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# Time-dependent relaxation dynamics of remanent strain in textured PMN–PZ–PT and PMN–PIN–PT piezoceramics

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### ABSTRACT

Crystallographically textured lead-based piezoceramics, particularly  $Pb(Mg_{1/3}Nb_{2/3})O_3-PbZrO_3-PbTiO_3$ (PMN–PZ–PT) and  $Pb(Mg_{1/3}Nb_{2/3})O_3-Pb(In_{1/2}Nb_{1/2})O_3-PbTiO_3$  (PMN–PIN–PT), play a pivotal role in electromechanical applications due to their enhanced field-induced strain responses. However, challenges in precision positioning arise from heightened remanent strain (S<sub>r</sub>) caused by embedded templates in the textured grains, impacting strain cycle consistency due to increased time delay for S<sub>r</sub> relaxation. This study investigates the timedependent relaxation dynamics of S<sub>r</sub> in textured PMN–PZ–PT and PMN–PIN–PT piezoceramics, focusing on experimentally determining relaxation time between cycles required for complete S<sub>r</sub> relaxation. Our findings reveal that the relaxation time in textured piezoceramics is notably longer than that in their non-textured counterparts, primarily due to pinning of domain walls by embedded templates. This work provides insights into the strain relaxation characteristics of piezoceramics and underscores the importance of time delay in designing precision positioning systems for improved reliability.

### Introduction

Piezoceramics with large unipolar strain response and consistent output stability under an applied electric field are crucial for high-precision positioning actuator applications [1,2]. To meet the need for practical application, piezoelectric single crystals such as Pb(Mg<sub>1/3</sub>Nb<sub>2/</sub>3)O<sub>3</sub>-PbTiO<sub>3</sub> (PMN-PT), Pb(Mg<sub>1/3</sub>Nb<sub>2/3</sub>)O<sub>3</sub>-PbZrO<sub>3</sub>-PbTiO<sub>3</sub> (PMN-PZ-PT), and Pb(Mg<sub>1/3</sub>Nb<sub>2/3</sub>)O<sub>3</sub>-Pb(In<sub>1/2</sub>Nb<sub>1/2</sub>)O<sub>3</sub>-PbTiO<sub>3</sub> (PMN-PIN-PT) have been widely investigated due to their ultra-high piezoelectric coefficient ( $d_{33} > 1400$  pC/N) and electromechanical coupling factor ( $k_p \approx 0.9$ ) [3–6]. However, the production of single crystals is quite demanding in terms of manufacturing cost, controlling the homogeneity of the composition, and limited shapes and sizes [7,8].

Crystallographically textured piezoceramics, fabricated through the templated grain growth (TGG) method, emerge as promising alternatives [7–10]. They offer comparable unipolar strain to single crystals and improved mechanical strength, along with versatility in shapes and sizes [7–10]. However, these textured piezoceramics exhibit notable strain hysteresis compared to single crystals [11]. This hysteresis, which is related to ferroelectric domain wall movement, is influenced by external factors, such as electric field and temperature, as well as internal factors

like stress, grain boundary, and defects [1,12–15].

As hysteresis can impede the material's applicability for precise positioning, it has been the subject of several studies [11,16–18]. One key aspect of this hysteresis is the persistence of remanent strain (S<sub>r</sub>) after removing the electric field, affecting the consistency of subsequent strain cycles [19]. The duration  $S_r$  remains in the material, termed relaxation time, is therefore crucial for achieving consistency in strain responses and remains an area that requires investigation. The time delay for Sr relaxation may vary for different material systems due to differences in domain wall motion, which depends on the material's nature and microstructure. In terms of the material's nature, domain wall motion is related to the coercive field  $(E_C)$ ; materials with higher  $E_C$ have more difficulty in domain wall motion [20]. As a result, materials with higher  $E_C$  may require a longer relaxation time than those with lower  $E_C$  [20–22]. In terms of microstructure, the grain boundary in polycrystalline ceramics can impede domain wall motion, leading to a longer relaxation time [12-15]. For textured piezoceramics, embedded templates can impede domain wall motion, further increasing the relaxation time [7,11,23]. For the single crystals, the relaxation time is expected to be short because there are no grain boundaries and embedded templates.

