

Abstract

Theoretical condensed matter research is plagued by a fundamental issue of complexity. The sheer amount of degrees of freedom in a material on any technologically relevant scale is overwhelming (e.g. $\sim 10^{23}$ electrons per cm^3), and makes it impossible to describe the quantum mechanical wavefunction exactly.

The Hamiltonian plays a central role in the description of crystals, the subject of this thesis. It can be decomposed into various parts, and their interactions. Depending on the physics under scrutiny it then often suffices to solve only one of those parts. This can be either because the energy scales and associated timescales that govern the constituents are very different, or because the interactions between them are small. One example, often put into practice, is the separation of electronic and phononic (lattice) degrees of freedom, leading to the well-known Born-Oppenheimer approximation, decoupling their respective motion. Another is the often neglected spin-orbit coupling, due to the tiny prefactor associated with its relativistic origin.

Solving these subproblems then allows for progress to be made in understanding the physics that govern them. However, there will inevitably be systems for which this interaction is not small and leads to fascinating new physics that manifestly depends on both subsystems combined. In this thesis we focus on these cases and how they arise in functional materials, with the occasional eye towards applications in technology.

The reason why these cross-order couplings can be interesting for technological applications, is that often one of the orders is more robust with respect to perturbations, and therefore more long-lived, but also harder to control efficiently. By exploiting the cross-order coupling in certain materials, one could potentially control the long lived order by applying perturbations to the more easily controllable order.

In giant Rashba effect systems, the coupling between spin and ferroelectric order leads to a linear spin-splitting of the band structure, whose sign depends on the orientation of the ferroelectric polarization. We show that, rather than the relativistic Rashba effect, a combination of electrostatics and atomic spin-orbit coupling lies at the origin of the large splitting.

The coupling between magnetism and ferroelectricity in multiferroic GdMn_2O_5 leads to a never before observed four-state hysteresis loop for the ferroelectric polarization, which depends on the magnitude, angle and history of the applied magnetic field. As we will show, this four-state hysteresis loop is accompanied by a full 360° rotation of spins in the material, which resembles the crankshaft of a car, converting the linear back-and-forth motion of the magnetic field into a rotational motion of the spins.

In a thin film of elemental Chromium, the ultrafast dynamics of a spin

density wave, coupled to a slower varying charge density wave, allows for a high degree of control of the latter through excitations of the former. This allows us to predict the sequence of optical pulses to be applied to the material in order to follow closely an enveloping signal function.

And finally, the coupling between ferroelectricity and strain in BaTiO_3 leads to a softening at purely ferroelectric domain walls, allowing for some mechanical control of the position of this wall.

We utilize both theoretical and computational tools to understand the nature of these interactions, how they lead to cross-order coupling in these materials, and how this then translates into the experimentally observed behavior.

Chapter 1

Mechanical softening in Ferroelectric domain walls

1.1 Introduction

So far, the focus of this work was largely on the behavior of, and interaction between various orders in uniform domains. A thorough understanding of these physics is of fundamental importance, since it provides the building blocks for more complex scenarios. Reality, however, presents to us almost exclusively these more complex scenarios. The reason for this is that while going through a phase transition with an associated spontaneous symmetry breaking, different parts of the material which are separated in space will separately (and usually randomly) choose one of the degenerate states with different values of the order parameters. One could argue that second order phase transitions could happen coherently throughout the entire system, but this assumes that the crystals are completely uniform and without defects, and that the temperature variation happens at an infinitely slow rate. [there is also the issue of dipolar fields, should I mention that?] In reality these assumptions are almost always invalid, inevitably leading to domains separated by domain walls (DWs) in which one or more order parameters interpolate between the values in the neighboring domains. The fundamental difference between DWs and the domains themselves has led to a great amount of research interest, both from the fundamental and technological point of view. The physics that govern the region of the DW is usually richer than that of the domains, since the domains force that region to not be at a local energetic minimum. This is often associated with a local inversion of the curvature of the energy potential, leading to a potential softening of the order parameters locally. DWs are also a manifestly two dimensional region embedded in a three dimensional medium, which can have a myriad of interesting consequences which can potentially be exploited technologically. One example is in ferroelectric information storage devices, where the small size is beneficial for information density, and ease of manipulation of their position allows for a great degree of flexibility. The materials that are usually used in this sense are those that can harbor charged conducting DWs, separating the manifestly nonconductive domains. By applying electric fields these can be created or destroyed at will, increasing or decreasing the conductivity of the

devicem. This conductivity can in turn be probed, effectively reading back what “data” was written previously[citation].

