

Innovative Method to Produce Large-Area Freestanding Functional Ceramic Foils

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ABSTRACT

Using thick and thin films instead of bulk functional materials presents tremendous advantages in the field of flexible electronics and component miniaturization. Here, a low-cost method to grow and release large-area, microscale thickness, freestanding, functional, ceramic foils is reported. It uses evaporation of sodium chloride to silicon wafer substrates as sacrificial layers, upon which functional lead titanate zirconate ceramic films are grown at 710 °C maximum temperature to validate the method. The freestanding, functional foils are then released by dissolution of the sacrificial sodium chloride in water and have the potential to be integrated into low-thermal stability printed circuits and flexible substrates. The optimization of the sodium chloride layer surface quality and bonding strength with the underlying wafer is achieved thanks to pre-annealing treatment.

Keywords: freestanding functional foils, NaCl sacrificial layers, PZT ceramic films, ferroelectric.

1. Introduction

Functional ceramics with ferroelectric properties have many applications in electronics as transducers, actuators and sensors [1] that can be used for example in mobile phones [2], structural health monitoring [3], medical imaging [4], robotic surgery [5], touch screens [6], micro-positioning [7] and active vibration control [8]. The use of thick and thin films as opposed to bulk functional ceramics has many benefits in terms of cost, size, weight and sensitivity, which greatly enhances the usability of transducers, actuators and sensors [1]. Therefore, having to integrate a bulky substrate, which serves no other purpose than growing the functional film, is an inconvenience for electronic components fabrication and an obstacle to their miniaturisation. And since growing functional ceramics generally requires a minimal temperature around 500 °C, it is preferable to integrate freestanding microscale foils on low-thermal stability printed circuits or polymers for flexible electronics, rather than growing them directly in-situ. Solutions such as thinning bulk functional ceramics [9], removing the growth substrate [10] or transferring the functional ceramic foil to a new carrier substrate [11,12] using

respectively grinding, etching or a laser process, have been suggested. However, thinning functional bulk materials to less than 30 μm is long, tedious and very challenging as they are generally friable ceramics and the process results in subsurface damage [9]. Moreover, procedures to remove the substrate such as isotropic etching are very costly in terms of time and equipment and consumables [10]. Finally, the laser transfer is time-consuming, limited in feature size and requires expensive transparent growth substrates, generally a glass material with a low coefficient of thermal expansion (α_{CTE}). Besides, there is the technical challenge to focus the beam of a very expensive-to-operate laser at the functional ceramic layer/glass growth substrate interface to transfer the functional layer to a new carrier without creating cracks in the ceramic film. Indeed, the high energy density produced by the laser and the difference between the α_{CTE} of the functional film and the growth substrate, are likely to generate stresses and cause cracks [11,12].

In the past, sodium chloride (NaCl), a non-oxide ceramic, has been used as a cheap, bulk, sacrificial material to achieve freestanding foils of materials such as Ni [13]. Besides, it was deposited onto a suitable carrier surface using an evaporation technique [14–16]. However, to the best of our knowledge, this work is the first successful attempt to deposit functional ceramic films on evaporated, sacrificial, NaCl layers and then release them as microscale, freestanding foils thanks to a simple dissolution of the NaCl interface in water. The chosen functional material was a lead zirconate titanate (PZT) ceramic, developed at Cranfield University and previously studied in literature [17,18], as it is one of the most widely used ferroelectric materials, commonly in the morphotropic phase boundary composition of $\text{Pb}_{1.1}(\text{Zr}_{0.52}\text{Ti}_{0.48})\text{O}_3$ [19,20].

2. Experimental section

A schematic summarizing the whole manufacturing process to obtain Ti/Pt/PZT freestanding foils is presented in Fig. 1. First, NaCl release layers were deposited onto 500- μm thick Si substrates covered with a 300-nm thick top SiO_2 layer, using an Edwards thermal evaporator with a tungsten basket heater and an alumina crucible (B8A-3X.030W from Testbourne Ltd, UK). In a typical procedure, the base pressure was maintained at 1×10^{-4} Pa and the current was successively set to 30 A for 60 s, 50 A, and 40 A for 1250 s to outgas, melt and deposit