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Fig. 1. Model representation of domain reorientation of poled ceramics (a), under electric field (b), immediately after removing electric field (c), and after a longer zero-field period (d). S<sub>T</sub>: total strain, S<sub>t</sub>: true strain, S<sub>t</sub>: remanent strain.

Recently, we have reported excellent electric-field induced strain and high  $T_c > 230$  °C obtained from textured 0.4PMN-0.22PZ-0.38PT and 0.37PMN-0.29PIN-0.34PT piezoceramics [24-26]. These textured piezoceramics are currently under consideration to be used for piezopumps to replace the mechanical pump installed in loop heat pipes operating at intermediate temperatures. The role of the mechanical pump is to prevent the occurrence of dry-out of the working fluid in the evaporator and achieve stable start-up of the loop heat pipes by supplying an optimal amount of additional working fluid to the evaporator. Because the mechanical pump is usually bulky and complex [27,28], the piezo-pump with simple and small sizes is expected to be a promising candidate to replace it. The amount of the working fluid supplied by the piezo-pump can be accurately controlled due to the precision positioning characteristic of the piezoceramics. This paper investigates the impact of embedded templates on Sr relaxation time in textured 0.4PMN-0.22PZ-0.38PT and 0.37PMN-0.29PIN-0.34PT piezoceramics for precision positioning applications. By comparing the relaxation time of the textured piezoceramics with those of their non-textured counterparts and single crystal PMN-PT, we aim to understand how microstructure influences Sr relaxation time and consequent strain response, offering insights for optimizing piezoceramics in precision positioning applications.

### Experimental

High-purity powders of PbO ( $\geq$ 99.9 %, Aldrich, USA), MgNb<sub>2</sub>O<sub>6</sub> (H. C. Strack, Germany), ZrO<sub>2</sub> (99 %, Aldrich, Germany), TiO<sub>2</sub> (99.8 %, Aldrich, Canada), In<sub>2</sub>O<sub>3</sub> (99.99 %, Aldrich, USA), Nb<sub>2</sub>O<sub>5</sub> (99.9 % Aldrich, China), Bi<sub>2</sub>O<sub>3</sub> (99.9 %, Aldrich, Germany), BaCO<sub>3</sub> ( $\geq$ 99.0 %, Aldrich, USA), NaCl ( $\geq$ 99 %, Aldrich, USA), and KCl ( $\geq$ 99 %, Aldrich,

USA) were used as raw materials.

The 0.4PMN–0.22PZ–0.38PT matrix powder was synthesized by solid-state reaction processing. Stoichiometric amounts of PbO, ZrO<sub>2</sub>, TiO<sub>2</sub>, and MgNb<sub>2</sub>O<sub>6</sub> powders were mixed in ethanol by conventional ball milling for 24 h using a polyethylene bottle and yttria-stabilized ZrO<sub>2</sub> (YSZ) balls. This mixture was then dried and calcined at 900 °C for 1 h, after which the calcined powder was subjected to planetary ball milling for 50 h to reduce the particle size. The 0.37PMN–0.29PIN–0.34PT matrix powder was synthesized *via* a two-step columbite precursor method [29]. To synthesize InNbO<sub>4</sub> powder, stoichiometric amounts of In<sub>2</sub>O<sub>3</sub> and Nb<sub>2</sub>O<sub>5</sub> powders were mixed by ball-milling for 24 h; following, the mixture was calcined at 925 °C for 6 h in a box furnace. To synthesize 0.37PMN–0.29PIN–0.34PT matrix powder, stoichiometric amounts of PbO, TiO<sub>2</sub>, MgNb<sub>2</sub>O<sub>6</sub>, and pre-synthesized InNbO<sub>4</sub> powders were mixed by ball-milling again for 24 h and then calcined at 850 °C for 6 h.

