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Domain wall engineering: mastering topological defects in ferroelectric and ferroelastic materials

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Abstract | Ferroelectric and ferroelastic domain walls are two-dimensional (2D) topological defects with thicknesses approaching the unit cell level. When this spatial confinement is combined with observations of emergent functional properties, such as polarity in non-polar systems or electrical conductivity in otherwise insulating materials, it becomes clear that domain walls represent a new and exciting state of matter. In this review, we discuss the exotic polarisation profiles that can arise at domain walls with multiple order parameters and the different mechanisms that lead to domain wall polarity in non-polar ferroelastic materials. The emergence of energetically degenerate variants of the domain walls themselves suggests the existence of interesting quasi-1D topological defects within such walls. We also provide an overview of the general notions which have been postulated as fundamental mechanisms responsible for domain wall conduction in ferroelectrics. We then discuss the prospect of combining domain walls with transition regions observed at phase boundaries, homo- and heterointerfaces, and other quasi-2D objects, enabling emergent properties beyond those available in today's topological systems.

Key points

- In ferroelectrics, the emergence of a second polarisation component leads to analogues of Bloch and Néel walls. The stabilization of these walls opens the possibility for quasi-1D topological defects separating wall regions of opposite polarities.
- Polar domain walls in ferroelastics rely on two mechanisms: a polarity imposed by the natural symmetry of strain-compatible domain walls, which can often be described by flexoelectric gradient coupling, and the emergence of a potentially switchable polarity when their natural symmetry is broken.
- Several mechanisms are responsible for domain wall conduction in ferroelectrics: extrinsic intra-bandgap defect states, intrinsic suppression of the conduction band and intrinsic shift of the band structure induced by local electric fields.
- Transition regions occurring at phase boundaries, homo- and heterointerfaces, and other quasi-2D objects probably exist at a smaller length scale, in the vicinity of domain walls, and could lead to exceptional properties and coupling phenomena.

Introduction

Ferroc materials (ferromagnets, ferroelectrics, ferroelastics, ferrotoroidics) exhibit a phase transition from a high-symmetry phase, known as the paraphase, to a low-symmetry phase that exhibits a new physical quantity, characteristic of the symmetry change, called the order parameter (magnetic moment, spontaneous polarisation, spontaneous strain, toroidal moment, respectively)¹⁻³. If a material exhibits several ferroic orders, it is referred to as multiferroic^{1,4,5}. Due to the energy degeneracy in possible orientations of the order parameter upon cooling from the paraphase, ferroic phases spontaneously exhibit domain structures. Regions of uniform order are called domains and the boundaries between adjacent domains, which are naturally occurring 2D topological defects, are called domain walls. The order parameters can be switched, and thus the domain wall structure engineered, by application of an external field (magnetic field, electric field, stress)¹⁻³.

Considering these four groups of materials under the same *ferroic* umbrella is often useful as they share many analogous behavioural characteristics, such as hysteresis curves, phase transitions and the occurrence of switchable domains. However, there are also important differences when considering their respective domain walls. For example, the contrasting relative magnitudes of dipolar exchange and anisotropy energies cause magnetic domain walls to be relatively broad, typically tens of nanometres; in comparison, ferroelectric and ferroelastic domain walls are extremely narrow, frequently with width at the unit cell level⁶⁻⁸. Domain walls in ferroelectrics and ferroelastics can therefore be much more readily considered as genuine 2D objects.

When dimensional confinement is considered in combination with now widespread observations of polar domain walls in non-polar ferroelastics and dramatically enhanced electrical conductivity in domain walls of ferroelectrics, it becomes clear that ferroelectric and ferroelastic domain walls represent a new 2D topological defect with exciting functionalities. In addition, polarisation variations within domain walls lead to quasi-1D topological defects, which differ from those of established polarisation vortex and skyrmion textures⁹⁻¹². Moreover, since domain walls can be created, eliminated, and moved by an external field, the 2D functionality they offer is not fixed but rather can be actively and dynamically deployed. The potential for mobile conducting ferroelectric domain walls to revolutionise nanoscale electronics and the entire way we think about dynamic circuitry is exciting, representing an intriguing electric alternative to concepts applied in domain-wall-based spintronics devices^{13,14}. For memory applications, the possibility to reach very high density of domain walls, up to 1% of the total volume¹⁵⁻¹⁷, with their own properties, implies that high information density could be realised, and that wall-related devices could outperform bulk devices by orders of magnitude.

These concepts were envisioned very early-on¹⁸ and formulated as ‘domain wall nanoelectronics’¹⁹⁻²¹ or ‘domain boundary engineering’²², but progress in developing this new technology has been relatively slow. In general, robust, and definitive information about all the fundamental mechanisms responsible for ferroelectric domain wall conduction (BOX 3) are still unclear for many materials, which is one reason why proof-of-principle devices have emerged only recently. These devices include diodes²³, memories²⁴⁻²⁹ and tunnel junctions³⁰. Another approach is to emulate the behaviour of electronic components, such as digital switches and rectifiers, utilising the complex electronic transport phenomena at domain walls^{31,32}. Polar domain walls in ferroelastics

are even further away from applications: thus far, it has merely been shown that the surface potential resulting from the polarisation in the walls can be read and modified with an electron beam³³ and be used, for example, to tune 2D electron gases at the nanoscale³⁴. In fact, it is likely that memory devices based on polar walls will require first a demonstration of ferroelectricity at the walls, i.e., a switching of the polarisation, before targeting specific application opportunities.

In this review, we discuss the intriguing and non-trivial polarisation configurations that can arise at domain walls and the established mechanisms that can lead to domain wall polarity in non-polar materials. We also provide an overview of the general notions which have been postulated as fundamental mechanisms responsible for domain wall conduction in ferroelectrics. We then suggest that the combination of domain walls with transition regions observed at phase boundaries, homo- and heterointerfaces, and other quasi-2D objects, gives access to novel emergent properties that are yet to be explored.

Our focus is on emergent key functionalities at ferroelectric and ferroelastic domain walls. For a more technical coverage on charged domain walls in these systems, we refer the readers to the review of Bednyakov *et al.*³⁵. We briefly discuss other topological structures in ferroics, which have been reviewed in detail, e.g., by Seidel *et al.*^{36,37}, and we refer the readers to the review of Meier for details on the interplay of ferroelectric and magnetic domain walls in multiferroics³⁸.

Domain wall polarisation in polar and non-polar materials

Phase transitions in ferroelectric domain walls

Intriguing and non-trivial polarisation configurations beyond the bulk properties can arise at domain walls due to their distinct symmetry. Until recently, the simplest Landau theory of only one order parameter (one-component) was enough to understand the available experimental observations and domain wall properties. Such textbook domain walls are approximated by a hyperbolic tangent (tanh) profile of the order parameter (see FIG. 1a, 1c and 1d), possibly with small deviations for higher order Landau potentials^{39,40}. Local alterations of the properties at the wall have been discussed in this context as early as 1970⁴¹, but no additional symmetry breaking and, in particular, no additional polarisation component, is allowed to emerge at these simple walls. In the special case of ferroelectrics, the standard 180° domain wall in this model gives rise to a typical polarisation profile, sometimes referred to “Ising wall”, whereby the polarisation cancels out at the wall (FIG. 1d).

On the other hand, it was proposed very early on that multiple order parameters, or multi-component order parameters, might open the door to a much richer energy landscape populated by exotic wall profiles leading to polarised – and potentially switchable – domain walls^{42–44}. Within this model, a spontaneous polarisation may occur at domain walls, even in an otherwise non-polar material, when the cancellation of the primary order parameter – or instability – at the centre of the domain wall allows a second instability to emerge (FIG. 1b). This domain wall state is bi-stable so that the wall splits spontaneously in two variants^{42,43}. In ferroelectrics, the emergence of a second polarisation component leads to Bloch and Néel walls in analogy to ferromagnets (FIG. 1e and 1f). In contrast to magnetic spins, however, the magnitude of the polarisation vector does not have to be constant and both wall profiles may also

combine polarisation rotations with a reduction of the polarisation, giving rise to complex polarisation profiles. In particular, Bloch- and Néel-type domain walls were heavily researched by various theoretical methods in the 2010s⁴⁵⁻⁵¹. The spontaneous symmetry breaking within the wall required for ferroelectric Bloch walls seems to be rare, because polarisation rotation in ferroelectrics comes with substantial costs in elastic energy. Still, it has been theoretically predicted even in standard perovskite ferroelectrics^{45,49,52-54}, like barium titanate (BaTiO_3), lead titanate (PbTiO_3) and lead zirconate titanate ($\text{Pb}(\text{Zr}_x\text{Ti}_{1-x})\text{O}_3$, PZT), whereby the special role of flexoelectric coupling has also been investigated^{50,51}. These polarisation profiles depend also on the crystallographic orientation of the domain wall normal^{45,46,53,54}.

The emergence of energetically degenerate variants of Bloch- and Néel-type walls themselves suggests the existence of interesting quasi-1D topological defects separating wall regions of opposite polarities, which have been discussed as a paradigm for memory devices^{9,10}. Other topological defects that recently attracted attention in ferroelectrics are polar skyrmions and bubble domains; however, the latter do not require domain walls (BOX 1).