250 nm of NaCl (Sigma Aldrich $\geq 99.5\%$), respectively. Afterwards, the NaCl-coated substrates were annealed in a Carbolite box furnace in ambient air at 450, 550, 650 or 750 °C, during 1 or 3 hours. Afterwards, a tape test and a dissolution test into H₂O were carried out to assess the NaCl/SiO₂ bonding strength and the ability to release a functional freestanding foil, respectively. Annealed NaCl layers were coated with electrodes composed of 8 nm of Ti and 250 nm of Pt, using a Nordico radio frequency/direct current (RF/DC) sputter system. Ti and Pt were respectively RF and DC sputtered under argon. Subsequently, 6- μ m thick PZT layers were deposited on the surface of the electrodes using a standard composite sol gel methodology described elsewhere [17,18]. Finally, Ti/Pt/PZT foils were released almost instantaneously from the wafers thanks to a simple immersion in H₂O. In some cases, a Nitto Denko - Revalpha 3195V tape strip was previously bonded to the surface of the PZT layer. The structure and microstructure, were characterised using X-ray diffraction (XRD, Siemens D5000) and scanning electron microscopy (SEM, Phillips XL30). Moreover, the Na and Cl content in the freestanding foils was determined by energy dispersive spectrometry (EDS, Oxford Instruments). Finally, the capacitance loss and ferroelectric properties were studied with a precision component analyser (Wayne Kerr 6425) and a Radiant Technology RT66C Ferroelectric tester, respectively. For the latter characterisation, the PZT film was either Si/SiO₂ wafer-supported and sandwiched between two Pt electrodes, or freestanding and sandwiched between a Pt electrode and a liquid metal eutectic drop of indium, gallium and tin, commonly referred to as Galinstan [21,22], and brought to the PZT surface with a hypodermic syringe. The samples were stored in a vacuum desiccator between each process and/or characterisation step to avoid absorption of atmospheric moisture.

3. Results and discussion

In a preliminary test, a Pt film was deposited on top of an NaCl non-oxide ceramic layer without any previous annealing of the latter, which caused the whole coating to delaminate from the SiO₂ surface. This meant that the stress generated at the Pt/NaCl interface exceeded the binding energy at the NaCl/SiO₂ one. Indeed, since the wafer was held at ambient chamber temperature during the NaCl deposition, the thermal energy was too low to

induce a satisfactory NaCl grain consolidation or a chemical bonding at the NaCl/SiO₂ interface. The influence of the water possibly contained in the hygroscopic NaCl layer was assumed to be non-significant as it should have been extracted when the chamber was vacuumed down to a pressure of 1.33×10^{-5} Pa before the Pt sputtering.

Hence, in order to optimise adhesion of NaCl to SiO₂, and its surface quality to grow a PZT ceramic film upon, post evaporation annealing was performed as described in Table 1. The corresponding samples are referenced as “NaCl T/t ”, with T the annealing temperature in °C and t the dwell time in hours. Three possible beneficial effects were expected [23–25]: a densification of NaCl improving the mechanical integrity of this layer, a grain growth lowering the surface roughness and giving a higher quality interface for the Ti/Pt/PZT film deposition, and the formation of a sodium silicate ceramic phase (Na_xSi_yO_z) promoting a strong chemical bond at the NaCl/SiO₂ interface.

According to the results summarised in Table 1, the samples annealed at 450 and 550 °C, regardless of the dwell time, were soluble in water but mostly peeled off the SiO₂ layer during a tape test, demonstrating weak adhesion. On the contrary, NaCl 650/3, NaCl 750/1 and NaCl 750/3 remained entirely attached to the substrate during the tape test but were not soluble in water, indicating a very strong chemical bonding. Finally, NaCl 650/1 was both soluble in water and unpeeled during the tape test, making it the most likely suitable layer for growing and releasing freestanding Ti/Pt/PZT foils.

The microstructure of NaCl 450/1 and NaCl 550/1 remained quite similar to the as-deposited NaCl one (Fig. 2a) but for NaCl 450/3 (Fig. 2c) and NaCl 550/3 (Fig. 2d), the grains became slightly bigger with rounded angles, more triple junction grain boundaries and fewer intergranular pores. Nonetheless, in both cases, the consolidation due to the densification was not sufficient to prevent the delamination from SiO₂ during the tape test. The microstructure of NaCl 650/1 (Fig. 2b) and NaCl 650/3 (Fig. 2e) exhibited considerably larger grains with almost no intergranular pores remaining, which indicated a higher densification accompanied by a significant grain growth. Therefore, NaCl 650/1 and NaCl 650/3 offered a smoother surface, better adapted to the Ti/Pt/PZT foil deposition. There was also a chemical bonding with SiO₂ as both samples could resist to

delamination during the tape test. In the case of NaCl 650/3 it was even too strong as it became insoluble in water. Finally, when the annealing temperature was increased up to 750 °C, the morphology was completely modified. NaCl 750/1 was composed of very large, smooth grains whereas only a few large grains embedded in a smooth underlying layer could be observed on the surface of NaCl 750/3 (Fig. 2f).

