

Giant piezoelectricity in thin films: beyond PZT

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Micro-electromechanical systems (MEMS) incorporating piezoelectric layers provide active transduction between electrical and mechanical energy, which enables highly sensitive sensors and low-voltage driven actuators surpassing the passive operation of electrostatic MEMS. Several different piezoelectric materials have been successfully integrated into MEMS structures, most notably $\text{Pb}(\text{Zr,Ti})\text{O}_3$. Piezoelectric materials with larger piezoelectric response such as the relaxor ferroelectric $\text{Pb}(\text{Mg}_{1/3}\text{Nb}_{2/3})\text{O}_3\text{-PbTiO}_3$ (PMN-PT) would enable further miniaturization. However, this has long been hampered by the difficulties of synthesis. This article will review recent successes not only in synthesizing high-quality epitaxial PMN-PT heterostructures on Si, but also in fabricating PMN-PT micro-cantilevers, which retain the piezoelectric properties of the bulk PMN-PT single crystals. These epitaxial heterostructures provide a platform to build MEMS and NEMS devices that function with large displacement at low drive voltage such as ultrasound medical imagers, micro-fluidic control, piezotronics and energy harvesting.

Keywords: piezoelectric, PMN-PT, epitaxial, MEMS, thin films, multifunctional oxide

Introduction

MEMS devices are continually pushed in the direction of smaller size and increased integration density with faster, larger range of motion, and more powerful actuating elements. Currently this is accomplished by incorporating passive mechanical elements of nano-scale dimensions within larger MEMS devices driven electrostatically. In this scheme, silicon is only a structural material driven by forces that are external to it. Incorporation of active materials directly into the NEMS structure enables direct coupling of the mechanical deformation to the internal electric fields for nano-mechanical actuation and for generation of electrical signals in response to ultra-small mechanical displacements.^{1,2,3,4,5,6} In this case, the sensing and actuating can be combined into the same system, using piezoelectric materials.

Piezoelectricity is an intrinsic, electromechanical coupling phenomenon arising from the non-centrosymmetric arrangement of atoms within the unit cell. Piezoelectric materials develop surface charges in response to an applied pressure (direct effect), and create mechanical displacements in response to an applied electric field (converse effect), which are useful for sensors and actuators, respectively. These active materials are highly desirable for a number of reasons. Such an integrated piezoelectric spring element can generate a large force with a small applied voltage in a linear fashion. This decreases the complexity, retains the integration density, and reduces the voltage burden on the integrated control electronics. In addition, robustness comes from the fact that high electric fields are confined inside solid state materials. A piezoelectric drive is always beneficial at low voltages compared to an electrostatic approach in MEMS actuators, as the force per unit area is linear in electric field (V/thickness) for the piezoelectric drive and quadratic for the electrostatic approach.^{3,4,5}

Applications of piezoelectric materials in MEMS structures typically take advantage of particular piezoelectric response modes, identified by numerical subscripts indicating the axis of the applied field and the axis of the strain or stress response. For instance, a strain induced in the same direction as an applied electric field is labeled as “33”, and a strain induced in a direction transverse to

the applied electric field labeled “31”. These are quantified with piezoelectric coefficients relating the strain to the applied electric field (d coefficients) and stress to the applied electric field (e coefficients). Thin films are also constrained by the boundary conditions imposed by the substrate. These effects are captured by effective piezoelectric coefficients for thin films, $e_{31,f}$ relating in-plane stress to out-of-plane electric field, and $d_{33,f}$, relating out-of-plane strain to out-of-plane electric field³. Both of these responses have been successfully applied in cantilever-type piezoelectric MEMS. For example, vertical capacitor structures with top and bottom electrodes allow the implementation of 31 responses, while the interdigitated electrodes on the top surface permit 33 responses. Both responses are directly measureable, and are used to characterize piezoelectric thin-film response for MEMS structures.

Piezoelectric MEMS

Piezoelectric materials have previously been successfully integrated into MEMS structures, most notably $\text{Pb}(\text{Zr,Ti})\text{O}_3$ (PZT), in both bulk ceramics and thin films.^{1,3,4,5} Although polycrystalline materials have dominated thin film forms, epitaxial films offer distinct advantages, for instance the possibility of lower leakage due to the absence of grain boundaries, and higher piezoelectric response.^{7,8} Recent advances in thin film growth have enabled epitaxial PZT films on yttria-stabilized zirconia (YSZ)^{9,10} and SrTiO_3 (STO)^{11,12} buffered Si. Such integration is essential in progress toward active MEMS, in order to take advantage of Si-based micromachining. As discussed above, use of the 31 piezoelectric response requires both top and bottom electrodes. Although a bottom electrode could have detrimental effects on the epitaxy required for single crystal films and their large piezoelectric response, the perovskite oxide SrRuO_3 has been found to be an excellent conducting material promoting epitaxial growth of a wide variety of functional oxides^{27,28}. Its good lattice match with most oxide materials, chemical and thermal stability, and metallic behavior over a wide temperature range, have positioned it as the workhorse electrode material for epitaxial oxide heterostructures.

