text{C}$ for 2 h in air. The densities of the sintered pellets were measured using the Archimedes method.

Ceramic pellets were crushed into powder and annealed 4 h at $650\text{ }^{\circ}\text{C}$ to remove any strain induced artefacts prior to performing X-ray diffraction. The phase purity and crystal structures of the ceramics were examined using a Bruker D2 phaser X-ray powder diffractometer (XRD) with a $\text{CuK}\alpha$ radiation ($\lambda = 0.15406\text{ nm}$). Full-pattern Rietveld refinement was performed using TOPAS 5 to obtain crystallographic information. The specimens were ground using 1200, 2500, 4000-grade SiC

paper, followed by a polishing using 6, 3 and $1\text{ }\mu\text{m}$ diamond pastes. Ceramic microstructures were studied using an FEI Inspect F50 scanning electron microscope equipped with back scattered electron (BSE) and energy dispersive X-ray spectroscopy (EDX) detectors. Powders of selected ceramics were analysed by high-resolution transmission electron microscopy (HRTEM) using a JEOL 2100.

Ceramic pellets were lightly ground and fired with gold electrode at $850\text{ }^{\circ}\text{C}$ for 2 h prior to electrical measurements. Temperature-dependent dielectric properties were studied using an Agilent 4184 A precision LCR meter from room temperature (RT) to $600\text{ }^{\circ}\text{C}$ at 10, 100, 250 kHz and 1 MHz. Impedance spectroscopy was performed using an Agilent E4980A impedance AC analyser from room temperature to $400\text{ }^{\circ}\text{C}$. Impedance data were normalised by a geometric factor (thickness/surface) and estimated resistivity was obtained using ZView software. Specimens for unipolar polarisation-field (P-E) and strain-field (S-E) measurements were ground to 0.2 mm and sputtered with Au electrodes using an Agar sputter coater. Unipolar P-E and S-E loops were obtained using AixACCT TF Analyzer 2000 ferroelectric tester at 1 Hz from room temperature to $150\text{ }^{\circ}\text{C}$.

3. Results and discussion

XRD patterns of powders from the KBT-BF-xSMN ceramics ($x = 0.00\text{--}0.10$) are shown in Fig. 1a. All samples with $x < 0.10$ could be indexed according to a single perovskite phase whereas a non-perovskite secondary phase was detected by XRD for $x = 0.10$. Full-pattern Rietveld refinements were performed using cubic ($Pm\bar{3}m$), rhombohedral ($R3c$) and tetragonal ($P4mm$) space groups. The best refinement results ($\text{GOF} = 1.05\text{--}1.15$) for all compositions were obtained based on a cubic $Pm\bar{3}m$ phase, Fig. 1b and Table S1 (SI). BSE images of polished KBT-BF-xSMN surfaces are shown in Fig. 1c and Fig. S1 (SI). A complex ‘‘core-shell’’ microstructure was observed for all compositions, with some