Even though the basic principles have been known for decades, the relevant length scales are just becoming accessible experimentally so that direct evidence for the theoretical concepts is still scarce. One remarkable recent result by Cherifi-Hertel *et al.* is the observation of non-Ising components of the domain wall polarisation in lithium tantalate (LiTaO_3) by optical Second Harmonic Generation (SHG)⁵⁵. Further work along this line is highly desirable in order to understand this signature in detail, and resolve the domain wall structure with atomic scale resolution, as done recently by high-resolution transmission electron microscopy (TEM)⁵⁶, scanning transmission electron microscopy (STEM)⁵⁷ and SHG^{55,57} to reveal Néel-like domain walls in PZT.

Ferroelastics and non-polar materials

While domain walls in ferroelectrics enable unusual polarisation configurations, their impact on the properties in non-polar materials, e.g., ferroelastics and antiferroelectrics, is even more fundamental. Here, domain walls can give rise to an electric polarisation in an otherwise centrosymmetric system. These studies of domain wall polarisations are usually rooted into two complementary approaches: domain wall symmetries and flexoelectricity. One way to describe the symmetries of domain walls is by a so-called layer group, derived systematically using pure geometrical arguments in real space as shown in several works by Janovec⁵⁸⁻⁶⁷. The layer group can also be derived by considering symmetries in order parameter space, which allows a natural connection with Landau theory^{68,69}. This approach notably yields the remarkable result that all strain-compatible ferroelastic domain walls are non-centrosymmetric and predict possible directions for the polar axis⁷⁰.

Experimentally, very few techniques are in fact able to directly determine the full symmetry of a domain wall as a real 3D object. So far, the point group symmetry has mainly been probed by SHG⁷¹⁻⁷⁵, performing a polarisation analysis of the incident and emitted light fields. This approach has produced remarkable results, starting with the experimental confirmation of the non-centrosymmetric symmetry of strain-compatible ferroelastic walls⁷¹⁻⁷⁵, but also observations that are at odds with the classical theoretical approaches. In calcium titanate (CaTiO_3), domain walls were found that have, according to the SHG analysis, a polar axis deviating from the predicted possible

directions (FIG. 3e)⁷¹; in lanthanum aluminate (LaAlO₃), a domain-wall symmetry was proposed ($3m$) that does not match the expected layer-group symmetry⁷². The latter case is particularly surprising as LaAlO₃ is not expected to exhibit any other instability or structural complexity. Future work will have to clarify these situations.

In addition to their polar point group symmetry, domain walls in ferroelastics come with a gradient of one tensor component of the spontaneous strain tensor. This gradient then creates a polarisation via the flexoelectric effect, which is always symmetry-allowed⁷⁶ (FIG 1c). Including this flexoelectric term in the free energy expansion gives a way to model and quantify domain wall polarisation, and many works have used this approach in the context of the recent surging interest in flexoelectric phenomena^{50,51,76,77}. Simulations have been reported at the level of a single isolated wall, as well as for larger volumes with the framework of complex ferroelastic patterns, involving several domain walls (FIG. 3a and 3b)⁷⁶. In the latter, it has been shown that junctions between domain walls and kinks in domain walls are key to obtain a macroscopic polarisation, given that, e.g., in the case of simple stripe domains the opposite direction of the polarisation in neighbouring domain walls cancels out and the macroscopic response vanishes⁷⁶.

All this is of course not exclusive to the symmetry breaking mechanisms described earlier (FIG. 1b). In fact, it is now well understood that two mechanisms exist as two possible reasons for the polarity at ferroelastic domain walls: first, a polarity imposed by the symmetry of strain-compatible domain walls⁷⁰, which can be often described by flexoelectric gradient coupling (FIG. 1c) and, second, the accidental and potentially switchable polarity induced by an additional symmetry lowering⁷⁸ (FIG. 1b). Both phenomena can be addressed together by Ginzburg-Landau type modelling^{69,79,80} in single walls or more complex structures. What will have to be clarified is the significance and respective importance of these phenomena with regard to the potential use of a domain wall polarisation as a functional element, i.e., if and how this polarisation can be detected, manipulated, or switched by electric fields.

Recent experimental works on polar domain walls have concentrated on demonstrating the loss of inversion symmetry^{34,71-75}, quantifying domain wall polarisation by structural studies⁸¹, and looking for evidence for characteristic electric or dielectric signatures of this domain wall polarisation^{11,33,34,82,83}.

In practice, most experimental studies have been devoted to the perovskites and ferroelastic strontium titanate (SrTiO₃) and CaTiO₃. Both are incipient ferroelectrics with a latent polar instability and are therefore prone to exhibit both possible mechanisms for domain wall polarisation (FIG. 1b and 1c). The experimental confirmation that polarity exists in the domain walls of CaTiO₃ came by high-resolution TEM conducted by Van Aert *et al.*⁸¹, where also the polar distortion amplitude was clarified (FIG. 3c), with off-centre displacements of Ti by 6 picometres at room temperature. The loss of inversion symmetry at the walls was later confirmed by optical SHG microscopy⁷¹ (FIG. 3e). Domain wall polarisation in CaTiO₃ and SrTiO₃ was as well invoked as the origin of a significant macroscopic signal in resonance piezoelectric spectroscopy^{11,82} (FIG. 3d), and the sharp contrast observed in low energy electron microscopy (LEEM)^{33,83} (FIG. 3f). In a more original way, domain wall polarisation in SrTiO₃ was also shown to change the LaAlO₃/SrTiO₃ interface current distribution³⁴.

The investigation of polar domain walls in non-polar materials requires the concerted application of these complementary techniques to understand the complex nanoscale physics involved. TEM is the only way to measure the displacement of the atoms away from the high-symmetry position in order to estimate the amplitude of the polar vector within the domain walls (FIG. 3c)⁸¹. Resonance piezoelectric spectroscopy measures frequencies of *macroscopic* mechanical resonances in a sample upon excitation by an AC electric field (FIG. 3d); in the case of materials with non-polar symmetries, these resonances can arise from the motion of polar domain walls^{11,82,84,85}.

Surface states are also generally polar and the polarity in the wall strongly interacts with surface polarity near the intersection between the domain wall and the surface. Furthermore, energy minimisation requires the strain compatibility between surface strains and the strains due to the domain walls. The initial conjecture that this would lead to funnel-like profiles was shown to be wrong and two characteristic excitations were found⁸⁶. These are long-wavelength relaxations in the surface layer and a ‘pinching’ movement of the domain wall, i.e., domain walls slightly narrower near the surface. LEEM is particularly sensitive to such surface effects^{87,88} and enables characterisation of polarisation profiles at the intersection of domain walls with the surface^{33,83}. The LEEM contrasts on the surface arise from polarisation charges, resulting from the polarisation inside domain walls, which deviate the incoming low energy electrons^{33,83}. Injection of higher energy electrons permits screening these charges and removing the contrast, which can be recovered by moderate annealing at 330°C (FIG. 3f).

Despite the recent progress in the characterization of polar domain walls in non-polar materials, the experimental demonstration of the bi-stable polarity of domain walls and its switching has not yet been realised. It remains the most challenging task in the field of polar domain walls in both ferroelectric and non-polar materials, and a crucial step needed towards their potential use as functional elements. Another critical aspect will be a detailed understanding of the control of the walls’ position and motions by means of electric or elastic stress fields. The effect of electric fields on ferroelastic wall motions^{89,90} and more generally the control of domain wall dynamics (BOX 2) will be the key.

Conductive domain walls in ferroelectrics

Enhanced electrical conductivity in ferroelectric domain walls has been reported in several materials, but we are just beginning to understand the underlying nanoscale physics. To reflect the state of the art, we discuss here key observations material by material, highlighting the specificities for each case. The established fundamental mechanisms that lead to (or co-determine) domain wall conduction in all these materials are summarized in BOX 3.

Bismuth ferrite

Bismuth ferrite (BiFeO_3) holds a special place in the history of research on functionally active domain walls: wall conductivity, and even superconductivity, had been reported in previous studies^{91–93} on copper chlorine boracite (Cu-Cl-B) and non-stoichiometric tungsten trioxide (WO_{3-x}), but the first spatially resolved measurements explicitly demonstrating locally enhanced conduction, using conductive atomic force microscopy

(c-AFM), were collected from 109° and 180° domain walls in the rhombohedral phase of BiFeO_3 (FIG. 5a)⁹⁴. A couple of years later, observations of conduction in 71° walls followed in different sets of thin film samples⁹⁵⁻⁹⁷.

Early density functional theory (DFT) based calculations⁹⁸ suggested that the relative changes in conduction between domain walls and bulk, and among the different domain wall types, were associated with a fundamental alteration in the local electronic band structure: models showed that conduction bands were depressed in energy at the walls, while there were only very minor shifts in valence band edges (FIG. 4b). This notion was supported by experiment: scanning tunnelling spectroscopy measurements⁹⁷ were used to directly probe the density of states as a function of location (traversing both 109° and 71° walls). The errors in band edge energy determination were significant, but a noticeable dip in the energy of the conduction band edge at the 109° domain wall was seen, while meaningful signs of valence band edge changes were not. These reports pointed to observations of conductivity being the result of local changes in band structure (and reduced band-gaps) at the domain walls. However, it has been contested as to whether these local changes are significant enough to cause the increase in conductivity observed⁹⁹.