The XRD patterns of NaCl 650/1, NaCl 650/3 and NaCl 750/3, normalised relatively to their respective maximum intensities, are shown on Fig. 3a. The intensity decrease of the NaCl peaks at 31.76 and 45.52° between NaCl 650/1 and NaCl 650/3 indicates that 650 °C was a temperature high enough to initiate a significant chemical bonding thanks to Na diffusion inside SiO₂. However, no sodium silicate phase was detected by XRD, which means that its crystallinity and/or quantity were too low. After a 3-hour annealing treatment at 750 °C, the NaCl phase could not be detected by XRD anymore but peaks corresponding the phases Na₂Si₂O₅, SiO₂ and Na₄SiO₄ appeared at 14.88, 21.84 and 23.12°, respectively. Hence, either all the NaCl reacted or there was not enough left to be detected by XRD. Moreover, as no XRD peak corresponded to a Cl-containing phase, a chlorinated species gas phase very likely accompanied the Na diffusion. The co-existence of the phases Na₂Si₂O₅ and Na₄SiO₄, situated respectively in the left, Na₂O-rich and the right, SiO₂-rich parts of the Na₂O-SiO₂ phase diagram [23] (Supporting Information, Fig. S1), could mean that a Na concentration gradient exists through the newly formed sodium silicate layer. In this case, the Na concentration would be lower in the deeper part of the ceramic layer, leading to the formation of the crystalline phases Na₂Si₂O₅ and SiO₂ (quartz), and higher in the outer part, leading to the formation of Na₄SiO₄. Another explanation would be an inhomogeneous Na diffusion in SiO₂ leading to different Na₂O-SiO₂ ratios and hence phases. A sodium silicate phase with an intermediate composition such as Na₂SiO₃ might also exist but in a proportion too low to be detected by XRD. In any case, the chemical bonding created by the Na diffusion inside SiO₂ explains why the layers annealed at 650 and 750 °C were not peeled during the tape test. Furthermore, the insolubility of NaCl 650/3, NaCl 750/1 and NaCl 750/3 suggests the presence of water-insoluble SiO₂ enveloping the water-soluble sodium silicates. Because of the good compromise between its strong bonding with SiO₂ and its ability to be dissolved into water to release a freestanding foil, NaCl 650/1 was defined as the optimal release layer. An EDS analysis was carried out on both sides of a Ti/Pt/PZT foil released from such a sacrificial layer (Fig. 3b).

Neither Na, nor Cl could be detected on the PZT surface, within the detection limit of the EDS apparatus (ca. 0.1 wt%). As for the surface of the Ti layer, which was in direct contact with the sacrificial layer, no Na and only 0.1 wt% of Cl were found. This means that the sacrificial layer was practically completely dissolved so no significant remaining amount could be identified. Being able to fully dissolve the sacrificial layer is very important because residues of non-conductive NaCl or sodium silicates on the surface of the functional foil would create a contact resistance, proportional to the amount of those insulating species.

Fig. 4a shows a Ti/Pt/PZT foil being released from a Si wafer into water as the NaCl layer is being dissolved. Since the totality of the NaCl release layer was not dissolved exactly at the same time, high stress areas occurred at the interface between the already released and the still bonded parts, thus initiating cracks. Then, given the brittle nature of the PZT ceramic and the high probability of minor defects inside this layer, the cracks could easily propagate throughout the foil. To overcome this problem, a thermal tape strip (Revalpha by Nitto) was bonded to the surface of the PZT layer before its release into water. This not only improved the foil overall mechanical integrity but also limited the angle of curvature at the high-stress released/attached foil interface. With this method, it was possible to release whole, crack-free, Ti/Pt/PZT films in one piece from the SiO₂ surface, as shown in Fig. 4b. The tape-supported foils can also be transferred to another substrate, and the tape finally removed from the PZT surface with a low temperature thermal cycle. The polished cross-section micrographs of a freestanding and a classical Si/SiO₂ wafer-supported Ti/Pt/PZT foil can be observed in Fig. 4c and 4d, respectively. Both of them were ca. 6-μm thick and exhibited a similar microstructure, with a compact stacking of grains, very few intergranular pores (large holes were due to grain removal during polishing) and no internal cracks. Fig. 4c also shows that there is an outstanding conformity between the bottom, dark PZT ceramic layer and the top, bright Ti/Pt metal electrode. Besides, the XRD pattern of a Si/SiO₂/NaCl-supported Ti/Pt/PZT foil, presented in Fig. 3a, confirms that PZT is the main phase constituting the functional film, with only minor traces of a ZrO₂ secondary phase. Firstly, this pattern proves that the structure of a Ti/Pt/PZT film is the same, whether grown directly on a Si/SiO₂ wafer or on an intermediate NaCl sacrificial layer. In the case of the latter, the higher intensity of the peaks corresponding to ZrO₂, and the lower intensity of the peak corresponding to Pt, are simply due to a thicker PZT layer. Secondly, the presence of ZrO₂ highlights the need

of an optimised PZT sintering process (time, temperature) to avoid the presence of this electrically insulating, oxide ceramic, which is expected to lower the overall conductivity of the PZT foil. As ZrO_2 is formed during the sintering of PZT, it is supposed to be homogeneously distributed through the PZT layer, and hence, the electrical resistivity of the ZrO_2 -containing PZT is likely to be proportional to the ZrO_2 content. As for the weak-intensity XRD peak around 45.52° , corresponding to the phase NaCl, it is due to the sacrificial layer sandwiched between the PZT film and the Si/SiO₂ wafer.