Another, more subtle, consideration in thin-film piezoelectrics is the effect of the substrate on the phase diagram. Materials with large piezoelectric coefficients are usually solid solutions of two compounds, with a compositional phase boundary between two low-temperature ferroelectric phases. At this morphotropic phase boundary, an electric field results in a large piezoelectric response. Theoretical work has suggested that epitaxial strain can dramatically modify the phase diagram of PZT thin films.¹³ This indicates that the optimal composition of thin-film piezoelectrics can depend on details of the film/substrate interaction, and need to be determined separately from bulk single crystal optimizations.

Although MEMS integration of piezoelectric materials has been successful to date, piezoelectric actuation with the current generation of materials, AlN, ZnO, or PZT, may not be amenable to further miniaturization due to their relatively small piezoelectric coefficients. For instance, in a piezoelectric based actuator structure such as a basic cantilever, in the simplest model the tip displacement is proportional to the square of the cantilever length, and to the first power of the bias voltage across the piezoelectric film. A shrinking length would require a dramatically higher bias voltage without changes in the piezoelectric properties. Moreover, the emerging field of piezoelectrically-driven electronic devices (piezotronics) requires much larger piezoelectricity beyond these piezoelectric materials.¹⁴ In other words, traditional active materials are likely not active enough!

Giant piezoelectricity of PMN-PT relaxor ferroelectric

The recent development of single crystal relaxor ferroelectrics such as $\text{Pb}(\text{Mg}_{1/3}\text{Nb}_{2/3})\text{O}_3\text{-PbTiO}_3$ (PMN-PT), yielding giant piezoelectric strain, propelled single crystal materials to the forefront of actuator research and development.¹⁵ Incorporation of such active materials with dramatically larger responses into MEMS results in *hyper*-active MEMS, and could extend the applicability of active structures to the nanoscale. The giant piezoelectric strain was attributed to an electric-field induced phase transition from rhombohedral to tetragonal in (001) oriented PMN-PT single crystals. These huge strain levels ($\approx 1.7\%$) and piezoelectric coefficients can be five to ten times those of PZT

ceramics, and the large electromechanical coupling coefficient of $k_{33} \approx 0.9$ promise exciting applications.^{15,16}

These giant piezoelectric properties arise only under specific conditions. First, as in other solid solution piezoelectric materials, the piezoelectric response depends on composition, and is largest near the morphotropic phase boundary (MPB).^{15,16} This phase boundary separates the rhombohedral (PMN-rich side) and tetragonal symmetry (PT-rich side) in the temperature vs. composition phase diagram. For bulk PMN-PT, this phase boundary occurs at 33 % PT in the PMN-PT solid solution system.¹⁶ At the morphotropic phase boundary, the piezoelectric response of PMN-PT is maximized as shown in Figure 1a. However it is not entirely clear how this phase boundary might be modified by strain in a thin film, or from thin-film boundary conditions. As discussed above, there are fundamental considerations that indicate different phase diagrams for thin films. There is also some experimental evidence that the mixed-phase region in MOCVD PMN-PT thin films grown on SrTiO₃ single crystal substrates is different from bulk¹⁷. This indicates that the phase diagram can be modified by the presence of the substrate, as well as corresponding epitaxial strain in the film.

Second, it should be single crystalline with (001) orientation rather than polycrystalline. (001) oriented single crystals exhibit large, non-hysteretic piezoelectric coefficients while (111) oriented ones show a lower, and hysteretic piezoelectric response.¹⁵ The unusually large strain level (≈ 1.7 %) achieved in bulk single crystals originates from the electric-field induced phase transition as described in Figure 1b. The applied electric field along [001], not parallel to the [111] direction of spontaneous polarization in rhombohedral PMN-PT, induces a rotation of polarization toward [001]. For PMN-PT with composition in the vicinity of the MPB, such polarization rotation toward [001] easily destabilizes the rhombohedral ground state, resulting in a phase transition to the tetragonal structure. When the tetragonal phase is driven by an electric field from the rhombohedral ground state, additional room for further strain with the longer c/a ratio is provided, increasing the maximum strain level. On the other hand, an electric field applied along the [111] direction, parallel to the polarization,

reinforces the original rhombohedral ground state without a structural phase transition, limiting the strain level.