The plate-like BaTiO<sub>3</sub> (BT) template was prepared via molten-salt synthesis followed by a topochemical microcrystal conversion process. A stoichiometric mixture of Bi<sub>2</sub>O<sub>3</sub> and TiO<sub>2</sub> (2:3 M ratio) was heat-treated in a salt (NaCl + KCl mixture, 1:1 M ratio) at 1000 °C for 1 h to obtain a plate-like Bi<sub>4</sub>Ti<sub>3</sub>O<sub>12</sub> (BiT) precursor. Then a mixture of the BiT precursor and BaCO<sub>3</sub> (1:5 M ratio) was heat-treated again in the same salt at 1000 °C for 1 h to synthesize the BT platelets. From the assynthesized BT platelets, the majority of BT particles smaller than 5 µm were removed utilizing a simple sedimentation technology and the residual BT platelets were used as the template for texturing.

Green compacts of 0.4PMN–0.22PZ–0.38PT and 0.37PMN–0.29PIN–0.34PT piezoceramics were prepared by tapecasting using 5 and 3 vol% of BT templates, respectively, followed by lamination, debinding, and cold-isostatic pressing. Finally, textured



Fig. 2. Unipolar strain response of textured 0.4PMN–0.22PZ–0.38PT ceramics at 2 kV/mm for the first and second cycles in a continuous operation (a) and in a discontinuous operation with a 1 s delay between the cycles (b).



Fig. 3. Electric field profiles with 0 to 0.7 s delays between the first and second cycles applied to textured 0.4PMN-0.22PZ-0.38PT (a) and unipolar strain response of the first and second cycles (b), (c).

0.4PMN-0.22PZ-0.38PT and 0.37PMN-0.29PIN-0.34PT piezoceramics were manufactured by sintering the green compacts at 1200 and 1175 °C for 10 h, respectively. Details of the matrix powder preparation, BT template synthesis and sedimentation, and tape-casting process are given in our previous papers [24–26].

For comparison, non-textured piezoceramics of both compositions were prepared by sintering at the same temperatures and time. A commercial PMN–PT single crystal, provided by Quintes (Korea), was also utilized for an additional comparison. For electromechanical analysis, a silver paste was printed on both sides of disc-shaped samples and fired at 650 °C for 30 min. The samples were poled by applying a direct current (DC) field of 3 kV/mm for 30 min in a silicone oil bath at 120 °C. A laser displacement sensor (aixACCT Systems GmbH 2013, Germany) was used to analyze the field-induced polarization and unipolar strain response.



Fig. 4. Five successive strain cycles of textured 0.4PMN-0.22PZ-0.38PT with 0.1 to 0.7 s delays between cycles (a), a comparison of the first and hundredth strain cycles with a 0.7 s delay (b), and  $S_T$  and  $S_T$  versus time delay between cycles (c).



Fig. 5. Five successive strain cycles of non-textured 0.4PMN-0.22PZ-0.38PT with 0.1 to 0.7 s delays.

### **Result and discussion**

Time delay required for remanent strain relaxation in textured and nontextured 0.4PMN-0.22PZ-0.38PT piezoceramics

When an electric field is applied above the  $E_C$  of ferroelectric materials, the domains align to the level of saturation along the direction of the field, resulting in total strain (S<sub>T</sub>) in the material, as shown in Fig. 1 (a) to (b) [30]. When the external electric field is removed, the aligned domains depolarize slightly but do not return immediately to their original position, resulting in a remanent strain (S<sub>r</sub>), as shown in Fig. 1 (b) to (c) [30]. For an extended zero-field period, as shown in Fig. 1(c) to (d), the depolarization may continue until the S<sub>r</sub> is fully relaxed. The rate of this depolarization can be affected by point defect dipoles, grain

boundaries, pores, etc. that impede domain wall motion, leading to longer relaxation time [12,13,31–35].

Fig. 2(a) shows the unipolar strain response of textured 0.4PMN–0.22PZ–0.38PT piezoceramics containing 5 vol% BT templates measured at 2 kV/mm for the first and second cycles in continuous operation, with a delay set to 0 s. In the first cycle, when the applied field is dropped to zero, the strain remains at a higher level (3) than its starting point (1). The second cycle starts from the endpoint of the first cycle (3) due to unrelaxed  $S_r$ . For the first cycle, the  $S_T$  is therefore the sum of unrelaxed  $S_r$  and the true strain ( $S_t$ ), while for the second cycle  $S_T$  equals  $S_t$ . In discontinuous operation, with delay > 0 s, the second cycle was measured 1 s after the first cycle. For both the first and second cycles,  $S_T$  is equal, as shown in Fig. 2(b), indicating that  $S_r$  fully relaxed in 1 s and the relaxation time therefore is  $\leq 1$  s.