While this kind of technological promise has driven much of the research of DWs towards their electronic properties, the coupling between electric polarization and internal strain puts the spotlight on their mechanical properties. This field is much less developed and has many unanswered questions. [mention piezoelectrics?] It has been shown that ferroelectric-ferroelastic (ferroelastic walls separate domains with differing strain textures) DWs can be moved by applying stress [Schneider2001]. This is not surprising since the different strain textures couple differently to the mechanical perturbation, causing an imbalance and associated movement of the separating DW. More interesting is the fact that the polarization itself can also be influenced by purely mechanical means, for example by the flexoelectric effect, with energy density:

$$f_{fl} = \frac{1}{2} f_{jklm} (\varepsilon_{jk} \partial_m P_l - P_l \partial_m \varepsilon_{jk}), \quad (1.1)$$

where repeated indices are summed over, ε denotes the strain and P the ferroelectric polarization. An applied strain gradient will act as an internal electric field $E_j = \frac{1}{2} f_{jklm} \partial_k \varepsilon_{lm}$, coupling to the dipoles that constitute P_l . While this effect is generally small (i.e. the elements of the flexoelectric tensor f_{jklm} are small), it scales inversely with the size of the sample, meaning that it becomes increasingly important at the nanoscale of current state of the art electronic devices. Thus, using a tip to apply a strain gradient to the surface of a ferroelectric material allows one to mechanically write domain patterns and DWs at will [Lu2012].

These developments have increased the research into the mechanical properties of these materials in recent years. Here we focus on the mechanical properties of purely ferroelectric, 180° , DWs. As it turns out, these walls appear mechanically softer than the domains they separate. A similar softening has been previously observed and studied for ferroelastic DWs [Lee2003], but purely electric DWs have largely flown under the radar of mechanical studies. One reason is that the size of ferroelectric domain walls is on the order of a couple of unit cells, and was thus perceived too small to be detected with mechanical means, seen as tip contact areas are generally at least 100 unit cells. As it turns out, however, there is a strain texture associated with these DWs, which extends much further than the region in which the primary order parameter changes. This can qualitatively be understood by considering the electrostriction energy density

$$f_q = -\frac{1}{2} \varepsilon_{jk} q_{jklm} P_l P_m, \quad (1.2)$$

which causes domains to be stretched along the direction of P . No such stress exists inside the wall itself, since $P = 0$, leading to an indentation with an associated long range strain texture, because sharp changes in the strain are very unfavorable from an elastic point of view. This structural texture associated with the DW can be picked up mechanically by the applied tips and allows to make the observations that are the subject of this Chapter.

1.2 Experimental

Before going into the theoretical details that describe the physics of this problem, we describe the experiments performed by the group of Prof. Catalan that were the impetus of this research. In order to determine the generality of the softening of ferroelectric DWs, three single crystal systems were investigated experimentally: LiNbO_3 , BaTiO_3 and PbTiO_3 . All measurements are based on Contact Resonance Frequency Microscopy, a scanning probe microscopy technique that uses the resonance frequency of an atomic force microscopy tip in contact with the sample to measure its local stiffness [Rabe2000]. Through Hooke's law it is known that the resonance frequency is given by $\omega = \sqrt{\frac{k}{m}}$, i.e. a higher sample stiffness k leads to a higher frequency. This allows for a direct translation of the measured frequency into sample stiffness, where the only limit on spatial resolution is the scanning speed, or the duration of the experiment. The results are shown in Fig. ??, where the main focus are the right panels, displaying a clear contrast between soft areas (brown) close to the wall, and harder areas (purple) inside the domains. One observation that points to the strain texture of the walls as the main actor is the relatively large width of the soft areas around the domain walls. It is also important to realize that the tip is conductive and in contact with the surface at all times during the experiment, ensuring charge neutrality and thus excluding electrical excitation. Another feature of the measurements, most clearly present in Fig. ??(f) is the difference in contrast between the up and down domains. This can be attributed to the flexoelectric coupling between the tip induced strain gradient and the polarization. Even though the applied forces were too small to reorient the polarization, the asymmetry in mechanical response can still be observed, and has been proposed as a mechanism for voltage-free mechanical reading of polarization [Cordero-Edwards2017, Cordero-Edwards2019, Abdollahi2015]. A similar explanation can not be used for the DWs, though, since they are markedly softer than either domain, resulting in a reduction of the stiffness of $\approx 19\%$.

[should I go more into detail of the models used by the experimentalists for the Young's modulus calculation?]