Attempts to elucidate fundamental band structure alterations through direct current-voltage measurements have often been deeply frustrated by the two-probe geometry inherent to the c-AFM technique: conduction has consistently been found to be dominated by the barrier resistance between the tip and the material, with both Schottky and Fowler-Nordheim behaviour being claimed^{95,96,100}. It has been suspected, particularly in work by Farokhipoor and Noheda^{95,96}, that barrier height reductions occur because of enhanced oxygen defect concentrations at the domain walls¹⁰¹ (FIG. 4a); Maksymovych *et al.*¹⁰² acknowledged that such vacancies should certainly be present, but consider that the multistate resistance levels induced under applied fields, in their work, had response times deemed too rapid to be caused by oxygen vacancy migration. Instead, they proposed that microscopically reversible distortion of the polarisation structure at the domain wall, induced by the voltage applied to the c-AFM tip during the measurement, might have been responsible for changing the conduction behaviour. Polarisation discontinuities and associated strong local electric fields have also been seen to play a role (FIG. 4c) and complex polarisation patterns have been observed using high-resolution STEM¹⁰³. Furthermore, c-AFM measurements on written domain structures, designed to generate abutting polarisation patterns, have shown that “head-to-head” walls (FIG. 4c) conduct while “tail-to-tail” walls (FIG. 4c) do not¹⁰⁴. In addition, it has been recently argued that strain-compatible domain walls in BiFeO_3 localize free excess electrons and create intrinsic polaronic in-gap states¹⁰⁵. This mechanism might dominate the thermally activated domain-wall conduction, e.g., in thin film materials with a high density of domains walls.

An overriding observation is that BiFeO_3 research has always been plagued by uncontrolled and often uncharacterised point defects: the presence of the transition metal (Fe) and the associated degree of freedom in valence state readily facilitate non-stoichiometry. Rojac *et al.*¹⁰⁶ used STEM techniques to reveal significant concentrations of Fe^{4+} ions at BiFeO_3 domain walls, resulting from the accumulation of bismuth vacancies. Mixed Fe^{3+} - Fe^{4+} valence states offer possibilities for conduction through electron hopping, suspected as key in conductivity measurements performed by Lee *et al.*¹⁰⁷. Perhaps the most compelling evidence for the dominant role of defect accumulation in generating enhanced conductivity at domain walls in BiFeO_3 can be

found in the work of Stolichnov *et al.*¹⁰⁸: they found that, when conducting domain walls were moved using applied electric fields, enhanced conduction persisted where the domain walls had been, prior to field-induced movement, indicating that the sheet conduction and the wall could be decoupled.

Actually, the enhanced electrical conduction in BiFeO₃ domain walls is likely the result of all the mechanisms discussed in FIG. 4, making it both a complex and versatile system. Going beyond just enhanced conductivity, magnetotransport^{107,109–111} was studied at domain walls in BiFeO₃, expanding possible domain wall applications into the realm of spintronics.

Hexagonal manganites

A breakthrough concerning the disentanglement of conduction contributions was the discovery of anisotropic transport properties at ferroelectric domain walls in hexagonal (h-) manganites [$RMnO_3$ ($R = Sc, Y, In, Dy$ to Lu)]^{112–114}. Hexagonal manganites exhibit uniaxial (improper) ferroelectricity and naturally develop all fundamental types of 180° domain walls, that is, neutral side-by-side, negatively charged tail-to-tail and positively charged head-to-head walls (FIG. 5c). The walls are stabilized by topologically protected six-fold anchor-points^{112,115}, which prevents the material from forming, e.g., simple stripe-like domains with neutral walls. In 2012, Meier *et al.* performed local transport measurements and showed that the conductivity at the charged domain walls originates from bound charges and the related electrostatic potential, locally modifying the electronic band structure (intrinsic)^{113,116,117}, analogous to the scenario depicted in FIG. 4c. It is a typical example where the wall conductivity depends on the crystallographic orientation of the domain wall normal.

At tail-to-tail domain walls, the mobile majority carriers (holes) feel the pronounced electrostatic fields from the negative bound charges, leading to hole accumulation and, hence, enhanced hopping conductance (p-type). Turner *et al.* revealed that the associated carrier mobilities are surprisingly high, in the order of $100 \text{ cm}^2 \text{ V}^{-1} \text{ s}^{-1}$ at room temperature, and amongst the highest reported in oxide systems¹¹⁸. Accordingly, the positive bound charges at head-to-head domain walls are screened by hole depletion, leading to reduced conductance. Recently, it was observed that head-to-head domain walls can also deliver enhanced conduction over bulk, but only under higher electric fields³¹. It is clear that, in this case, the active carriers are n-type^{31,118}. Such carriers arise from a significant band bending with the conduction-band minimum dipping below the Fermi energy, which leads to the formation of an electronic inversion layer³¹ (FIG. 4c). Thus, the origin of conduction at charged domain walls in hexagonal manganites is well-understood. At the quantitative level, however, there are still some discrepancies between the calculated and measured carrier densities and their participation in transport^{31,118}. Such discrepancies likely arise from variations in the domain-wall charge state “hidden” within the bulk¹¹⁹ and secondary contributions due to interactions between the domain walls and point defects.

In contrast to the charged domain walls, neutral side-by-side domain walls in h- $RMnO_3$ (FIG. 5c) show a similar electronic structure as the bulk¹²⁰. Their transport properties are extrinsic and arise from the complex defect chemistry at domain walls, ranging from insulating¹¹² to conducting¹²¹ (FIG. 4a). For example, it has been demonstrated that oxygen interstitials have a propensity to form at neutral domain walls³², giving rise to enhanced DC conduction (p-type) and increased AC conductivity at frequencies in the

kilo- to megahertz range³². Higher drive frequencies in the terahertz regime lead to dynamical domain-wall displacements, resulting in anomalous microwave AC conductivity at neutral domain walls¹²².

Barium titanate

Another ferroelectric where the high conductivity is explained within the scenario of FIG. 4c is BaTiO₃. In this system, Sluka *et al.* have used macroscopic current measurements to show that, at head-to-head domain walls, the static conductivity can increase by 9 orders of magnitude¹²³. These strongly charged domain walls are more difficult to obtain than in improper ferroelectrics, like hexagonal manganites, because of their high energy cost¹²⁴, but they are stabilized by the strain-compatibility of the adjacent ferroelastic domains and an almost perfect compensation of the bound polarisation charges by free carriers, which are at the origin of the high conductivity. In BaTiO₃, and ferroelectrics in general, such compensation cannot be provided by the free carriers readily available in the material but originates from electron transfer across the bandgap or external charge injection. At low voltages (< 8 V), the free carriers in the wall are separated from the electrode by an insulating gap due to the flattening of the electric potential profile by the equipotential electrodes and large voltages are required to observe the electron transfer and an increase in conductivity¹²³. At tail-to-tail walls, experiments show that the conductivity is like the bulk values, which might arise from electron depletion^{124,125}. Also, oxygen vacancies that are screening tail-to-tail walls, in particular in samples with high oxygen deficiency¹²⁶, exhibit a low mobility compared to electrons and hence lead to a low conductivity.

Lithium niobate

Thus far, the higher static conductivity of domain walls in the uniaxial ferroelectric lithium niobate (LiNbO₃), observed in an early study of Aristov *et al.*¹²⁷, has been attributed to partially charged domain walls, analogous to the scenario depicted in FIG. 4c, but where the domain walls are inclined by a few degrees instead of being vertical. On a macroscopic scale, Schröder *et al.* have shown using c-AFM on both faces of single crystals that domain walls in LiNbO₃ are inclined with respect to the ferroelectric polarisation direction and that this inclination can be controlled by a change in dopant concentrations, e.g., magnesium (inclination up to 0.3°)¹²⁸. The inclination, confirmed by Čerenkov second-harmonic generation measurements (FIG. 5d)^{129,130}, results from an incomplete poling of the material and can be much larger in sub-micrometre thick samples (inclination up to 20°)^{27,131}. Similarly, inclined domain walls have also been observed in LiTaO₃ by introducing a composition gradient along the polarisation axis, which changes the temperature of the transition to the paraelectric phase and the coercive field^{132,133}. Such a strategy should be successful in LiNbO₃ as well but has not been demonstrated yet.

Inclined domain walls lead to partial head-to-head configurations with an accumulation of free electronic carriers, leading to an intrinsic rise in conductivity¹³⁴. The Landau-Ginzburg-Devonshire theory predicts an increase of conductivity by one order of magnitude at small angles (0.07°)¹³⁴, which could be obtained in the first experiments only with additional carriers provided by illumination at an energy larger than the bandgap¹²⁸. The predicted conductivity across tail-to-tail walls, which should be at least an order of magnitude smaller than the one for head-to-head walls due to the low mobility of holes compared to electrons¹³⁴, has not been observed experimentally yet. It was

realised later by Eng and Gruverman that, close to the surface, there is an additional bending of the upper segment of the wall, away or toward the polarisation direction, which can be controlled by the application of sub-coercive voltage (FIG. 5d – right inset)^{131,135}. This surface effect leads to strong variations in carrier density in the first 100 nm and dominates the overall conductivity measured by c-AFM, which can then be tuned by 2 orders of magnitude^{131,135}. Similar domain wall bending is expected in most ferroelectrics but has not been investigated yet. Additional free carriers that are excited while probing the walls with c-AFM can further give rise to large AC conductivity¹³⁶.