To determine the minimum time delay for achieving full  $S_r$  relaxation in the textured 0.4PMN–0.22PZ–0.38PT piezoceramics, the first and second strain cycles were measured with  $\leq 1$  s delay, as shown in Fig. 3. Fig. 3(a) shows the applied electric field waveforms with various delay intervals between the first and second cycles. It can be observed from Fig. 3(b) and (c) that the gap between the starting point of the first cycle, indicated by the ring, and the second cycle, indicated by the arrow, gradually decreases as the delay between pulses increases. Eventually, the gap disappears at 0.7 s delay, indicating that  $S_r$  fully relaxes and therefore the first and second strain cycles are identical.

To investigate the consistency of the S<sub>T</sub> level, five successive cycles were measured with 0.1, 0.3, 0.5, and 0.7 s delays between the cycles, as shown in Fig. 4(a). For the 0.1 s delay, compared to the starting point of the first strain cycle (indicated by the ring), the starting points of the following strain cycles are at a higher level (indicated by the arrow). At 0.3 and 0.5 s delays, there is noticeable variability, where some cycles' starting levels align with the first cycle (indicated by the ring), while others differ (indicated by the arrow), indicating that full Sr relaxation is not consistently achieved. However, when the delay is extended to 0.7 s, the starting points of all strain cycles converge, signifying consistent full Sr relaxation between successive cycles and resulting in a stable ST output. This is confirmed by comparing the first and hundredth strain cycles with a 0.7 s delay, as shown in Fig. 4(b). Further extension of the delay to 0.8, 0.9, and 1 s, as shown in Fig. 4(c), continues to demonstrate consistent S<sub>r</sub> and S<sub>T</sub> values, highlighting that a delay of 0.7 s or longer is necessary for reliable relaxation.

For the textured 0.4PMN–0.22PZ–0.38PT ceramics, the time delay for achieving full S<sub>r</sub> relaxation could be be influenced by the presence of the embedded BT template, the mechanical stress induced by the crystal structure difference between the 0.4PMN–0.22PZ–0.38PT matrix and BT template [8,25], and a slight Ba-ion diffusion within a limited distance from the embedded BT template to the surrounding 0.4PMN–0.22PZ–0.38PT shell of the textured grains [9]. These three factors are expected to hinder the domain wall movement which results in a prolonged time delay for full S<sub>r</sub> relaxation.

Non-textured piezoceramics may show a shorter delay to  $S_r$  relaxation upon the removal of the applied external field because they do not contain the embedded BT templates. Fig. 5 shows five strain cycles of the non-textured 0.4PMN–0.22PZ–0.38PT piezoceramics measured with 0.1 to 0.7 s delays between cycles. Similar to the textured piezoceramics, the



Fig. 6. SEM micrographs of textured (a) and non-textured (b) 0.4PMN-0.22PZ-0.38PT piezoceramics.



Fig. 7. An SEM micrograph of textured 0.37PMN–0.29PIN–0.34PT piezoceramics (a), P-E loops of textured 0.4PMN–0.22PZ–0.38PT and 0.37PMN–0.29PIN–0.34PT piezoceramics (b), and five successive strain cycles of textured 0.37PMN–0.29PIN–0.34PT piezoceramics with 0.1 to 0.9 s delays (c).

non-textured piezoceramics also exhibit decreasing  $S_r$  with increasing delay. Eventually, the  $S_r$  drops to zero between successive cycles with a 0.7 s delay, resulting in a consistent level of  $S_T$  output. Despite the relatively shorter delay for  $S_r$  relaxation expected for the non-textured piezoceramics owing to the absence of the embedded BT template, the relaxation time was the same as that obtained in the textured piezoceramics.