Electrical measurements were performed on both Si/SiO₂ wafer-supported and freestanding PZT films. In the first case, the PZT layer was sandwiched between two similar Pt electrodes. In the second case, the freestanding Ti/Pt/PZT foil was in contact with a liquid drop of Galinstan [21,22], which circular contact area with PZT constituted the top electrode, as seen in Fig. 5a. The obtained electric field vs. polarization curves are plotted on Fig. 5b and represent a symmetrical hysteresis loop for the wafer-supported PZT film and an asymmetrical one for the freestanding PZT foil. In the first case, the hysteresis loop symmetry is explained by the fact that the PZT layer is situated between two similar Pt electrodes, thus creating conduction and valance band bendings of the same magnitude in the space charge regions (SCRs) at both PZT/electrode interfaces [26,27]. In the second case, the difference between the Pt and the Galinstan electrodes causes a difference in the band bending magnitude on either side of the PZT film and thus a difference in the length of the SCR into the PZT bulk, which leads to an asymmetrical hysteresis loop. It is also possible that the level of band bending at the Galinstan/PZT interface is not consistent in both orientations when the domain is switched, which also causes an asymmetry in the curve. The freestanding PZT foil had a dielectric constant of 461 at 1 kHz and a higher remnant polarisation than that of the constrained film.

4. Conclusion

In summary, PZT functional ceramic films were successfully synthesised on Si/SiO₂ wafer-supported NaCl sacrificial layers and released as *ca.* 6- μm thick freestanding foils by a simple dissolution of the NaCl layer in water. An annealing of the NaCl non-oxide ceramic prior to the PZT films growth was necessary to avoid

delamination from the wafer during this process. The influence of the temperature and dwell time on NaCl microstructure and interaction with SiO₂ was carefully studied. The annealing parameters offering the best compromise between the sacrificial layer dissolution and its bonding strength to the SiO₂ layer were 650 °C during 1 h and no significant amount of residual NaCl was found on the surface of the freestanding foils after their release. The Ti/Pt/PZT foil grown on the NaCl sacrificial layer had a crystallographic structure and a microstructure very close to those of a Ti/Pt/PZT film grown directly on a Si/SiO₂ wafer. Moreover, the ferroelectric properties of the freestanding Ti/Pt/PZT foil were slightly higher than those of a classically grown, wafer-supported Ti/Pt/PZT film, as evidenced by the higher remnant polarization of the freestanding foil. Due to the good thermal stability of NaCl ($T_f = 801$ °C), the method presented here could be generalised to a wide variety of freestanding functional foils and find numerous applications in electronics, particularly when components miniaturisation or flexible substrates are required.

Figure captions

Fig. 1. Schematic of the manufacturing process of the Ti/Pt/PZT freestanding foils.

Fig. 2. SEM top-views of a) as-deposited NaCl, b) NaCl 650/1, c) NaCl 450/3, d) NaCl 550/3, e) NaCl 650/3 and f) NaCl 750/3.

Fig. 3. a) XRD patterns of NaCl 650/1, NaCl 650/3, NaCl 750/3, a Ti/Pt/PZT film classically grown directly on a Si/SiO₂ wafer, and a Ti/Pt/PZT foil grown on a wafer-supported, NaCl sacrificial layer; b) EDS spectra carried out on both sides of a Ti/Pt/PZT freestanding foil.

Fig. 4. Optical images of a) a freestanding PZT foil being released in water as the NaCl layer is being dissolved and b) a tape-bonded, released PZT foil; and SEM polished cross-sections of c) a freestanding PZT foil and d) a Si/SiO₂-supported PZT film.

Fig. 5. a) Optical image of the measurement for the ferroelectric characterisation of the PZT freestanding foil and b) electric field vs. polarization curves for a wafer-supported PZT film and a freestanding PZT foil.

Table caption

Table 1. Annealed NaCl samples ID, annealing temperature and time, and dissolution and tape test results.