In fact, the conductivity at domain walls is likely to be even higher than the value obtained through these c-AFM measurements (conductance ~ 10 pS) where substantial contact barriers at the tip lead to non-ohmic behaviour^{128,137}. As such, the work of Werner *et al.*¹³⁷ is very instructive. It shows a 4 orders of magnitude difference between the current mapped with c-AFM (~ 1 μ A) and the current recorded through macroscopic electrodes contacting several domain walls (~ 10 mA), leading to a real enhancement of the conductivity at domain walls by 13 orders of magnitude compared to that of the bulk.

On the nanoscale, Gonissen *et al.* have shown with STEM that domain walls in LiNbO₃ are not straight but exhibit meanders and kinks, which result in local head-to-head or tail-to-tail sections where bound charges accumulate (FIG. 5d – left inset). Compared with PZT⁷ (FIG. 5b), these sections are spatially less extended (< 10 nm), but it has been argued that the charges could be screened by stabilized defects^{138,139}, which could play an extrinsic role in the enhanced conductivity of LiNbO₃. Further work is highly desirable to investigate this effect.

Lead zirconate titanate

Another system that underlines the complexity of domain-wall related transport phenomena in ferroelectrics is PZT. In 2008, Jia *et al.* showed that, at the atomic scale, both neutral side-by-side and charged head-to-head 180° domain walls arise in SrTiO₃/Pb(Zr_{0.2}Ti_{0.8})O₃/SrTiO₃ trilayers⁷. In particular, they observed that domain walls exhibit local head-to-head sections, where the wall tilts away from the ideal neutral structure (FIG. 5b). As discussed in the previous sections on, e.g., hexagonal manganites and LiNbO₃, this tilting results in domain wall bound charges that require screening (FIG. 4c), for example, by oxygen vacancies (extrinsic) or electronic carriers (intrinsic), possibly stabilized by a local change in the Ti valence state from Ti⁴⁺ by Ti³⁺ (REF.^{7,140}). **These excess electrons on Ti sites, which experience a strong on-site Coulombic repulsion, could take the form of localized small polarons¹⁴¹.**

The formation of domain walls with neutral and head-to-head sections later became the key to understand the thermally activated transport at 180° walls in simple Pb(Zr_{0.2}Ti_{0.8})O₃ thin films^{142–144}. Guyonnet *et al.* proposed that the charged sections exhibit an increased defect density (oxygen vacancies), providing trap states for hopping mediated transport along walls (FIG. 4a)¹⁴². To enable *extrinsic* defect-mediated conduction, a certain threshold density of oxygen vacancies is to be reached, as the same team demonstrated by controlling growth conditions and atmosphere¹⁴⁵. Here, the highly localized electric fields, which are used to create the domain walls with c-AFM, come into play. Such fields may drive the reorganization/injection of oxygen vacancies, which then preferentially segregate to newly formed walls. Additional free

carriers that are excited while writing the walls with c-AFM can further enhance the DC conductance or, at high frequencies (gigahertz), oscillate between barriers, giving rise to large AC conductance (~ 100 nS)¹⁴⁶.

Even higher, metallic conductance, was achieved by writing nanodomains with strongly curved walls and, hence, a large fraction of charged segments¹⁴⁷. The work on strongly curved walls showed that both electron tunnelling and metallic conductivity play a role in PZT and confirmed the strong correlation between domain wall conduction and curvature – a concept that was then expanded to nominally neutral 90° domain walls⁷: by strain engineering, 90° domain walls in PZT were forced to bend away from their mechanically compatible position, inducing partially charged walls with metallic-like conductivity^{148,149}. Importantly, the 90° domain walls were generated at $T = 4$ K, that is, a temperature at which defects cannot migrate, which led to the conclusion that the conduction arises from *intrinsic* band bending effects (FIG. 4c). More recently, state-of-the-art microscopy studies provided additional insight, revealing the emergence of complex Néel-like polarisation rotations (FIG. 1f) across domain walls in PZT, which can locally violate charge neutrality and promote domain wall conductance^{55–57}.

The research on PZT highlights the importance of domain wall bound charges for conductivity to arise (see FIG. 4c). However, the charged sections are usually “hidden” within the bulk so that it is challenging to establish a quantitative relation between their density and the domain wall conduction. In addition, beyond walls with charged sections, more complex polarisation rotations emerge, and both intrinsic and extrinsic contributions appear to play role for the domain wall conduction, critically depending on small variations in the domain wall orientation and growth conditions.

In summary, the electronic transport at domain walls is studied intensively. So far, however, most application oriented studies focus on controlling the domain wall behaviour via electric fields and defect densities; other intriguing phenomena like the interaction of light with domain walls is still a rather overlooked topic that has only been studied in LiNbO_3 ¹²⁸ and BiFeO_3 ^{17,111,150–153}. Further studies, taking advantage of near-field techniques, e.g., tip-enhanced Raman spectroscopy, which overcome the diffraction limit and give the possibility to probe domain walls with a reduced bulk contribution, are desirable¹⁵⁴. Another promising field that deserves further attention concerns the local interaction of electric and magnetic degrees of freedom, leading to unusual correlation phenomena¹⁵⁵ and domain wall magnetotransport^{107,109–111}.

Combining domain walls and transition regions for emerging properties

Beyond conductivity and polarity at domain walls

From a more conceptual point of view, domain walls represent transition regions of the order parameter. As such, they can exhibit a wide range of unusual properties, going way beyond the changes in conductivity and polarity discussed in the previous sections. For example, at simple Ising-type walls (FIG. 1d), the polarisation is suppressed and the free energy exhibits a local maximum with a negative curvature¹⁵⁶. This specific profile of the free energy across a domain wall, predicted by Landau theory, has been observed experimentally through a precise mapping of polarisation and local electric fields in a $\text{SrTiO}_3/\text{PbTiO}_3$ superlattice¹⁵⁶. As a direct consequence of the unusual curvature of the energy landscape, the walls exhibit negative permittivity. The effect

can be boosted by engineering dielectric/multidomain ferroelectric superlattice, leading to a situation where the negative domain wall permittivity dominates the macroscopic response of the system¹⁵⁷. The approach allows for designing superlattices with improved permittivity, which is higher than for the individual constituent dielectric (SrTiO_3)^{156,158}, enabling a new generation of low-energy field-effect transistors¹⁵⁹.

Furthermore, at the atomic scale, transition regions of the order parameter correlate with structural changes, altering interatomic distances and, hence, the electronic bonds and the local strain state¹⁶⁰. Such structural inhomogeneities associated with domain walls have a direct impact on the propagation of phonons. Royo *et al.*¹⁶¹ have demonstrated that domain walls can act as phonon polarizers and filter phonons depending on their polarisation¹⁶¹. In the model case of tetragonal PbTiO_3 , they demonstrate that 180° domain walls scatter phonons associated with atomic displacements along the direction transverse to a heat flux, while phonons associated with atomic displacements along the longitudinal direction are transmitted¹⁶¹. Thus, a single domain wall can lead to a reduction of the thermal conductance by $\sim 43\%$, essentially through the scattering of low-frequency phonons. Beyond changes in thermal conductance, such a domain-wall-based phonon polarizer could find applications in devices relying on nonradiative energy relaxation and inter-band recombination processes, which are governed by polarisation-dependent selection rules.

Strain gradients at domain walls can also lead to the emergence of chemically novel 2D-phases that have not yet been synthesised by standard chemical routes and remain a specificity of domain walls¹⁶². Such unusual chemical phase has been reported at the ferroelastic domain walls of multiferroic orthorhombic TbMnO_3 thin films¹⁶². It arises from the local substitution of Tb atoms by Mn atoms on almost all atomic sites. Remarkably for applications in spintronics¹⁶³ and magnonics¹⁶⁴, the corresponding local increase in Mn content changes the local magnetic properties and leads to a positive in-plane magnetic moment per atomic plane parallel to the domain wall, while the domains are antiferromagnetic¹⁶². Another example taking advantage of a different chemistry at domain walls is the ‘nano-capacitors’ engineered by defect-chemistry in strained SrMnO_3 thin films¹⁶⁵. A comparison of c-AFM conductance maps with SHG imaging measurements reveals electrically decoupled conducting nanoarrays of polar domains delimited by insulating walls, which can be individually charged. Complementary density functional theory calculations indicate that the walls are insulating and act as a barrier to conduction because of the accumulation of oxygen vacancies driven by local strain¹⁶⁵.

Thus, the transition region of order parameters associated with domain walls is highly relevant both from a fundamental as well as a technological point of view. This has also been realised by other communities, which is currently leading to an expansion into adjacent research fields with intriguing synergetic effects. As such, there has been a strong interest in properties of phase boundaries, homo- and heterointerfaces, and other quasi-2D objects, all described here-after as transition regions.