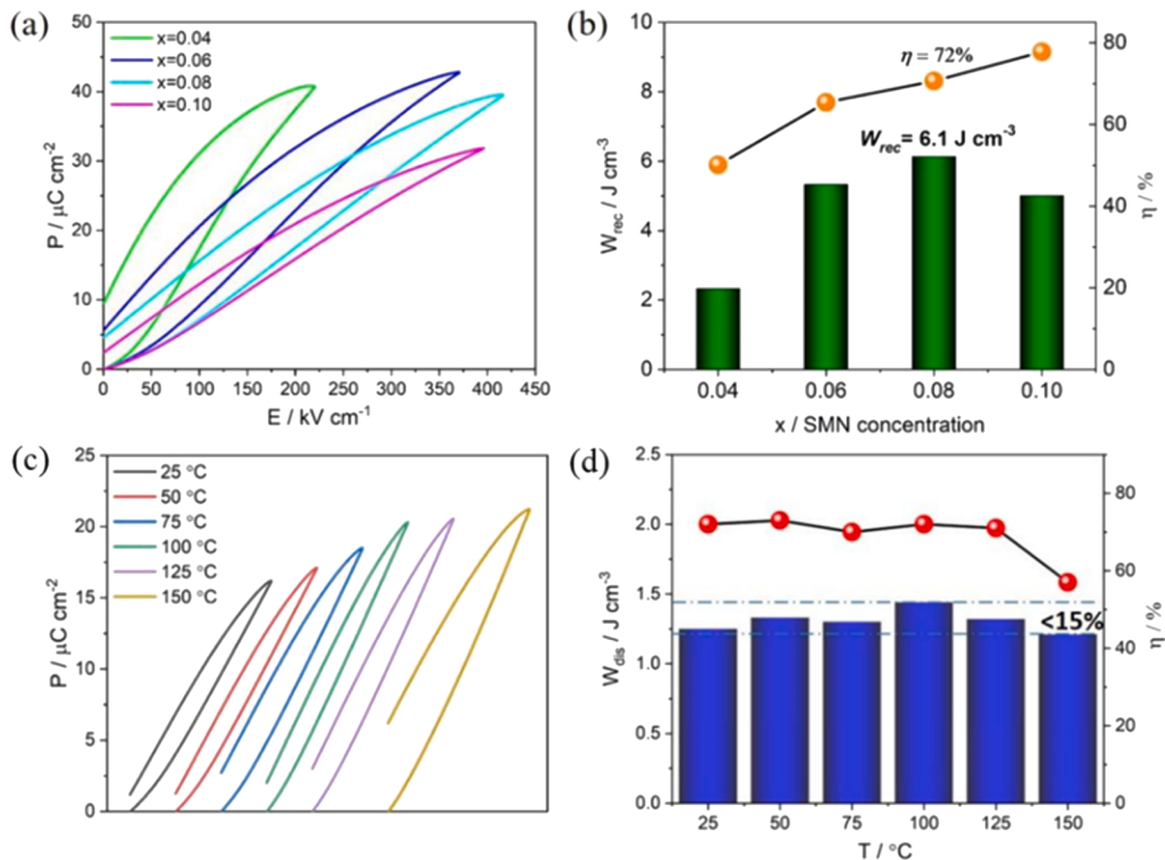


Fig. 3. (a) Unipolar P-E loops at E_{\max} and (b) W_{dis} and η values for KBT-BF-xSMN ($x = 0.04$ – 0.10) ceramics. (c) Temperature-dependent unipolar P-E loops and (d) calculated W_{dis} and η for $x = 0.08$ ceramics.

exhibiting dark Bi-deficient and others bright Bi-rich cores (Fig. S2 SI). This unique microstructural behaviour is formed due to kinetic limitation of the diffusion process and immiscibility on cooling from high temperature. It is reminiscent of that observed by Wang and co-workers [27] in their BF-BT based compositions which despite their chemical inhomogeneity were electrically homogeneous. It has also been reported in other lead-free perovskites, such as NaNbO_3 and BT based bulk ceramics [65,66]. An additional grain boundary (bright) phase was also present for $x = 0.10$, presumably corresponding to the impurity secondary peaks in XRD patterns (Fig. 1a). Unlike in BF based ceramics, the average grain size of the KBT-BF-xSMN solid solutions remained broadly the same irrespective of the value of x , $2.95 \pm 0.56 \mu\text{m}$ for $x = 0.00$ and $2.14 \pm 0.28 \mu\text{m}$ for $x = 0.08$, Table S2 (SI). The local microstructure of $x = 0.08$ was further examined by TEM with lamellar-like nano-domains ($\sim 10 \text{ nm}$) observed within grains, Fig. 1d. Although longer in length than the nano-regions expected for true relaxor ferroelectrics, the lamellar domains indicate a reduced coherence length of polarisation compared with KBT [67].