Third, they should be purely perovskite phase without any second phases. The pyrochlore phase of $\text{Pb}_2\text{Nb}_2\text{O}_7$ is the most prominent second phase that appears in PMN-PT thin film synthesis. It can be easily formed due to the comparatively low stability of the perovskite phase.¹⁸ This non-piezoelectric second phase degrades the piezoelectric properties of PMN-PT thin films.

Fourth, the stoichiometry of the five constituent elements of PMN-PT should be exact. In particular, the high volatility of lead oxide (PbO) at the growth temperature leads to off-stoichiometry in the films, resulting in not only electrically leaky thin films but also formation of lead-deficient second phases and detrimental grain boundaries.

Epitaxial PMN-PT heterostructures on silicon

Integration of high piezoelectric coefficient PMN-PT films on Si substrates has been pursued for more than a decade, but was realized only recently due to the complexities discussed above.¹⁹ The major difficulty in the thin film synthesis is the extreme sensitivity of film properties to growth parameters. It can be difficult not only to determine optimized growth conditions, but also to reproducibly fabricate high-quality PMN-PT films. Here we describe in some detail recent advances that demonstrate epitaxial integration of PMN-PT with Si, discuss its fabrication into MEMS cantilevers, and discuss broader applications.

Control of growth conditions in high-rate off-axis sputtering^{20,21} allows reproducible synthesis of high quality epitaxial PMN-PT heterostructures on Si using two critical approaches: (1) an epitaxial SrTiO_3 buffer layer on Si and (2) a miscut Si substrate. The miscut substrate has been identified as the key to grow phase-pure PMN-PT films without lead-deficient pyrochlore phase. The role of substrate miscut likely is to maintain film stoichiometry by decreasing the propensity for volatile species (PbO) to desorb. This vicinal approach has been demonstrated to work well for the growth of epitaxial oxide thin films with volatile species.^{20,22}

SrTiO₃ buffer layers on Si were grown by molecular beam epitaxy (MBE) technique.^{23,24,25} These buffer layers enable the growth of epitaxial oxide films on the substrates with a different crystal structure, *i.e.*, perovskite films on diamond cubic Si substrate. Even though the original symmetry of each layer is different in the PMN-PT heterostructure, epitaxy is maintained by the small lattice mismatch of pseudocubic unit cells, as shown in Table I and Figure 2a. As discussed above, SrRuO₃ can provide an oxide bottom electrode, and can also be grown by off-axis sputtering.^{26,27} SrRuO₃ is an ideal bottom electrode²⁸ for epitaxial piezoelectric heterostructures since it is a conductive perovskite with a good lattice match with PMN-PT and the SrTiO₃ template layer. Finally, the PMN-PT films were deposited by high-rate off-axis sputtering.

Figure 2b shows θ - 2θ scans of 3.5 μm thick PMN-PT films on 4° miscut Si substrates. It has an excellent epitaxy of (001) perovskite phase with no detectable pyrochlore phase. In contrast, the PMN-PT films on exact Si grown under the same deposition conditions exhibit a large volume of pyrochlore phase with a polycrystalline perovskite phase.¹⁹ This indicates the importance of substrate miscut in controlling film stoichiometry.

Azimuthal ϕ scans of this phase-pure PMN-PT film in Figure 2c show in-plane epitaxy with a cube-on-cube epitaxial relationship (Fig. 2a), $[100]_{\text{pc}} \text{ PMN-PT} // [100]_{\text{pc}} \text{ SrRuO}_3 // [100] \text{ SrTiO}_3 // [110] \text{ Si}$. (“pc” stands for pseudo-cubic.) The full width at half maximum (FWHM) of the 002_{pc} ω scan, and of the 101_{pc} ϕ scan are 0.26° and 0.6°, respectively. Commercial PMN-PT bulk single crystals have FWHM of 0.14° and 0.27°, respectively. The high-resolution TEM image (Fig. 1d) exhibits an atomically sharp interface between the SrRuO₃ and PMN-PT layers; the epitaxial match between the layers is clear. Also, chemical composition analysis by wavelength dispersive spectroscopy (WDS) shows that the PMN-PT films are stoichiometric with 0.67PMN-0.33PT composition within experimental error. These analyses confirm that PMN-PT thin films on miscut Si satisfy the necessary conditions for giant piezoelectricity shown in the bulk single crystals in terms of stoichiometry, phase-purity, composition, and orientation.