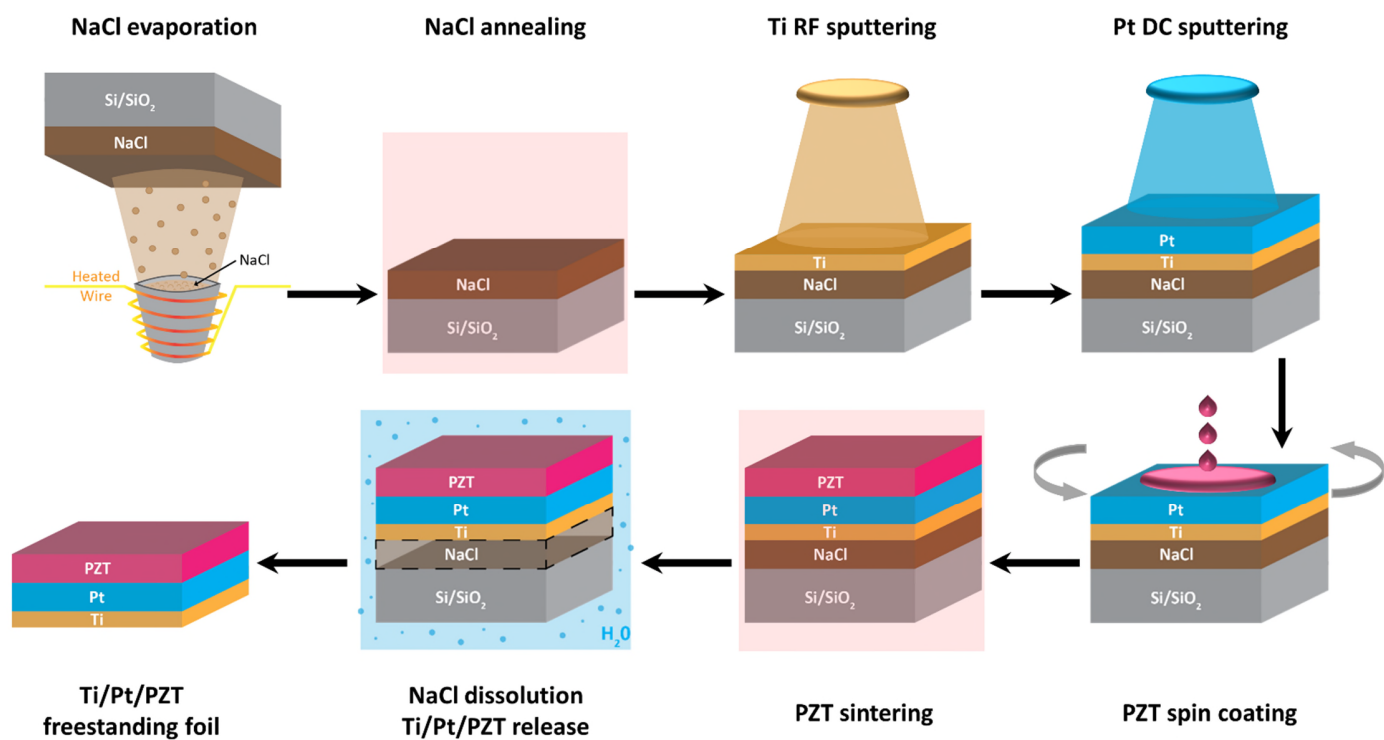


Fig. 1. Schematic of the manufacturing process of the Ti/Pt/PZT freestanding foils.