Bipolar P-E loops at 100 kV cm^{-1} for KBT-BF-xSMN ($x = 0.00$ – 0.10) ceramics are shown in Fig. 2a. The highest $P_{\max} \sim 29 \mu\text{C cm}^{-2}$ and $P_r \sim 17 \mu\text{C cm}^{-2}$ are for $x = 0.00$ with both decreasing with increasing x . The highest $\Delta P \sim 15 \mu\text{C cm}^{-2}$ was obtained for $x = 0.02$, followed by a gradual decrease with increasing x , Fig. 2c. Bipolar S-E loops at 100 kV cm^{-1} for KBT-BF-SMN ($x = 0.00$ – 0.10) ceramics are displayed in Fig. 2b. Electrostrictive-like S-E behaviour is dominant for compositions with $x > 0.00$, suggesting compositions on the whole have a relaxor-like response. We note however, that some negative strain is observed for $x = 0$, consistent with the appearance of a ferroelectric hysteresis loop. $x = 0.00$ showed the highest electric field-induced strain of $\sim 0.22\%$ which decreased with increasing x . For $x > 0.04$, essentially linear P-E loops (P_r close to zero) and negligible electric field-induced

strain ($S_{\max} < 0.05\%$) were obtained (Figs. 2b and 2c), both of which are desirable for energy storage performance and reliability. The temperature-dependent dielectric permittivity (ϵ_r vs. T) and dielectric loss ($\tan \delta$ vs. T) data at 100 kHz for KBT-BF-xSMN ($x = 0.00$ – 0.10) ceramics are shown in Fig. 2d. A frequency-dependent broad peak in ϵ_r at ~ 410 °C (100 kHz data) (Fig. S3 SI) was observed for $x = 0.00$, confirming relaxor-behaviour. As x increases, the ϵ_r vs. T plots become flatter with $\epsilon_r(\max)$ decreasing from ~ 5500 – 900 , the latter varying by $< \pm 15\%$ between 130 and 600 °C. For $\tan \delta$ vs. T , $\tan \delta (< 0.05)$ remains low from RT to ~ 350 °C for all compositions, followed by a sharp rise above 400 °C.

The unipolar P-E loops up to maximum E_{pulse} and the calculated energy storage performance for $x = 0.04$ – 0.10 are shown in Fig. S4 and S5 (SI). For $x = 0.00$ and 0.02 , energy storage performance was not evaluated due to low E_{pulse} and η . The maximum E_{pulse} for the three single phase compositions above $x = 0.02$ increases with increasing x , with the highest $E_{\text{pulse}} = 410 \text{ kV cm}^{-1}$ obtained for $x = 0.08$, as shown in Fig. 3a. The P-E loops become slimmer with increasing x , resulting in the highest $W_{\text{dis}} \sim 6.1 \text{ J cm}^{-3}$ and $\eta \sim 72\%$ for $x = 0.08$ at 410 kV cm^{-1} . The highest $\eta \sim 82\%$ was obtained for $x = 0.10$, as shown in Fig. 3b, yielding $W_{\text{dis}} = 5.2 \text{ J cm}^{-3}$ at 400 kV cm^{-1} . The temperature-dependent unipolar P-E behaviour at 200 kV cm^{-1} for $x = 0.08$ was also evaluated to investigate temperature stability of the bulk ceramics, Fig. 3c. W_{dis} (Fig. 3d) exhibits relatively good temperature stability ($< 15\%$ variation at 200 kV cm^{-1}) from ~ 25 – 150 °C, which is important for practical applications during operation. The widening of the P-E loop > 150 °C here has been found in other BF based system [27,28] and is generally attributed to either the diffusion oxygen vacancies or an electric field-induced phase transition. In previous studies the latter has been postulated as more likely due to the inherently low strain [27,28].