1.3 Theory

Since the mechanical DW softening seems a general property of 180° ferroelectric DWs, we focus on BaTiO_3 (BTO) in the following. Due to the relatively large size of the strain texture associated with domain walls, and the tip that is pressed into the material, we adopt the continuum Ginzburg-Landau-Devonshire model as described in [Marton2010]. The free energy density in terms of primary order parameter for the ferroelectric polarization, P , and associated strain ε

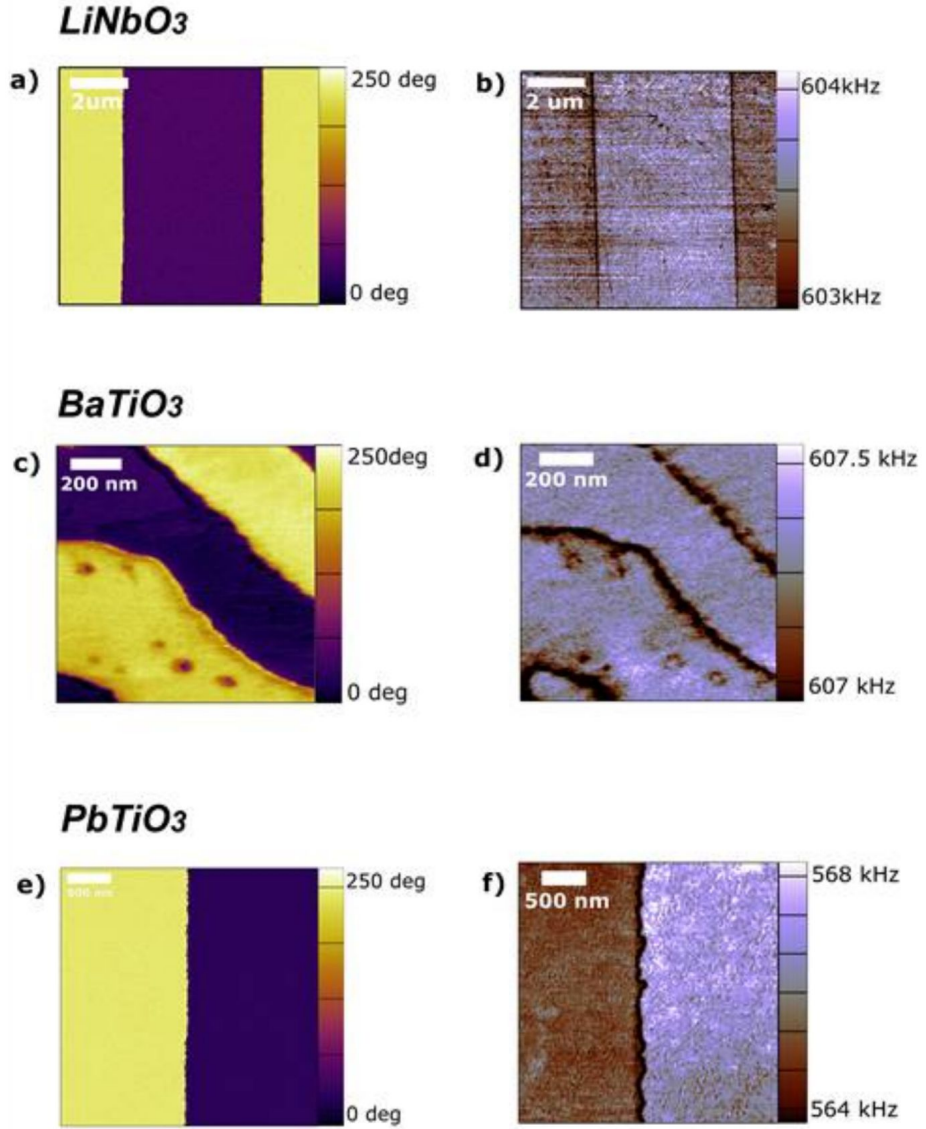


Figure 1.1: **Ferroelectric polarization and stiffness** (a-b) Periodically poled LiNbO₃ single crystal, (c-d) BaTiO₃ spontaneously polarized and (e-f) PbTiO₃ thin film. The contrast in figures (a,c,d) demonstrate the two opposite out-of-plane orientations of the ferroelectric polarization, and the differing stiffness in (b,d,f). The main focus are the DWs which appear most dark and are thus markedly softer than the domains.