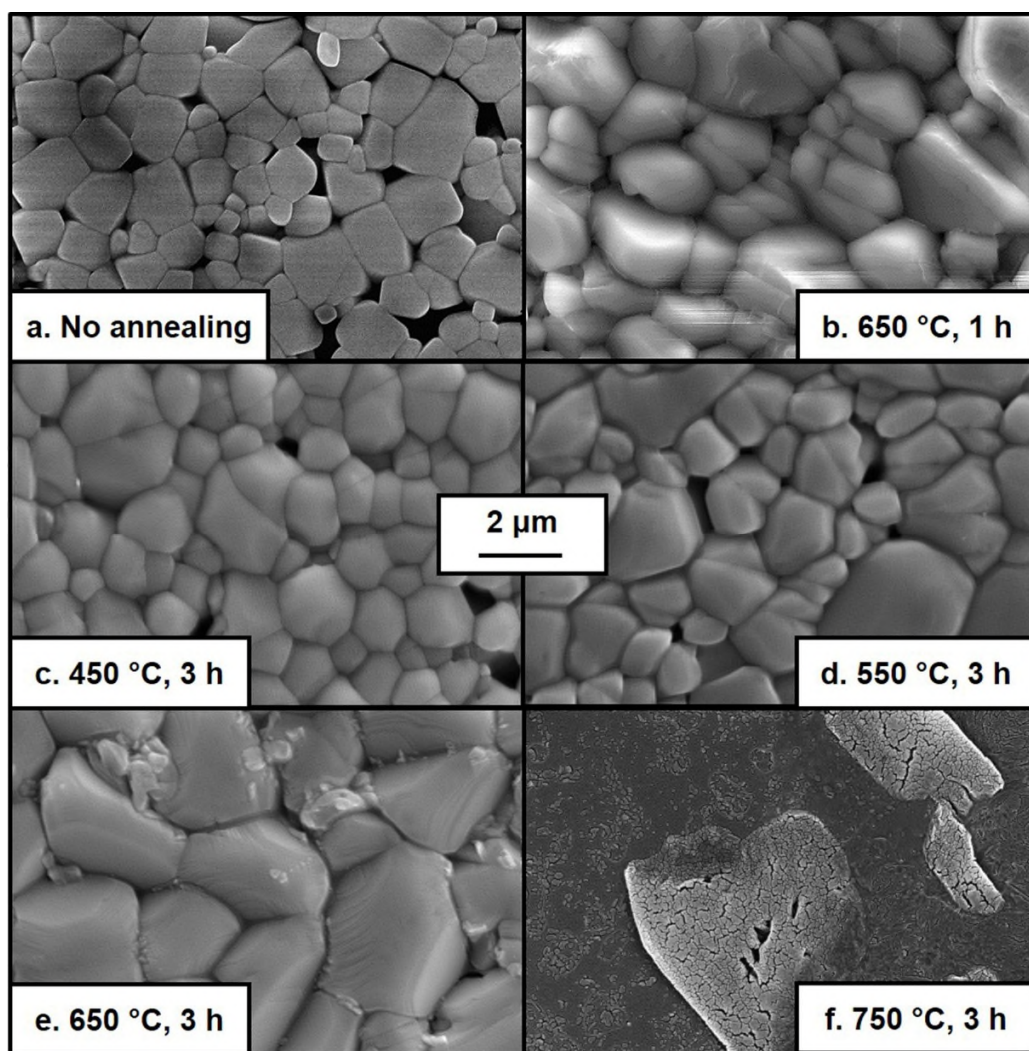


Fig. 2. SEM top-views of a) as-deposited NaCl, b) NaCl 650/1, c) NaCl 450/3, d) NaCl 550/3, e) NaCl 650/3 and f) NaCl 750/3.