As reported for previous BF-based compositions, electrical

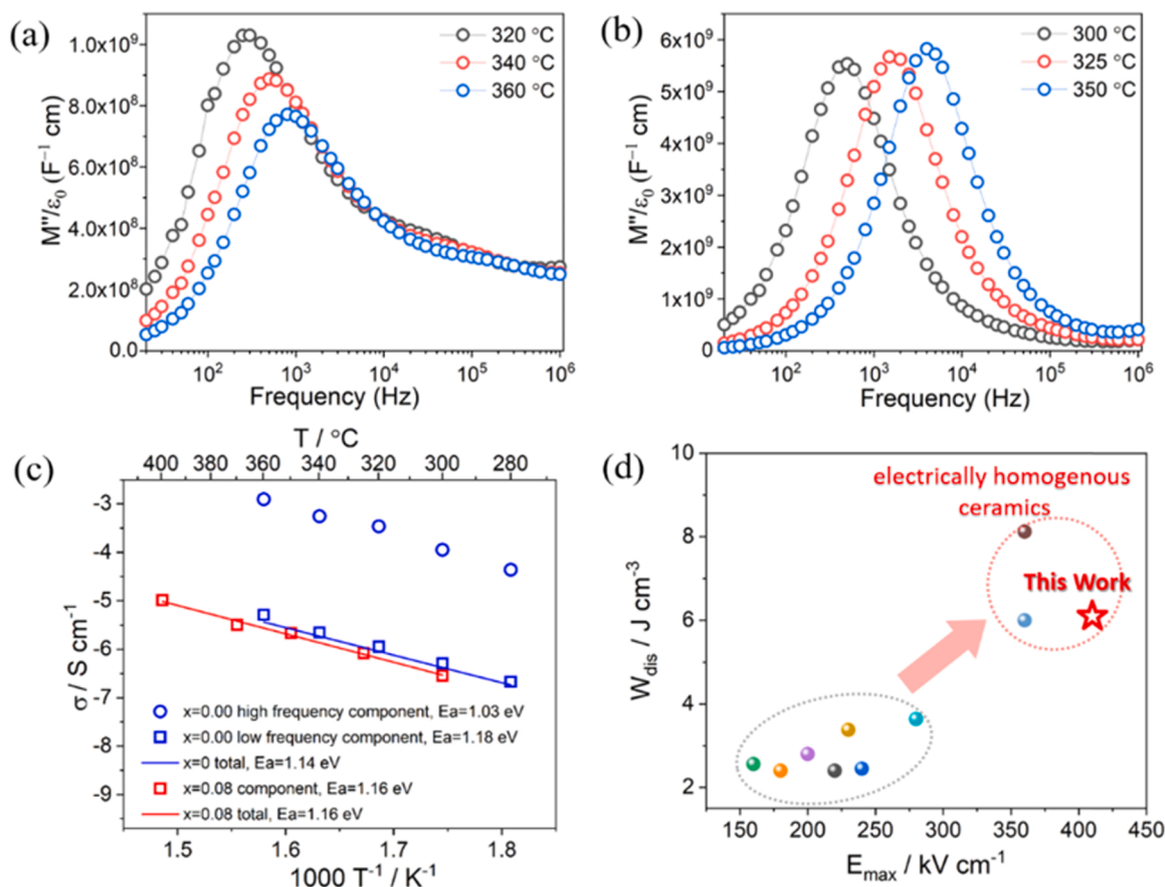


Fig. 4. Temperature-dependent M'' spectroscopic plots for (a) $x = 0.00$ and (b) $x = 0.08$. (c) Arrhenius plots for $x = 0.00$ and $x = 0.08$. (d) Comparison of W_{dis} and E_{max} for BF-containing ceramics reported in the literature [25,27–32,34].