Transition regions

One example of a conceptually new type of transition region that falls into this category is the epitaxial trilayer heterostructure $\text{PZT}/\text{La}_{0.7}\text{Sr}_{0.3}\text{MnO}_3/\text{PZT}$, which can be described as an homogeneous artificial material with inversion-symmetry properties¹⁶⁶. Indeed, the interlayer ($\text{La}_{0.7}\text{Sr}_{0.3}\text{MnO}_3$) offers the possibility to switch the polarisation

direction of the top PZT layer without changing the polarisation direction of the bottom PZT layer. When the polarisations of the two PZT layers are antiparallel, the material exhibits an inversion symmetry while when they are parallel, the inversion symmetry is broken. Such trilayer systems can thus act as nano-sized light emitters, where a SHG output signal – correlated with the loss of inversion symmetry – can be controlled by applying a voltage¹⁶⁶. Subsequent experiments have demonstrated the generality of this approach, which is robust against an exchange of the control ferroelectric layers as well as the oxide interlayer. The strategy opens the possibility to alter other properties of the interlayer or its interfaces and transfer what has been learnt from charged head-to-head and tail-to-tail walls to thin film heterostructures and multilayers. To optimize such structures, the control of interfaces will be critical and could rely on the accumulated knowledge acquired through the study of multiferroic heterostructures. Here, ionic, electronic, strain and exchange couplings at interfaces result in transition regions, often down to a couple of atomic layers only, both in the ferroelectric and ferromagnetic materials^{167–174}.

Symmetry breakings have also been observed in homogeneous thick films^{175,176}. In this regard, the recent discussion in a work of Noheda on so-called “transitional ferroelectrics” is appealing¹⁷⁶. The denomination comes from the observation of a gradual change of structure from a tetragonal symmetry at the top of a thick film of BaTiO₃ to an orthorhombic symmetry at the bottom¹⁷⁶. The transition region between both phases is characterized by a reduced symmetry where the polarisation rotation is coupled to the strain gradient (FIG. 6a). The ease with which the polarisation rotates in this transition region under small electric fields leads to huge and temperature-independent dielectric responses (> 4000) and large piezoelectric responses¹⁷⁶. These exceptional properties arise from the unusual characteristics of the transition region, which is not a polycrystal but also not a single crystal, not a relaxor but also not a homogeneous ferroelectric.

A remarkably similar observation has been reported in free standing single-crystal thin membranes of BaTiO₃. In these systems, under increasing stress due to the mechanical bending of the membrane, the tetragonal domains evolve from *c*-domains (out-of-plane) to *a*-domains (in-plane), with a continuous rotation of the dipoles between both domains (FIG. 6b)¹⁷⁷. This transition region where the dipoles rotate is thought to be at the origin of the exceptionally large flexibility and super-elasticity (recoverable strain ~10%) of these membranes since it could eliminate most of the mismatch stress between domains¹⁷⁷. Also, it resembles the transition region discussed in “transitional ferroelectrics” but with tetragonal nanodomains as building blocks, instead of a phase with reduced symmetry.

It is highly likely that these transition regions will additionally exhibit unusual evolutions with temperature. In PZT, ab-initio calculations have been used by Nahas *et al.* to characterize the evolution of the ferroelectric domain structure with temperature¹⁷⁸. On heating, an initial labyrinthine domain pattern, which consists of convoluted stripes and meandering domains, evolves into a less-symmetric parallel-stripe state before to undergo the transition to the more-symmetric paraelectric phase (FIG. 6c). The demonstration of such an inverse transition¹⁷⁹, i.e., a transition to a less-symmetric domain state on heating, is a first in ferroelectric oxides and is in agreement with the experimental evolution of domains in BiFeO₃ upon cooling from different high temperatures¹⁷⁸.

The remarkable communality between the different transition regions described above and their evolution with driving fields (temperature, stress) is that they lead to exceptional properties and coupling phenomena. Also, it is highly likely that they can be triggered by light and will react to mechanical strain, and thus provide interesting piezo-photovoltaic properties¹⁸⁰. The existence of such regions relates to different origins or driving forces, ranging from chemistry, phase coexistence, frustrations, fluctuations, and chiral textures, all of which can be in principle encountered in or around domain walls. Therefore, most of these transition regions probably exist in the vicinity of domain walls, at a smaller length scale, and shall receive more attention to foster mutually beneficial research. The fact that the inverse transition described previously finds its origin in the diffusion and relaxation of topological defects localized at the junction of different domains¹⁷⁸ is a striking example of the possibilities offered by such cross-fertilisation. One could also ask if the recent superconductivity observed in an “infinite-layer” nickelate of specific chemistry¹⁸¹ could be observed as well in the domain walls of rare-earth nickelates, which constitute 2D objects potentially stabilizing defects, and have been less studied than in other oxides¹⁸². No doubts that superconductivity, super-elasticity, giant dielectric properties at domain walls are in reach.

Outlook

For future investigations, we believe that insights into the full 3D structure of the domain walls, down to the nanoscale, are highly desirable in order to answer open questions about the microscopic origin of the intriguing functionalities. Experimental techniques, such as dark-field X-ray microscopy, which can map lattice distortions around domain walls in bulk samples¹⁸³, or 3D atom-probe tomography, which could provide access to the chemical identity of individual atoms at domain walls¹⁸⁴, would be relevant. In particular, they might help to understand the role of point defects interacting with domain walls and disentangle extrinsic and intrinsic effects to gain reliable quantitative information. The latter is important when moving towards domain-wall based devices, where well-defined and deterministic behavior is critical.

Regarding applications, the standard idea is to create, move and eliminate domain walls with an external field to control the functionality. This approach, however, relies on classical device geometries, e.g., where conductive domain walls are used as interconnects between electrodes. A conceptually new approach is to keep the walls static and use their versatile properties to control electronic signals. Going beyond just conductance, by driving, for example, metal-insulator transitions or magneto-resistive changes within domain walls represents a promising pathway. Different domain walls could emulate the behavior of different electronic components^{31,32}, representing the building blocks of domain-wall based networks and circuitry, where the atomic-scale dimension associated with the domain walls is fully taken advantage of instead of using them as active components in much larger devices. With this, we believe that domain walls could play a key role in the transition from nano- to atomic scale electronics.

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Author contributions

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Competing interests

The authors declare no competing interests.

Box 1 | Topological defects in ferroelectrics

A topological defect is a localized discontinuity of the order parameter that cannot be removed by a local continuous change in the order parameter configuration^{11,185}. When localized in one direction but extended in the two other directions, e.g., as a domain wall, it is a *planar defect*. The line separating regions of opposite polarities within a domain wall, e.g., in a Bloch wall, is a *line defect* (FIG. 2a)^{9,10}. Domain walls, as well as line defects within domain walls, are not the only topological defects that attract the attention of the ferroelectrics' community. Inspired by magnetic textures, researchers explore now properties of other non-trivial ferroelectric structures – often described as topological defects – consisting of arrays of individual polarisation flux closure vortices (FIG. 2b)¹⁸⁶, polar-skyrmion bubble domains (FIG. 2c)¹⁸⁷ and perhaps also ferroelectric Bloch skyrmions¹⁸⁸, that could be the most direct counterparts of the magnetic Bloch skyrmions known from thin films of chiral magnets¹⁸⁹.

In thermally activated systems, simulations reveal highly mobile polar vortices¹⁹⁰. Their lifetime is short, and these vortices are prone to be destroyed by applied electric fields¹⁹⁰, except when they are attached to domain walls, surfaces or defects^{191,192}. Most intriguingly, junctions between domain walls and kinks in domain walls also generate polarisation vortices^{76,193,194}. Under motion of the wall, such vortices are expected to generate a magnetic field in nonmagnetic materials by rotating polar vectors inside domain walls¹⁹⁵. Experimentally, only very weak magnetic signals have been observed¹⁹⁶ and more research is urgently needed.

Box 2 | Domain wall dynamics: avalanches in ferroelectrics

The dynamical behaviour of domain walls is subject to intense research because it is at the core of ferroelectric switching^{197–200}. One approach relies on the study of individual domain wall movement on long-time scales. This is the initial phenomenological method of Ishibashi and Takagi^{201–203} and the developments that followed^{204,205}, along with optical microscopy observations^{154,206} and more recently piezoresponse force microscopy measurements^{207–211}.

Another approach aims to understand collective movements across short-time scales. It considers discrete, impulsive jumps, occurring during the motion of domain walls, as indicators of avalanches on a broad range of scales. A typical example is the movement of domain walls in BaTiO₃ during ferroelectric switching²¹². In this experiment, the pinning and depinning of domain walls generates strain relaxation. This relaxation is fast so that the strain forms waves which can be observed as acoustic signals at the sample surface. These strain waves (coined ‘jerks’) are detected by piezoelectric sensors of high sensitivity and broad frequency response²¹². The transmitted signal is then statistically analysed. The individual jerks show energy and amplitude distributions that indicate avalanche processes^{213–216}, and both parameters are power law distributed with an exponent near 1.7 for the energy. This exponent is close to the typical exponent found in other ferroelastic materials^{217–221}. A similar energy exponent has been confirmed through simultaneous displacement current and birefringence measurements²²².