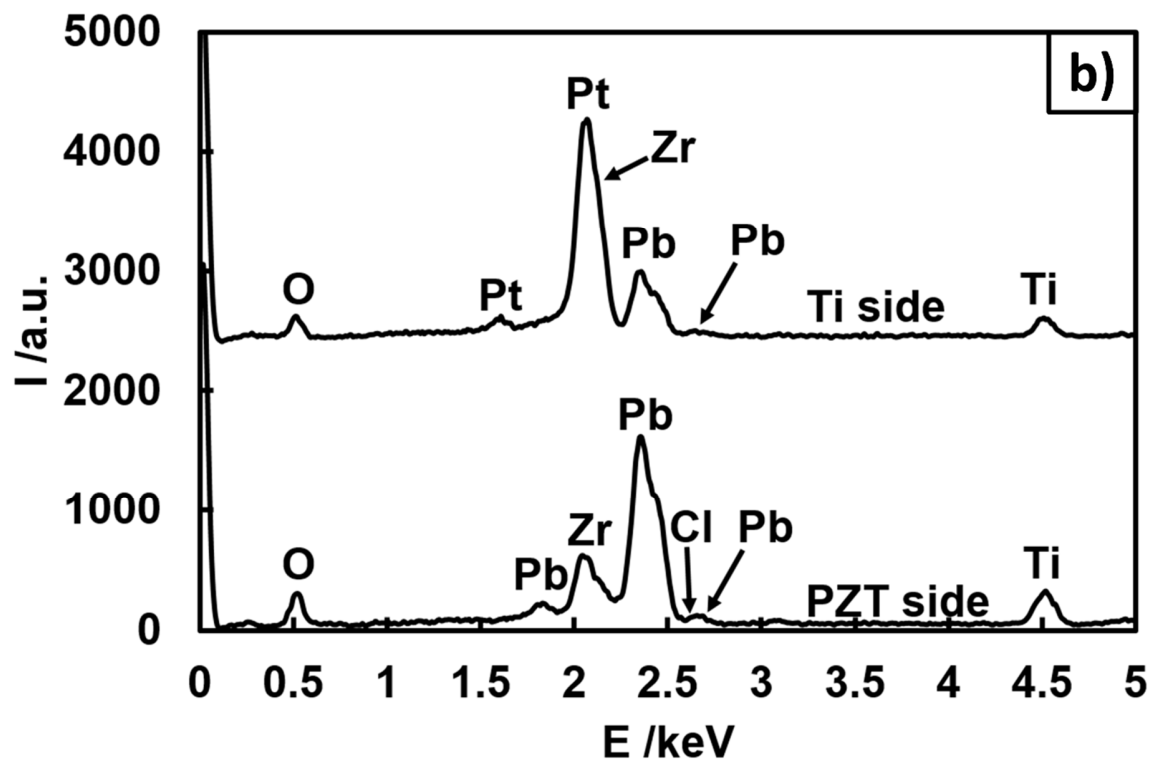
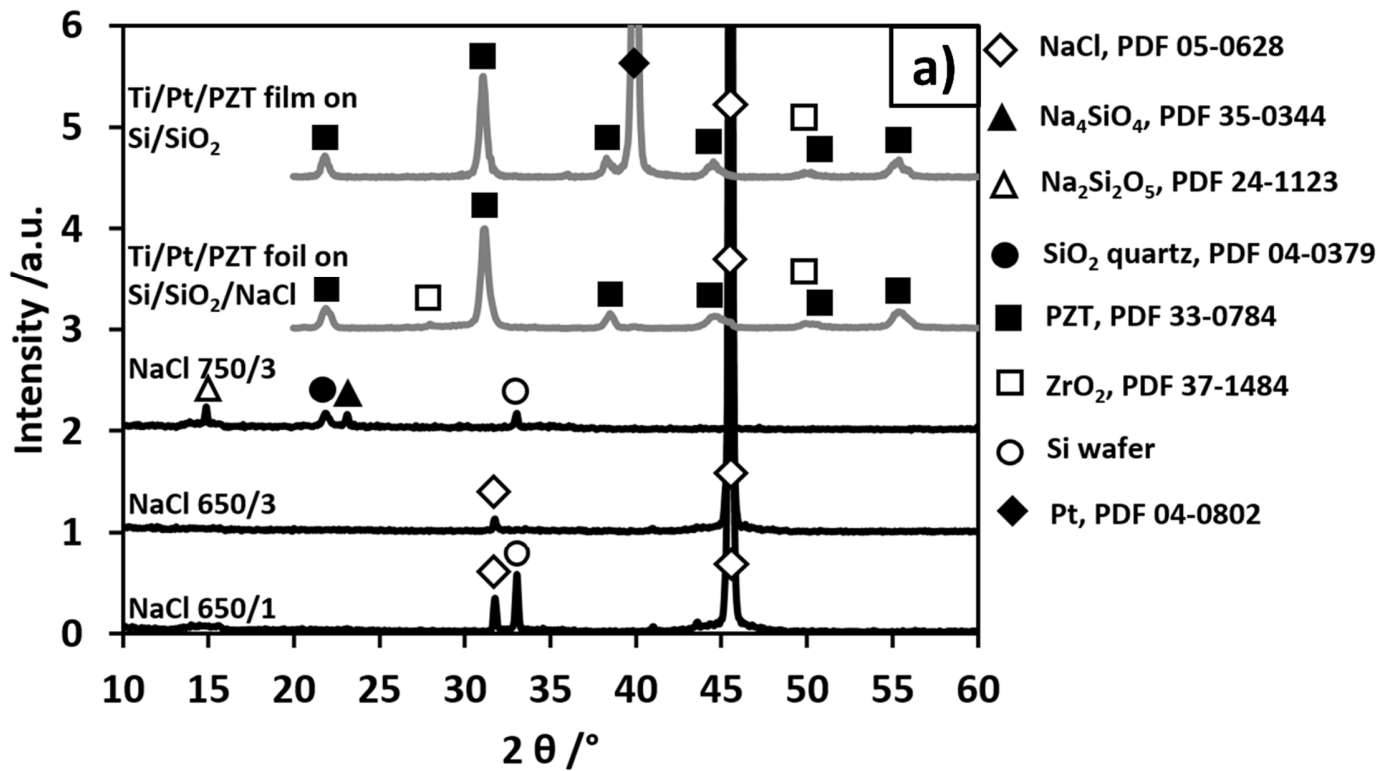


Fig. 3. a) XRD patterns of NaCl 650/1, NaCl 650/3, NaCl 750/3, a Ti/Pt/PZT film classically grown directly on a Si/SiO₂ wafer, and a Ti/Pt/PZT foil grown on a wafer-supported, NaCl sacrificial layer; b) EDS spectra carried out on both sides of a Ti/Pt/PZT freestanding foil.