Electrical and piezoelectric properties

One of the intriguing features of these PMN-PT films is a strong downward built-in bias in the as-grown state. As a result, the polarization vs. electric field (P - E) hysteresis loops are shifted (indicating imprint) with a built-in bias of magnitude -38 kV/cm. This can also be seen in the dielectric permittivity vs. electric field measurement (Fig. 3a). The built-in bias is larger than the coercive field of PMN-PT films (≈ 17 kV/cm), resulting in self-poling in the downward direction. This strong imprint benefits the physical property of PMN-PT films in many ways. First, the built-in bias enhances the magnitude of the remanent polarization (P_r): the positive P_r is increased from ≈ 6 $\mu\text{C}/\text{cm}^2$ to ≈ 19 $\mu\text{C}/\text{cm}^2$. This in turn increases the piezoelectric response at zero-applied field. Secondly, the negative built-in bias stabilizes the polarization of PMN-PT in a downward direction. As a result, piezoelectric devices built from these films can be more robust against depolarization due to voltage or temperature excursions, enabling the reliable application of wider range of electric field. Note that such a strong imprint cannot be achieved using bulk single crystals. Thirdly, the built-in bias decreases the permittivity at zero field. This increases the figure of merit for sensors operating in a voltage sensing mode, as well as piezoelectric energy harvesting systems.⁴

As discussed above, this strong imprint is beneficial to applications of PMN-PT thin films. There could be several origins of this relatively strong imprint. Defect dipoles, trapped carriers, strain gradients, nonswitching domains, and structural asymmetries have all been suggested to lead to imprint. In this case, comparison of interdigitated, top-surface only electrodes with asymmetric top and bottom electrode configuration suggests that in these films it arises at least in part from the asymmetric interfaces between Pt / PMN-PT and PMN-PT / SrRuO₃, although further investigations are needed to identify the exact mechanism.¹⁹

The transverse piezoelectric coefficient, e_{31f} , provides the figure of merit for the majority of micromachined piezoelectric sensors and actuators. PMN-PT films on Si shows the maximum e_{31f} value of -27 C/m² ± 3 C/m² measured by wafer flexure²⁹, which are the highest reported for any piezoelectric thin films.

(Uncertainties quoted throughout the paper are one standard deviation statistical uncertainties). This directly reduces the required driving voltage for piezoelectric MEMS actuators.

Moreover, due to the combination of large piezoelectric response and low dielectric permittivity by imprint, the figure of merit for MEMS piezoelectric energy harvesting,^{30,31,32,33,34,35} $\frac{e_{31,f}^2}{\epsilon_0 \epsilon_{33}}$, (ϵ_{33} , relative dielectric permittivity),

shows the highest value. This is also indicative of an enhanced figure of merit for the voltage-driven sensors. Figure 3b compares PMN-PT film's figure of merit with PZT and other piezoelectric materials, demonstrating their advantages for energy-harvesting applications as well as piezoelectric MEMS. These improved responses for actuating and energy-harvesting in PMN-PT films are also confirmed by comparative simulations of PMN-PT and PZT cantilevers using self-consistent data⁹, confirming that the figure of merit is a good indicator of device performance.

The combination of high piezoelectric response with the ability to drive the materials hard without depolarization will enable better resolution and deeper penetration depth in film-based ultrasound imaging.³⁶ Furthermore, the ability to achieve large displacements in small structures with high resonant frequencies offers the possibility for faster switching networks for low-power CMOS circuits³⁷, lower voltage fully mechanical logic devices.^{38,39} Moreover, the integration of multifunctional materials on epitaxial structure with giant piezoelectricity will open a new field of the dynamically strain-controlled devices^{40,41}, which enables controlling optical, electrical, and magnetic properties through induced strain.

PMN-PT microcantilevers

Development of microfabrication processes which do not degrade the large piezoelectric coefficients of PMN-PT thin films is essential for technological applications of these devices⁴². Cantilevers are often fabricated as prototypical structures for electromechanical devices utilizing the e_{31} coefficient, such as

actuators and energy harvesters. Figure 4 shows a successful fabrication process, along with the resulting device performance.