homogeneity plays an important role in optimising energy density.[27] The electrical microstructure of KBT-BF ($x = 0.00$) and KBT-BF-0.08SMN ($x = 0.08$) ceramics were examined using impedance spectroscopy. M'' spectroscopic plot of $x = 0.00$ exhibit one main M'' peak at lower frequency (lower- f) of 10^2 – 10^3 Hz and a second small peak at the higher frequency (higher- f) of 10^4 – 10^5 Hz (Fig. 4a), indicating an inhomogeneous electrical microstructure. At 360 °C, the corresponding capacitance values of the two M'' peaks are 6.47×10^{-10} and 1.65×10^{-9} F cm^{-1} , respectively, and are interpreted as bulk components [68]. The resistivity of the lower- f and higher- f components are estimated to be 307.7 and 0.8 k Ω using the relationship $\omega RC = 1$. In contrast, a single peak is observed in the M'' spectroscopic plot at 350 °C, corresponding to single bulk component with $C \sim 8.58 \times 10^{-11}$ F cm^{-1} and resistivity of 463.7 k Ω , indicating an electrical homogeneous microstructure (Fig. 4b).

The total resistivities of $x = 0.00$ and $x = 0.08$, obtained from the intercept of the arc from Z^* plots, were 91 and 97 k Ω cm at 400 °C (Fig. S6 SI), respectively. The total activation energies, obtained from the intercept of complex Z^* plots, were similar for $x = 0.00$ ($E_a = 1.14$ eV) and $x = 0.08$ ($E_a = 1.16$ eV), Fig. 4c, which is also comparable to other reported compositions, such as BF-BT and BF-ST based ceramics (1.14–1.18 eV) [28,30,69]. It is worth noting that the conductivity for low- f component (resistive shell) of $x = 0.00$ is approximately level up with the total conductivity for $x = 0.08$. However, the higher- f component (conductive core) of $x = 0.00$ is more conductive (at least two order of magnitude) with lower $E_a = 1.03$ eV compared to the lower- f bulk conductivity, leading to potential early breakdown at lower electric field. In contrast, the higher $E_a \sim 1.16$ eV coupled with the electrical homogeneity of $x = 0.08$ favour increasing E_{pulse} . Both compositions are observed to exhibit core-shell microstructure but perhaps

the difference on conductivity between conductive higher- f component (BF-rich core) and resistive lower- f component (KBT-rich shell) is significant for $x = 0.00$. While Nb^{5+} from SMN doping could possible suppress the conductivity in BF-rich conductive core, allowing electrical microstructure to be homogeneous for $x = 0.08$. The electrical homogeneity demonstrated above suggests that the fabrication of MLCC prototypes with high E_{max} is feasible due to the absence of conductive pathways. Although it may seem counterintuitive that compositions with chemical inhomogeneity are electrically homogeneous, these observation are consistent with other BF-based ceramics [25,27–32,34,35] that have been used to form high E_{pulse} (1 MV cm^{-1}) prototype MLCCs. Apart from BF-based ceramics, achieving electrical homogeneity also enhances W_{dis} in other lead-free systems, such as BT- [70] and NBT-based [23,71] bulk ceramics.

4. Conclusions

KBT-BF- x SMN ($x = 0.00$ – 0.10) ceramics were successfully synthesised by solid-state reaction. All compositions had an average (pseudo) cubic structure (XRD), albeit with evidence of lamellar ferroelectric nanodomains by TEM, and exhibited a complex, chemically inhomogeneous “core-shell” microstructure (SEM). However, the electrical microstructure of $x = 0.08$ was homogeneous with an E_a value of 1.18 eV, allowing the composition to achieve excellent energy storage performance, $W_{\text{dis}} = 6.1$ J cm^{-3} and $\eta = 72\%$, under $E_{\text{pulse}} \sim 410$ kV cm^{-1} . In addition, to the high discharging density, KBT-BF- x SMN show a temperature stable ϵ_r from 130 to 600 °C with very little frequency dependence from 10 kHz to 1 MHz and retain their pulsed discharge characteristics until 150 °C at 200 kV cm^{-1} . This material is therefore a potential candidate for next-generation energy-density

prototype MLCCs, provided a low-cost internal electrode system can be found.

Declaration of Competing Interest

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

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Appendix A. Supporting information

Supplementary data associated with this article can be found in the online version at doi:10.1016/j.jeurceramsoc.2022.08.020.

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