This can be explained by examining the grain size difference between the textured 0.4PMN–0.22PZ–0.38PT with an average grain size of 23  $\mu$ m and the non-textured counterpart with an average grain size of 4  $\mu$ m, as depicted in Fig. 6. The significant grain size difference between the textured (0.4PMN–0.22PZ–0.38PT) ceramics and the non-textured counterpart can be attributed to the use of TGG in the textured ceramics, which promotes anisotropic grain growth and results in larger grains. In contrast, the non-textured ceramics, processed without templating, exhibit smaller grains due to the absence of such directed growth, as explained in our previous work [26]. This is because the domain wall can be pinned by the grain boundary and the intensity of pinning depends on the grain boundary area [36–41]. As the grain size becomes smaller, the grain boundary area increases, which possibly results in an increased grain boundary pinning effect on the domain wall [37,41,42]. In addition, because the domain size decreases with the square root of the grain size, domain wall-to-domain wall pinning could also be increased [41–45]. Hence, it can be claimed that the embedded 5 vol% BT templates and the large grains in the textured 0.4PMN–0.22PZ–0.38PT counterbalances the small grains in the nontextured counterpart on the domain wall pinning. This suggests that the time delay for S<sub>r</sub> relaxation in the non-textured piezoceramics may not be solely determined by the absence of BT templates, but also by the grain size.

### Time delay required for remanent strain relaxation in textured and nontextured 0.37PMN-0.29PIN-0.34PT piezoceramics

Fig. 7(a) shows the microstructure of textured 0.37PMN–0.29PIN–0.34PT piezoceramics, which contains 3 vol% BT templates embedded in the textured grains. Unlike PMN–PZ–PT, PMN–PIN–PT material systems are known for their high  $E_C$ , indicating that domain wall motion in this material system is relatively difficult [22,23]. The  $E_C$  of the textured 0.37PMN–0.29PIN–0.34PT



Fig. 8. An SEM micrograph of non-textured 0.37PMN-0.29PIN-0.34PT piezoceramics (a) and five successive strain cycles of non-textured 0.37PMN-0.29PIN-0.34PT piezoceramics with 0.1 to 0.5 s delays (b).



Fig. 9. Five successive strain cycles of PMN–PT single crystal with 0.1 to 0.3 s delays.

piezoceramics is higher than that of the textured 0.4PMN-0.22PZ-0.38PT piezoceramics, as shown in Fig. 7(b). As a result, it is expected that the textured 0.37PMN-0.29PIN-0.34PT piezoceramics will take a longer time delay for Sr relaxation than the textured 0.4PMN-0.22PZ-0.38PT piezoceramics. The relaxation of Sr in the textured 0.37PMN-0.29PIN-0.34PT piezoceramics, as shown in Fig. 7(c), follows a similar trend to that observed in the textured 0.4PMN-0.22PZ-0.38PT piezoceramics. However, complete Sr relaxation takes a longer time delay of 0.9 s likely due to its higher  $E_C$  relative to the 0.4PMN-0.22PZ-0.38PT material system.

Fig. 8(a) shows an SEM micrograph of non-textured 0.37PMN-0.29PIN-0.34PT piezoceramics. Compared to the textured 0.37PMN-0.29PIN-0.34PT piezoceramics in Fig. 7(a), the non-textured counterpart in Fig. 8(a) exhibits almost identical average grain size. Therefore, unlike the case in the 0.4PMN-0.22PZ-0.38PT piezoceramics, the grain boundary pinning effect on the domain wall of both the textured and non-textured 0.37PMN-0.29PIN-0.34PT piezoceramics is expected to be nearly the same. As a result, a stronger domain wall pinning is expected in the textured 0.37PMN-0.29PIN-0.34PT ceramics mainly due to the presence of embedded BT templates inside the textured grains. Since there are no templates in the grains of the nontextured 0.37PMN-0.29PIN-0.34PT piezoceramics, a full relaxation of  $S_r$  took only 0.5 s as shown in the strain versus time curve in Fig. 8 (b), therefore, it can be said that the longer delay, 0.9 s, for the textured 0.37PMN-0.29PIN-0.34PT ceramics could be due to the effect of embedded BT templates in the textured grains.