Depth-controlled Ar^+ ion milling with a liquid nitrogen-cooling stage was used to etch metal and oxide layers to define the cantilever structure. An inductively-coupled plasma with a 1:1 mixture of SF_6 and O_2 gases was used to etch the Si substrate beneath PMN-PT heterostructures, where the plasma etches Si from the exposed silicon side walls, releasing the PMN-PT cantilevers. No significant etching of other layers during the Si undercut was found.¹⁹ Various length cantilevers can be fabricated in this way to evaluate the effect of processing on the piezoelectric layer, and to determine in-place piezoelectric characteristics of the thin film layer.

These characteristics are most commonly determined from the overall mechanical response of the cantilever, measured by white light interferometry, which maps out displacements over the entire surface of the cantilever. This method can determine both dc response, where the cantilever profile as a function of dc bias voltage is determined, as well as ac response, where the cantilever resonant frequency is determined by scanning the frequency of an ac bias. For AC measurements, strobed white light interferometry can be used.

In the case of these PMN-PT cantilevers, a sine wave of 50 mV peak to peak was applied without DC bias, and the frequency was changed until the resonance was found. The resonance frequency was independent of voltage amplitude for voltages below 50 mV. The statistical measurement uncertainty is less than ± 0.5 kHz, limited by the mechanical quality factor of the cantilever resonance in ambient air, with the typical full width at half maximum (FWHM) peak width of 4 kHz. Figure 4c shows resonance frequency versus cantilever length. It scales well as $1/\text{length}^2$ as expected from simple mechanical models, indicating that PMN-PT films are spatially homogeneous in terms of thickness and composition over distances from a few micrometers to the inter-cantilever distance of a few hundred micrometers.

Figure 4d shows the mechanical response of 34 μm long cantilever as a function of the applied DC voltage. The tip of the cantilever moves linearly with

voltage with the sensitivity of $0.375 \mu\text{m}/\text{V} \pm 0.005 \mu\text{m}/\text{V}$. This cantilever motion is consistent with the finite element simulation which used the material parameters reported for bulk single crystal PMN-0.33PT.⁴³ This indicates that the PMN-PT films on Si still exhibit comparable physical properties to bulk single crystal unaffected by the fabrication processes.

These results also demonstrate that active MEMS incorporating piezoelectric layers with giant piezoelectric response are viable competitors to electrostatic actuation. The major drawbacks of electrostatic actuation are the required high control voltage and the nonlinear response. The former often prevents scaling down and denser integration of the electronic drivers for actuator arrays. The latter hampers precise control of positioning, and limits the range of operating voltage. The PMN-PT cantilever substantially lowers this driving voltage and exhibits linear displacement with voltage compared to the electrostatic cantilever simulated with comparable structure in Figure 4d. Note that the electromechanical performance of PMN-PT cantilevers is expected to be further enhanced with optimized and precisely controlled thicknesses of all layers for required stiffness and displacement⁴⁴, for instance using silicon on insulator (SOI) wafers.

MEMS applications of epitaxial PMN-PT thin films

It is particularly interesting to consider the potential impact of this technology on applications requiring dense integration of many MEMS actuated devices to provide novel system functionality. When scaling to hundreds or thousands of elements, it is essential that these mechanical systems can be controlled by low voltage signals that can be provided by small, cheap, densely integrated analog or digital circuits. This is particularly important when the density is such that electronics has to be monolithically or hybridly integrated directly above or below the MEMS, as in some optical micromirror array applications,^{45,44} as well as potential future applications of MEMS-tunable nanophotonics^{46,47,48}. At the same time it is desirable, and for some cases essential, to increase the speed of these mechanical elements, which is accomplished by scaling down their size. This scaling to lower drive voltages, smaller elements and higher speeds often cannot

come at the expense of decreased actuator range, which is set by some external parameter, e.g. the wavelength of light in an optical application. This results in a challenging set of system constraints. Integration of high energy density piezoelectric actuators may help overcome this challenge. In addition to increased electro-mechanical coupling, the lower dielectric constant will decrease the capacitive load and typically lower the power dissipation in the drive circuit for high frequency actuation. Finally, better quality single crystal piezoelectric films may lower intrinsic materials losses, improving performance of resonant transducers and decreasing thermal mechanical noise in MEMS sensors.

Conclusions

Active MEMS incorporating piezoelectric thin films as the central element can be reliably fabricated, but even the highest performance traditional piezoelectric materials constrain potential applications. Recent breakthroughs in synthesizing high quality epitaxial PMN-PT films on Si has provided new opportunities to build MEMS with dramatically higher performance, and to extend piezoelectric actuators to the nanoscale. Integrating giant piezoelectric active materials could generate the required large linear forces at small drive voltages with fast actuation, provide accurate displacements at high integration densities, reduce the voltage burden on the integrated control electronics, and decrease NEMS complexity. These heterostructures have tremendous potential to develop the necessary actuators, are compatible with silicon nanofabrication processes, and can be integrated with silicon-based electronics. The strong piezoelectric activity of these PMN-PT thin films promises dramatically greater design freedom for realization of smaller electromechanical devices with even better performance, enabling device applications such as ultrasound medical imaging, micro-fluidic control, energy harvesting, and piezotronics.