Thus, the collective movement of domain walls as avalanches is scale invariant, which means that even extremely fast movements – below the cut-off frequency of the experimental equipment – are likely. This puts some limitations on the use of domain walls at high frequency (typically GHz) where random noise may prevent switching operations. One possibility is to extend the operating temperature to low temperatures. Measurements in ferroelastic SrTiO₃ in the quantum critical state show that domain movements do indeed change dramatically with temperature²²³. Between ~40 K and 25 K the domain configuration becomes frozen into a glass state where the domains slide over a complex defect landscape similar to a spin-glass behaviour^{224,225}. Below 25 K a quantum domain solid emerges where the domains form elastic solid structures which interact weakly with the underlying matrix. Moving these structures requires weak thermal activation, which vanishes when the absolute zero temperature is approached²²³.

Box 3 | Conduction mechanisms

Enhanced conductivity at ferroelectric domain walls finds its origin in distinct mechanisms that may operate either in concert, or individually, depending on the specific ferroelectric involved, the domain wall type, and the processing used for synthesis. First, domain walls aggregate (or repel) point defects that mediate conduction either through the creation of intra-bandgap electronic donor or acceptor states or through ionic motion of the defects themselves along the wall (FIG. 4a). Second, the intrinsic electronic band structure of domain walls is fundamentally different from the bulk (irrespective of point defects), because the atomic and associated electronic configurations are locally distinct – this could allow meaningful reductions in the bandgap to facilitate significant increases in thermally excited conduction electrons (FIG. 4b). Third, local electric fields associated with polarisation discontinuities across the wall, such as head-to-head and tail-to-tail configurations, uniformly shift the absolute energies of the electronic band structure locally (through band-bending) allowing conduction bands to dip below the Fermi energy of the system as a whole, or valence bands to peak above it (FIG. 4c).

The three mechanisms mentioned above, which occur in the bulk of the material, can also arise and be enhanced at the junction between domain walls and the surface. For ferroelastic domain walls, it has been shown that the spontaneous strain gradient that exists in the vicinity of the domain wall/surface junction decreases the local band gap²²⁶ (mechanism of FIG. 4b). This gradient also creates a polarisation via the flexoelectric effect (similar to the mechanism described in FIG. 1c in the bulk) and hence an electric field that leads to an increase of free carriers²²⁶. Furthermore, at domain wall/surface junctions, the accumulation of defects is favoured²²⁶ (mechanism of FIG. 4a). In ferroelectrics, the thickness of the walls changes at the surface with potential implications on the local distribution of bound charges and free carriers (mechanism of FIG. 4c): the polarisation discontinuity at the surface leads to a broadening of the domain walls²²⁷, which depends also on the inclination angle for partially charged domain walls²²⁸.

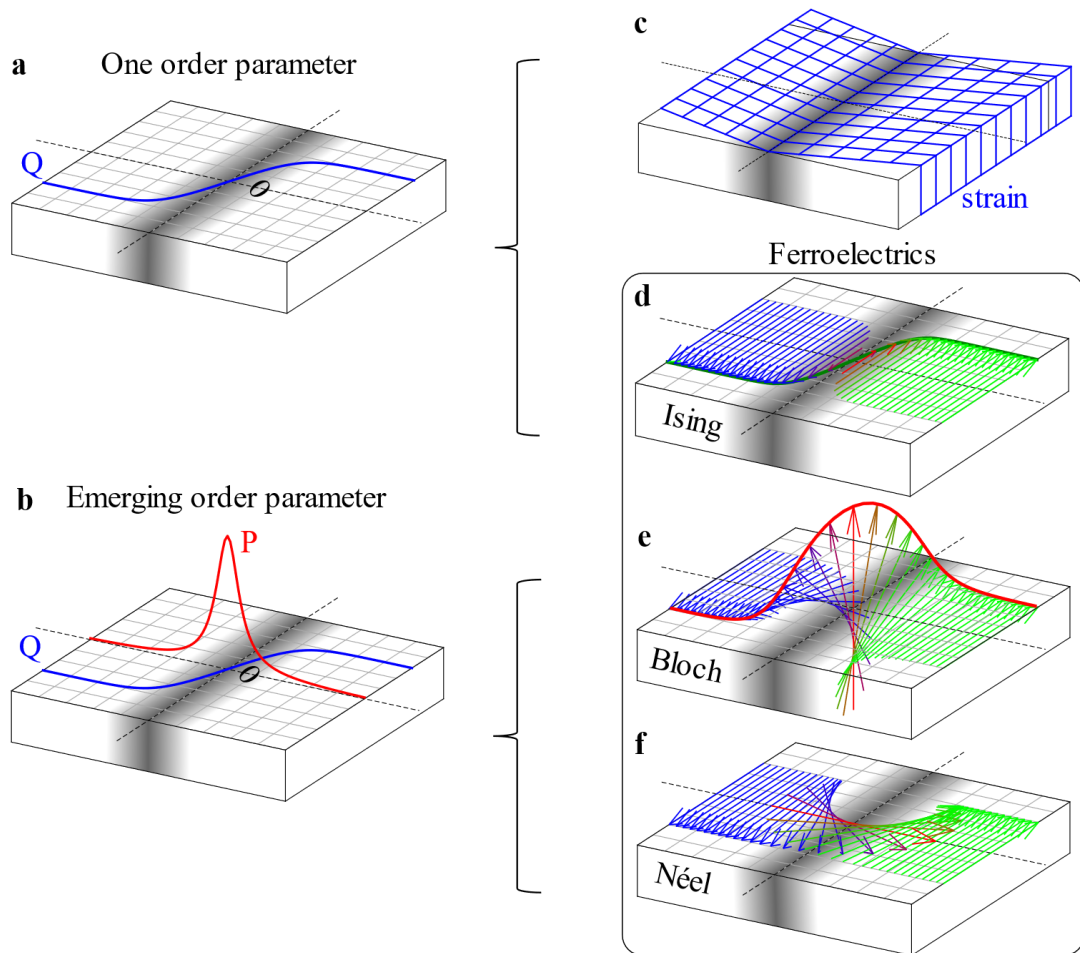
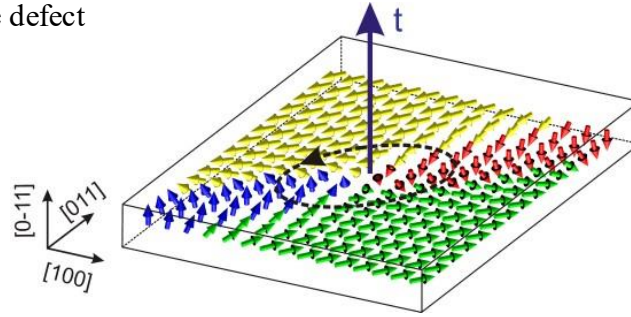
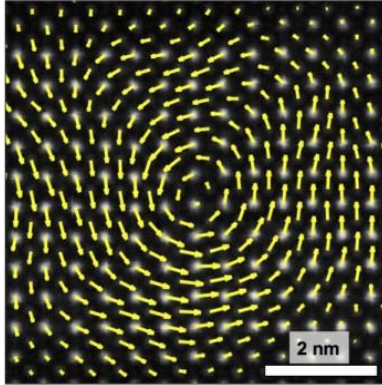


Fig. 1. **Mechanisms leading to polar domain walls.** **a** | Schematic of an elementary domain wall with a single order parameter and tanh profile. **b** | Phase transition in a domain wall, with the emergence of an additional instability, which may be a polarisation component P . **c** | Sketch of a ferroelastic wall, with a discontinuity (or gradient) in spontaneous strain. **d** | Ferroelectric elementary Ising wall. **e** | Ferroelectric Bloch wall, where the polarisation rotates in the plane of the wall. **f** | Ferroelectric Néel wall.

a Ising line defect



b Polarisation flux closure vortex



c Polar-skyrmion bubble

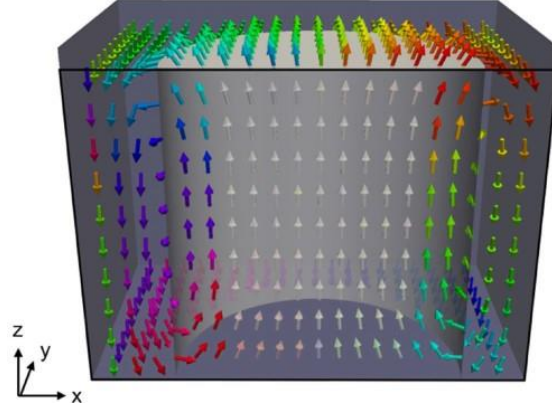


Fig. 2. **Examples of topological defects in ferroelectrics beyond domain walls.** **a** | Ising line defect in rhombohedral BaTiO_3 as obtained from the model described in REF.¹⁰. When circumventing the core of this defect following a closed path (dashed line) around its axis t , the polarisation on this path rotates around the $[-211]$ axis, which is perpendicular to t . Reproduced with permission from REF.¹⁰. **b** | STEM polar displacement vector map of a single vortex in a $(\text{SrTiO}_3)_{10}/(\text{PbTiO}_3)_{10}$ superlattice. Reproduced with permission from REF.¹⁸⁶. **c** | Three-dimensional cross-sectional view of a polar-skyrmion bubble. Reproduced with permission from REF.¹⁸⁷.