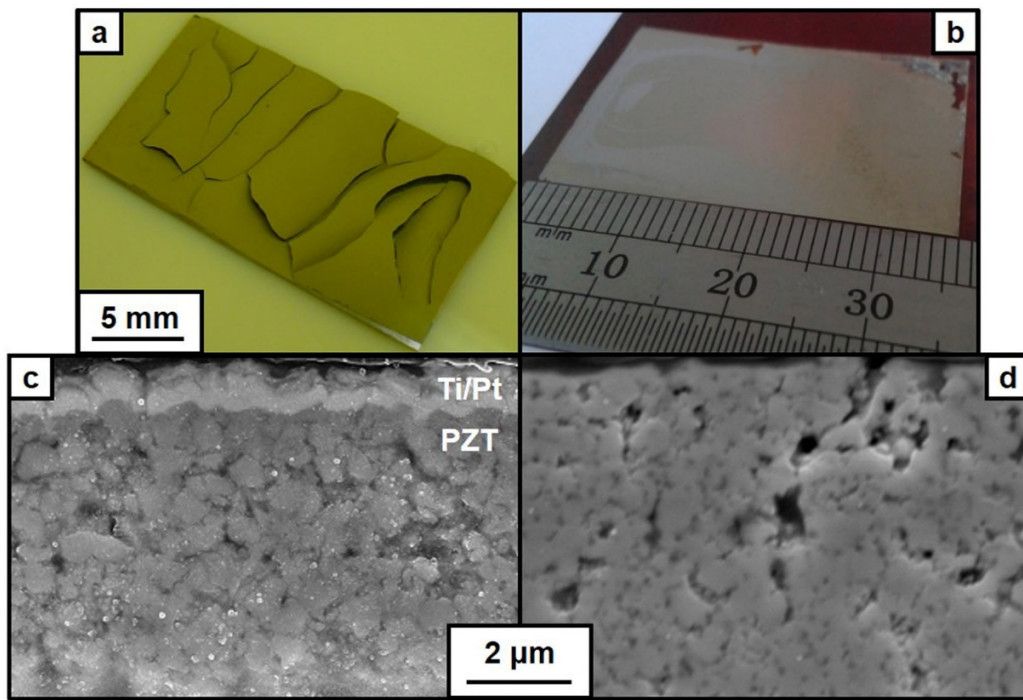


Fig. 4. Optical images of a) a freestanding PZT foil being released in water as the NaCl layer is being dissolved and b) a tape-bonded, released PZT foil; and SEM polished cross-sections of c) a freestanding PZT foil and d) a Si/SiO₂-supported PZT film.

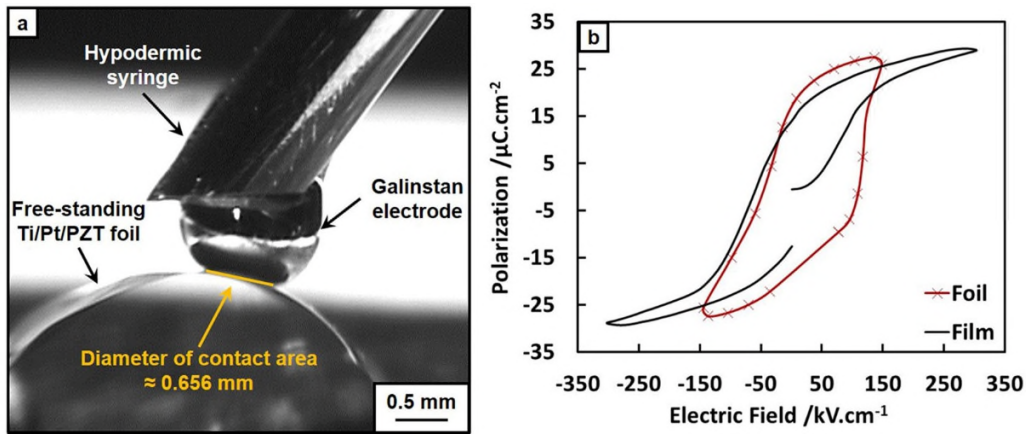


Fig. 5. a) Optical image of the measurement for the ferroelectric characterisation of the PZT freestanding foil and b) electric field vs. polarization curves for a wafer-supported PZT film and a freestanding PZT foil.

Table 1.

Annealed NaCl samples ID, annealing temperature and time, and dissolution and tape test results.

Sample ID	T /°C	t /h	Dissolution into H ₂ O	Peeling during tape test
NaCl 450/1	450	1	Yes	Yes
NaCl 450/3	450	3	Yes	Yes
NaCl 550/1	550	1	Yes	Yes
NaCl 550/3	550	3	Yes	Yes
NaCl 650/1	650	1	Yes	No
NaCl 650/3	650	3	No	No
NaCl 750/1	750	1	No	No
NaCl 750/3	750	3	No	No

Appendix A. Supplementary data

Supplementary data associated with this article can be found in the online version.

Conflict of interest

The authors declare no competing and/or financial interest.

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