However, there remain unanswered questions about this new platform: how can the piezoelectric response of PMN-PT thin films be optimized? How can we design PMN-PT films composition, thickness, strain state, and substrate clamping to produce the optimum device performance for a given device geometry? This

requires an understanding of the fundamental physics in this system. For example, the interplay between strain, substrate clamping, defects, and possible shift of MPB composition needs to be understood. For this, a wide range of interdisciplinary works would be desirable from materials science, physics, mechanics, to computational science. Successful integration of giant piezoelectric materials on Si provides a new challenge, and will stimulate fruitful collaborative researches in solid state physics as well as technology.

Acknowledgements

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Figure Captions

Figure 1. (a) Schematic phase diagram of giant piezoelectric relaxor-ferroelectrics. The solid and dotted blue lines represent the longitudinal piezoelectric coefficients in (001) and (111) relaxor-ferroelectric single crystals, respectively. (b) Schematic illustration of electric-field induced phase transition via polarization rotation. from rhombohedral to tetragonal structure as an origin of huge strain level ($\approx 1.7\%$ in bulk single crystals). Figures are drawn based on Ref. 7.

Figure 2. Epitaxial PMN-PT (3.5 μm) heterostructure on Si with 4° miscut along [110]. (a) Schematic of the PMN-PT heterostructure; (b) X-ray diffraction pattern measured using a 2-dimensional area detector of PMN-PT heterostructure on Si; (c) ϕ -scan of the 101 PMN-PT and 202 Si diffraction peaks. (d) High-resolution cross-sectional TEM image of PMN-PT and SrRuO_3 interface; Pseudo-cubic notations are used for both PMN-PT and SrRuO_3 peak indexing. Figures are adapted from Ref. 9.

Figure 3. Electrical and piezoelectric properties of PMN-PT thin films on Si. (a) Polarization and relative dielectric permittivity versus electric field measurements on 1 μm PMN-PT thin film on Si; (b) Comparison of figure of merit for piezoelectric actuators ($e_{31,f}$) and energy harvesters ($e_{31,f}^2/\epsilon_o\epsilon_{33}$) of PMN-PT film with other reported values. Figures are adapted from Ref. 9.

Figure 4. Fabrication and characterization of PMN-PT microcantilevers. (a) Microfabrication process for PMN-PT cantilevers with Pt (60 nm) / PMN-PT (270 nm) / SrRuO_3 (100 nm) / SrTiO_3 (13 nm); (b) Scanning electron microscopy (SEM) image of PMN-PT cantilever; (c) Resonance frequency vs. cantilever length measurement. The inset is resonance frequency vs. $1/\text{length}^2$; (d) Tip displacement of 34 μm long PMN-PT cantilever vs. applied DC voltage measurement. All measurements were performed at room temperature and

ambient pressure. Error bars represent one standard deviation, and are not shown where statistical uncertainties are smaller than the symbol size. Figures are adapted from Ref. 9.

Tables

Materials	Symmetry	(Pseudocubic) bulk lattice parameter	Growth technique
PMN-PT	Rhombohedral & Tetragonal	$\approx 4.02 \text{ \AA}$	Sputtering ⁹ ($\leq 4 \text{ }\mu\text{m}$)
SrRuO₃	Orthorhombic	$\approx 3.93 \text{ \AA}$	Sputtering ^{17,18} (100 nm)
SrTiO₃	Cubic	3.905 \AA	MBE ¹³⁻¹⁵ ($\approx 150 \text{ \AA}$)
Si	Cubic	3.84 \AA	Czochralski

Table I. Material information of each layer in the epitaxial PMN-PT heterostructure on Si.

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Figure 1.