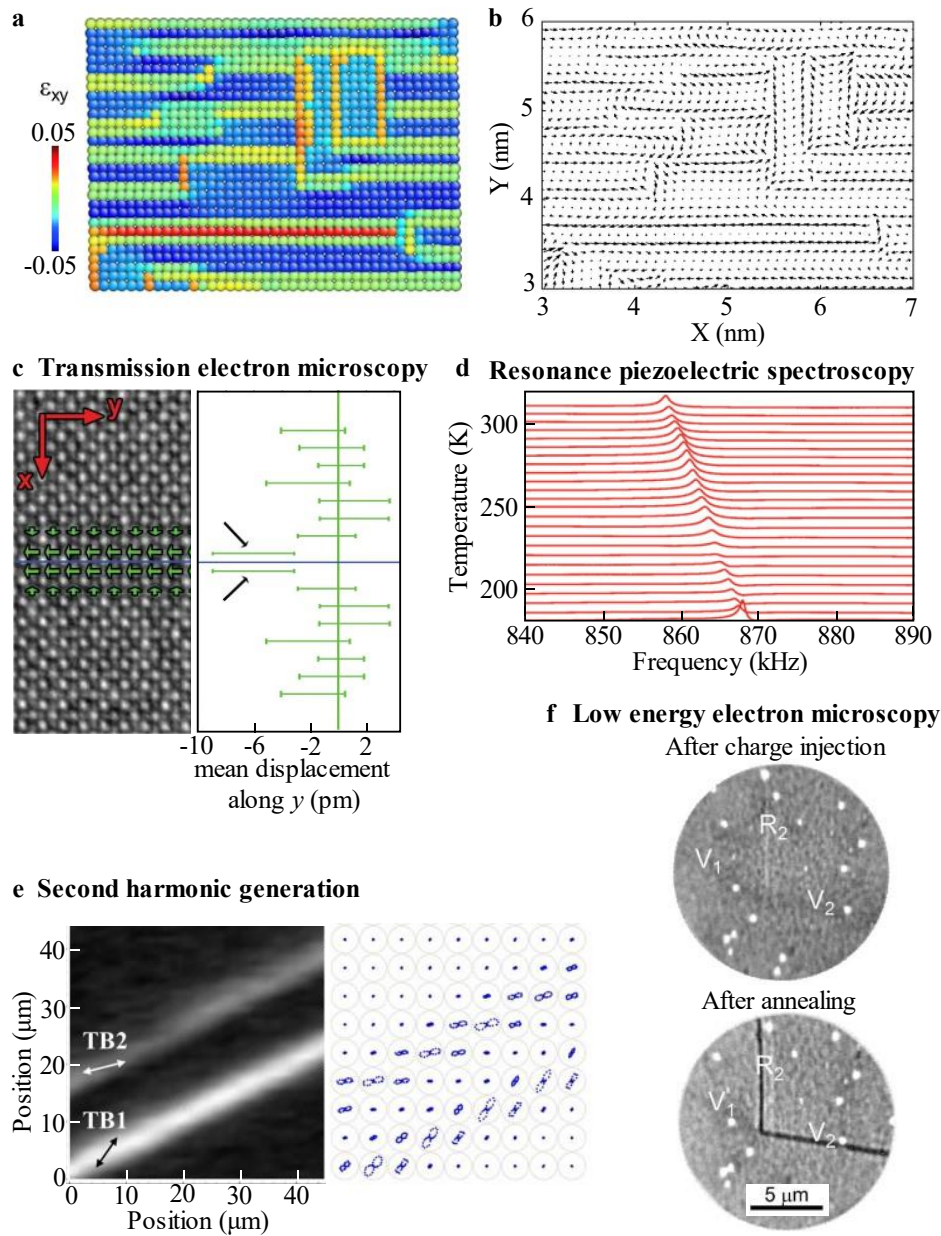


Fig. 3. Polar domain walls in non-polar CaTiO_3 . **a** | Atomistic toy model of a domain pattern with multiple domain walls in view of the atomic shear strain ϵ_{xy} (indicated by the colour code). **b** | Resulting dipole configuration (indicated by black arrows) along domain walls and vortices near domain wall intersections. Dipole displacements are amplified by a factor of 25. The junction density is high leading to extended vortex structures in P . Reproduced with permission from REF.⁷⁶. **c** | On the left, phase of the TEM reconstructed exit wave where the mean displacements of the Ti atomic columns from the centre of the four neighbouring Ca atomic columns are indicated by green arrows. On the right, displacements of Ti atomic columns in the y-directions averaged along and in mirror operation with respect to the twin wall together with their 90% confidence intervals. Adapted with permission from REF.⁸¹. **d** | Resonance piezoelectric spectra of a single crystal CaTiO_3 sample. An AC electric field of 50 V mm^{-1} was applied to obtain the spectra. The spectra are vertically translated so that the axis is labelled as temperature. Adapted with permission from REF.³³. **e** | SHG image of an area that contains two SHG active domain walls: TB1 and TB2. The corresponding polar diagrams mapping is shown in the right panel. Adapted with permission from REF.⁷¹. **f** | Low energy electron microscopy images containing three domain walls (V_1 , V_2 , R_2) after 10-min exposure to 20 V e-beam, and the same regions after annealing at 330°C . Adapted with permission from REF.³³.

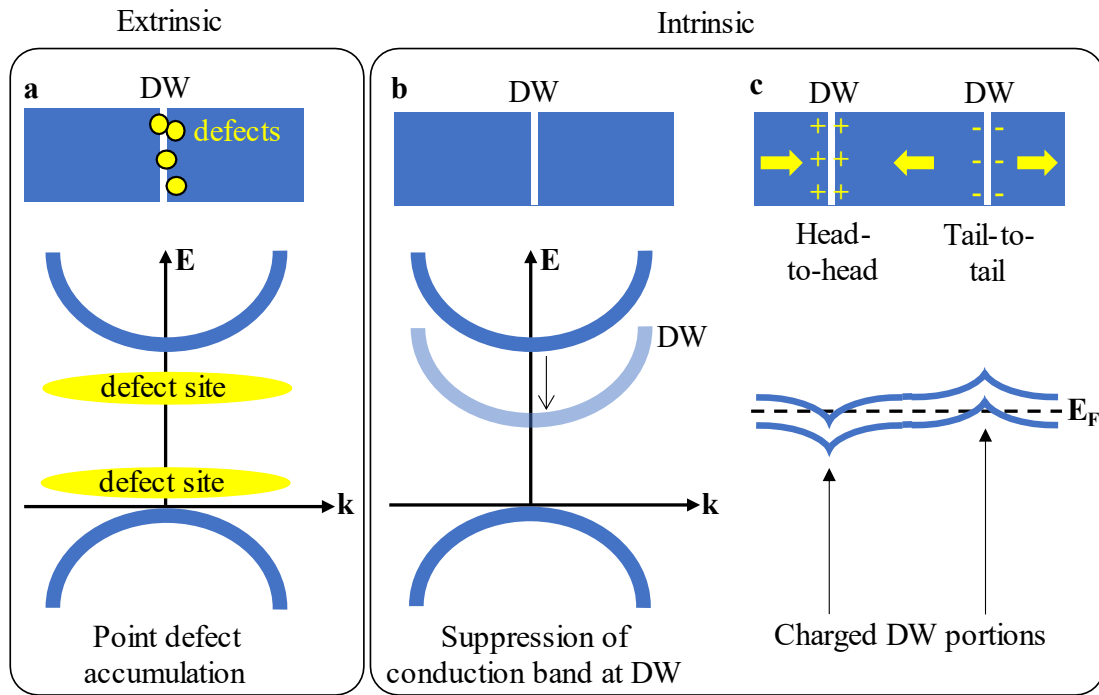


Fig. 4. **Typical mechanisms of domain wall (DW) conduction.** **a** | Intra-bandgap defect states between the conduction band (top semi-circle) and the valence band (bottom semi-circle). Usually, intra-bandgap states are caused by defects but it may also be intrinsic to the pristine domain wall¹⁰⁵. **b** | Intrinsic different electronic band structure of the domain wall leading to a reduction in the bandgap. **c** | Shift of the band structure induced by local electric fields associated with local head-to-head (giving positive bound charges "+") and tail-to-tail (giving negative bound charges "-") configurations of the polarisation (indicated by yellow arrows). E and k indicate the energy and the wave vector, respectively. E_F is the Fermi energy.