## Time delay required for remanent strain relaxation in a commercial PMN–PT single crystal

Unlike textured or non-textured piezoceramics, single crystals have no grain boundaries and embedded templates. Consequently, the absence of these features is expected to result in relatively weak pinning of domain walls in single crystals, which may lead to a shorter delay for Sr relaxation compared to textured or non-textured piezoceramics. However, in single crystals, the reorientation of domain walls upon removal of the external field can still be influenced by the presence of local point defect dipoles [46-48]. Fig. 9 displays strain versus time of the single crystal PMN-PT, showing a similar S<sub>r</sub> relaxation tendency as the textured and non-textured 0.37PMN-0.29PIN-0.34PT and 0.4PMN-0.22PZ-0.38PT piezoceramics. Despite this, the time delay required for full Sr relaxation for five successive strain cycles of single crystal PMN-PT is 0.3 s. This is much shorter than that required for full Sr relaxation of the textured (0.9 and 0.7 s) and non-textured (0.5 and 0.7 s) 0.37PMN-0.29PIN-0.34PT and 0.4PMN-0.22PZ-0.38PT piezoceramics, respectively. The shorter time delay in the single crystal PMN-PT can be attributed to the absence of grain boundaries and embedded BT templates that pin the domain wall.

Regardless of textured sample, non-textured sample, and single crystal, all relaxation curve showed the decay shape. Further investigation into the decay shape of these relaxation curves is crucial for understanding the creep behavior in ferroelectric materials, which may significantly influence the performance and reliability of piezoelectric devices. To further elucidate the complex domain behaviour in piezo-ceramics, incorporating computational modelling [49] can be instrumental. Models such as phase-field modelling offer a robust framework for simulating domain formation and evolution under various electrical and mechanical boundary conditions. These models can simulate domain wall motion, domain switching, and intergranular interactions that are critical for interpreting the material's macroscopic electrome-chanical properties.

In this study, we have explored the relaxation dynamics of remanent strain in textured and non-textured piezoceramics, focusing on the impact of embedded templates and grain sizes. An interesting area for future research is the investigation of how slight variations in template content might influence the relaxation time of real strain, particularly in highly textured ceramics. While our current analysis touches upon this, a more in-depth study could provide valuable insights into optimizing the performance of piezoceramics for specific applications. This could involve both experimental investigations and modeling approaches to understand the microstructural changes induced by varying template concentrations and their effects on domain wall pinning and relaxation dynamics.

### Conclusion

We have studied the effect of time delay between electric field pulses on the remanent strain relaxation in textured and non-textured 0.37PMN-0.29PIN-0.34PT and 0.4PMN-0.22PZ-0.38PT piezoceramics and a PMN-PT single crystal. A remanent strain was observed when the time delay between pulses was shorter than each material's characteristic relaxation time. For 0.37PMN-0.29PIN-0.34PT ceramics of comparable grain size, the relaxation time was shorter for the nontextured ceramics, while no difference in the relaxation time could be detected in the 0.4PMN-0.22PZ-0.38PT ceramics where the grain size of the textured ceramics was significantly larger than that of the nontextured counterpart. The PMN-PT single crystal had a shorter relaxation time than all the ceramics. Based on this, we can conclude that both embedded templates and grain boundaries prevent domain reorientation, either by directly pinning the domain walls or by increasing the domain wall density. Therefore, both the microstructure and the time delay between pulses should be carefully controlled to achieve consistent electric field-induced unipolar strain response from not only

lead-based piezoceramics but also lead-free piezoceramics for improved reliability in precision positioning applications.

### CRediT authorship contribution statement

**Temesgen Tadeyos Zate:** Writing – original draft, Methodology, Investigation, Data curation, Conceptualization. **Astri Bjørnetun Haugen:** Writing – review & editing, Funding acquisition. **Dariusz Mikielewicz:** Writing – review & editing, Funding acquisition. **Jae-Ho Jeon:** Writing – review & editing, Writing – original draft, Supervision, Funding acquisition, Conceptualization.

### 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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### Data availability

Data will be made available on request.

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