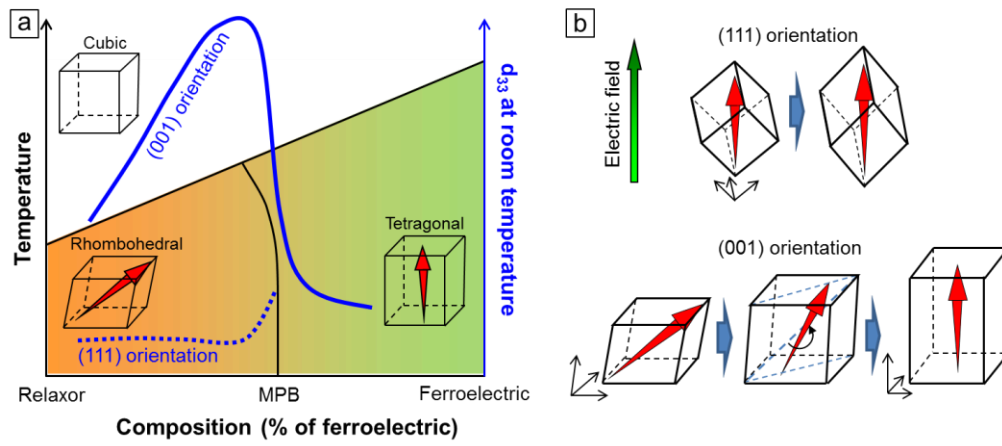


Figure 2.

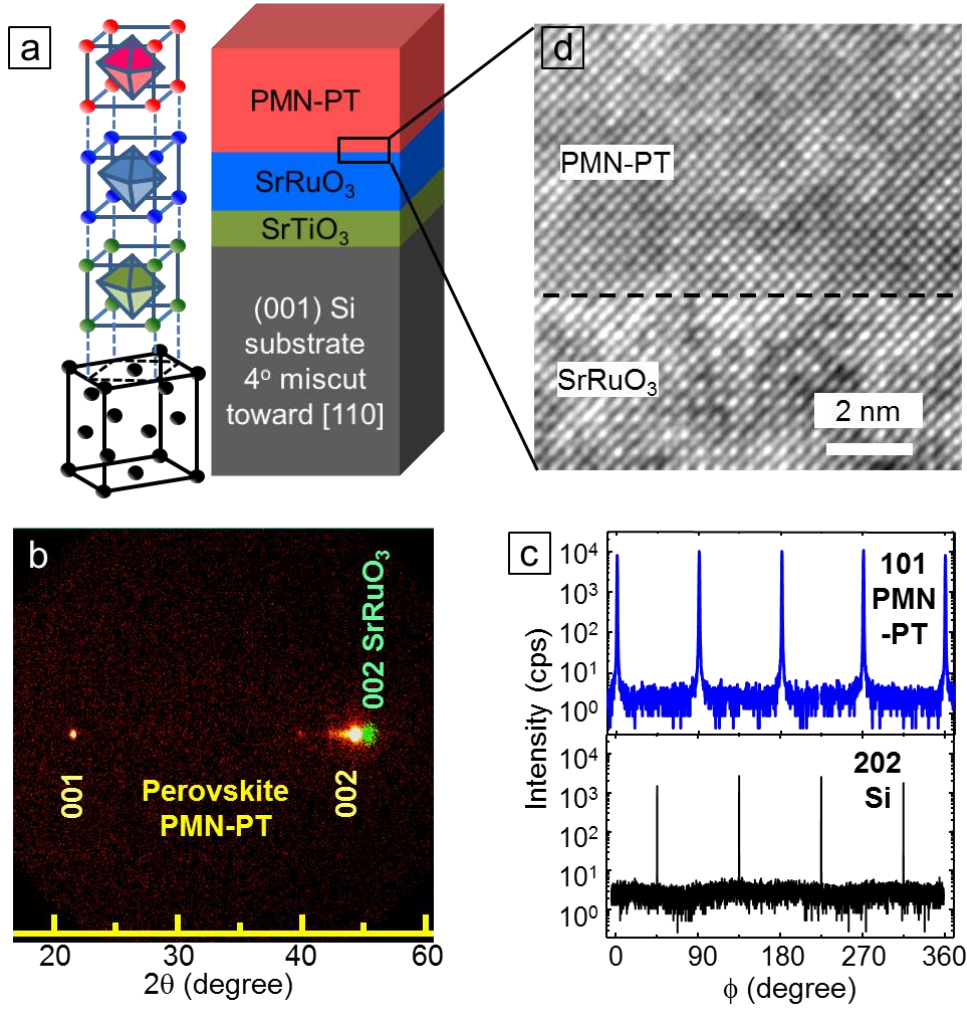


Figure 3.