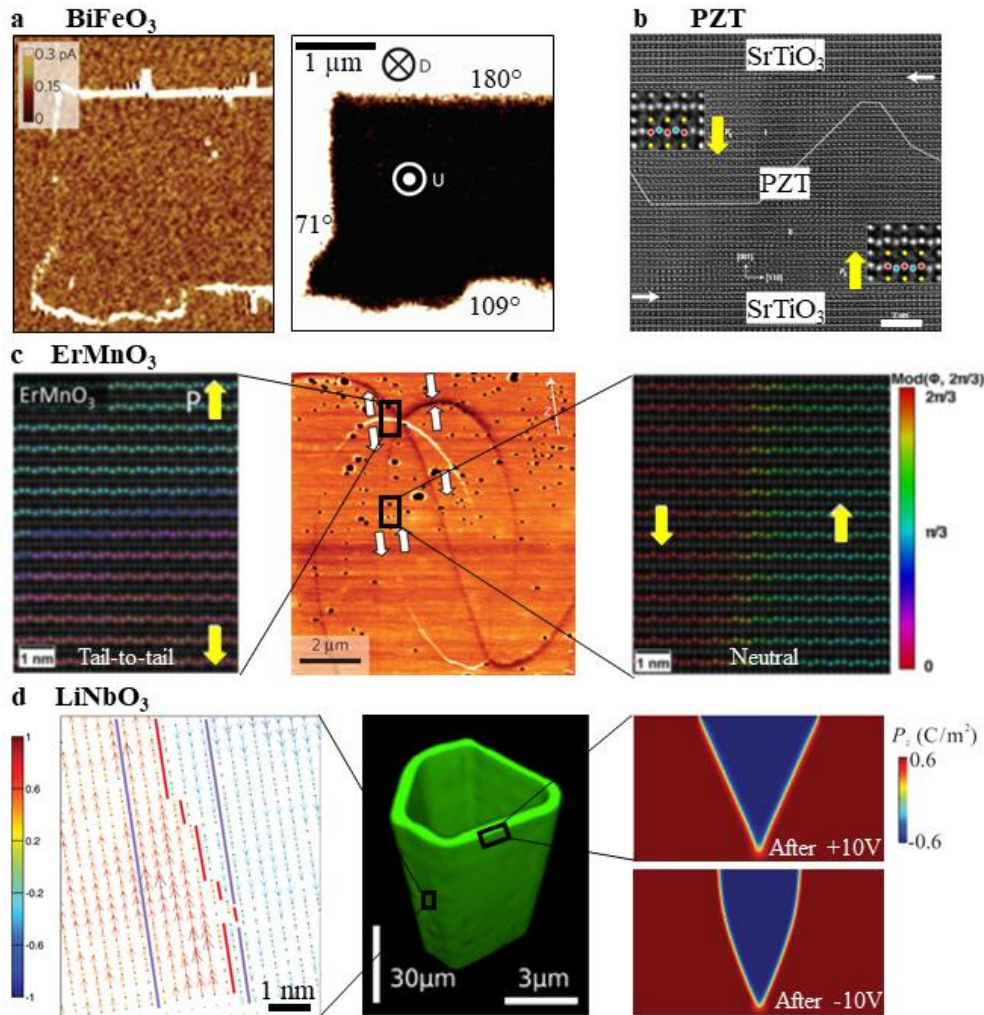


Fig. 5. Conduction at 180° domain walls. **a** | *c*-AFM (left) and out-of-plane piezoresponse force microscopy (right) images of a written domain pattern in a monodomain BiFeO₃ film showing the out-of-plane polarisation component of the domains to be either down, labelled as ‘D’ (white), or up, labelled as ‘U’ (black), with conduction at 109° and 180° domain walls, but not at 71°. Adapted with permission from REF.⁹⁴. **b** | TEM image of a SrTiO₃/PbZr_{0.2}Ti_{0.8}O₃/SrTiO₃ thin film heterostructure. The image is recorded under negative spherical-aberration imaging conditions with the incident electron beam parallel to the [-110] direction. The atom columns appear bright on a dark background. The horizontal arrows denote the horizontal interfaces between the PZT and the top and the bottom SrTiO₃ film layers. The dotted line traces the 180° domain wall. The yellow arrows show the directions of the polarisation in the 180° domains. The insets show magnifications of the dipoles formed by the displacements of ions in the unit cells (yellow: PbO, red: Zr/Ti, blue: O). The white scalebar corresponds to 2 nm. Adapted with permission from REF.⁷. **c** | *c*-AFM image of two neighbouring singularities in the *yz*-plane of ErMnO₃ with polarisation directions in-plane as indicated by the arrows. The inset on the left shows high-angle annular dark-field imaging-STEM (HAADF-STEM) images of negatively charged tail-to-tail domain walls and the inset on the right HAADF-STEM images of a neutral domain wall. The colour overlay of the phase describes the Er displacement. Adapted with permission from REF.^{113,115}. **d** | Three-dimensional profile of an inclined domain wall in LiNbO₃ by Cherenkov second-harmonic generation (centre). The inset on the left shows the analysis of 25 atomic rows from a STEM image: grey dots correspond to the fitted atom column positions, red and blue arrows indicate the displacement of the fitted atomic positions with respect to the base line and pointing in the direction of the displacement (the larger the displacement, the darker and longer the arrow). The inset on the right is a phase-field modelling of the domain wall profile after the application of +/- 10 V. Bending of the upper segments of wall close to the surface can be clearly seen. Adapted with permission from REF.^{6,129,131}.

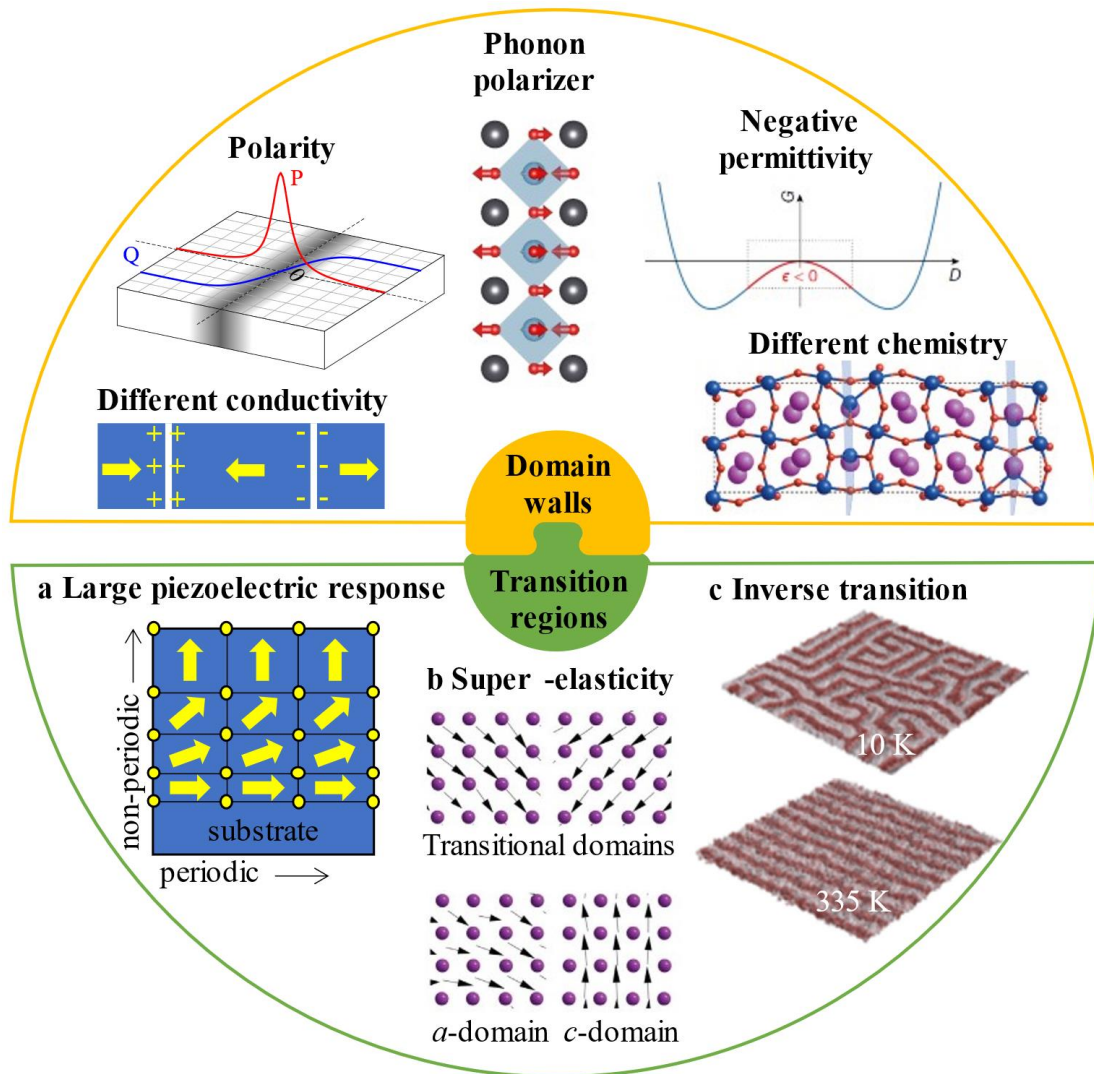


Fig. 6. Properties of domain walls and transition regions. Schematic view of the properties of domain walls (top semi-circle) and transition regions (a-c). The images for domain walls are adapted from literature: phonon polarizer¹⁶¹, negative permittivity¹⁵⁶, different chemistry¹⁶². **a** | Schematic of a transitional ferroelectric with a gradual change in symmetry. Adapted with permission from REF.²²⁹. **b** | Dipole configurations in four selected areas of a bent membrane by atomistic simulations, featuring a -domains, c -domains and transitional domains. Adapted with permission from REF.¹⁷⁷. **c** | Evolution of the labyrinthine pattern (10 K) to a stripe pattern (335 K) with increasing temperature. Adapted with permission from REF.¹⁷⁸.

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