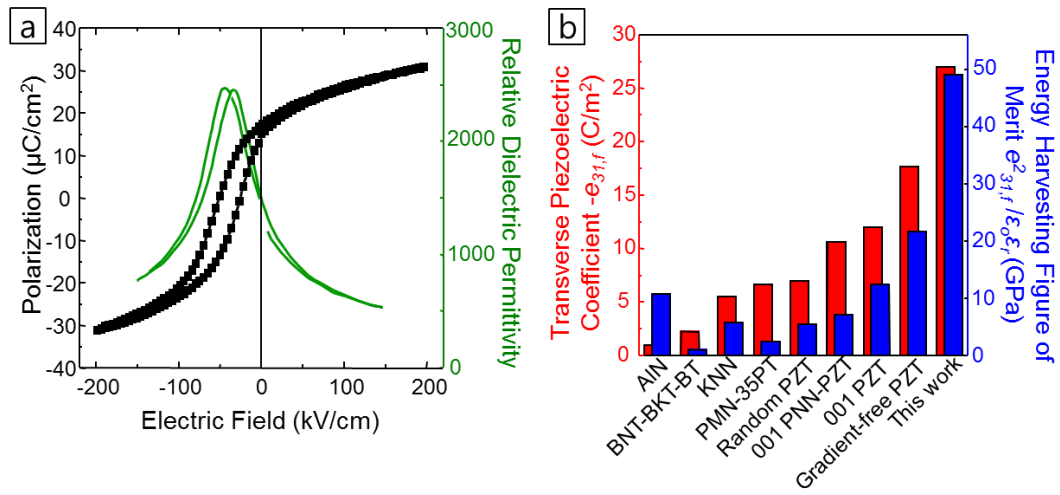
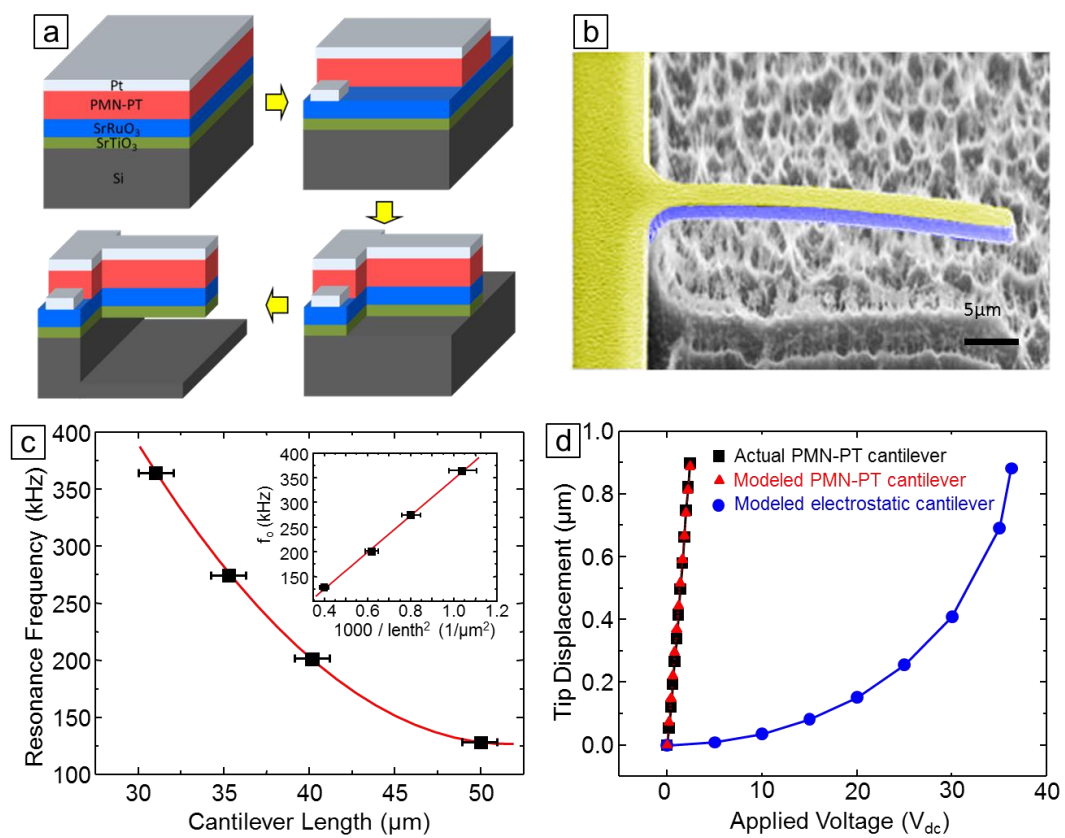


Figure 4.



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