α		G		q	
α_1	$3.34 \cdot 10^5 (T - 381)$	G_{11}	$51 \cdot 10^{-11}$	q_{11}	$14.2 \cdot 10^9$
α_{11}	$4.68 \cdot 10^6 (T - 393) - 2.02 \cdot 10^8$	G_{12}	$-2 \cdot 10^{-11}$	q_{12}	$-0.74 \cdot 10^9$
α_{12}	$3.23 \cdot 10^8$	G_{44}	$2 \cdot 10^{-11}$	q_{44}	$1.57 \cdot 10^9$
		C		f	
α_{111}	$-5.52 \cdot 10^7 (T - 393) + 2.76 \cdot 10^6$	C_{11}	$27.5 \cdot 10^{10}$	f_{11}	0.85
α_{112}	$4.47 \cdot 10^9$	C_{12}	$17.9 \cdot 10^{10}$	f_{12}	-0.5
α_{123}	$4.91 \cdot 10^9$	C_{44}	$5.43 \cdot 10^{10}$	f_{44}	0.072

Table 1.1: **Ginzburg-Landau-Devonshire model parameters** The parameters are reported in Voigt notation, e.g. $C_{11} = C_{1111}$, $C_{12} = C_{1122}$ and $C_{44} = C_{1212}$. SI units are used, and $T = 300\text{K}$ in the simulations.

can be written as:

$$f = f_L + f_G + f_c + f_q + f_{fl}, \quad (1.3)$$

$$f_L = \alpha_{ij} P_i P_j + \alpha_{ijkl} P_i P_j P_k P_l + \alpha_{ijklmn} P_i P_j P_k P_l P_m P_n, \quad (1.4)$$

$$f_G = \frac{1}{2} G_{ijkl} \partial_i P_j \partial_k P_l, \quad (1.5)$$

$$f_c = \frac{1}{2} C_{ijkl} \varepsilon_{ij} \varepsilon_{kl}, \quad (1.6)$$

$$f_q = -q_{ijkl} \varepsilon_{ij} P_k P_l, \quad (1.7)$$

$$f_{fl} = \frac{1}{2} f_{ijkl} (\varepsilon_{ij} \partial_k P_l - P_i \partial_j \varepsilon_{kl}), \quad (1.8)$$

where the indices run through x, y, z , and a summation is performed over the repeated ones. The first term is the Landau free energy for a uniform ferroelectric polarization, where terms up to sixth order have to be included to bound P , since in BTO both α_{ij} and α_{ijkl} are negative below the transition temperature. The second term denotes the Ginzburg part, i.e. the energy penalty for spatial variations of the polarization. This term influences the width of DWs, where a larger G leads to wider walls and vice versa. The elastic energy density is described by f_c with stiffness tensor C_{ijkl} , which has the form of the standard Hooke's law. f_q signifies the electrostriction, the main term that couples the polarization to the strain, and causes the domains to be stretched along the direction of the polarization [add some panels like in the discussion of the powerpoint]. Lastly, to be complete, we can include the flexoelectric contribution, f_{fl} , but ultimately we neglected it in the simulations, since it leads to effects which are small and did not markedly influence the results. The parameters for the model for BaTiO₃ are given in Tab. ???. From symmetry, only a limited set of parameters is unique, for example $C_{1111} = C_{2222} = C_{3333}$, and $C_{1212} = C_{2121} = C_{2323} = C_{3232} = C_{1313} = C_{3131}$, etc.

To find the static equilibrium conditions in the most general sense, the integrated free energy $F[P] = \int d^3x f(P(x))$ needs to be minimized, for which the variational method can be used. This leads to the well known Euler-Lagrange

equations [Cao1991, Marton2010]:

$$\frac{\partial}{\partial x_j} \left(\frac{\partial f}{\partial \partial_j P_i} \right) - \frac{\partial f}{\partial P_i} = 0, \quad (1.9)$$

$$\frac{\partial}{\partial x_j} \frac{\partial f}{\partial \varepsilon_{ij}} = 0. \quad (1.10)$$

In the homogeneous case of a single domain with P_z (which we write P in the following) and ε_{zz} (written as ε) and no external forces, Eq. ?? is reduced to

$$\frac{\partial f}{\partial \varepsilon} = 0 \Leftrightarrow \varepsilon = \frac{q_{11} P^2}{C_{11}}, \quad (1.11)$$

$$\frac{\partial f}{\partial P^2} = 0 = \alpha_1 + (2\alpha_{11} - \frac{q_{11}^2}{C_{11}}) P^2 + 3\alpha_{111} P^4 \quad (1.12)$$

$$\Leftrightarrow \quad (1.13)$$

$$P^2 = \frac{-(2\alpha_{11} - \frac{q_{11}^2}{C_{11}}) + \sqrt{(2\alpha_{11} - \frac{q_{11}^2}{C_{11}})^2 - 12\alpha_{111}\alpha_1}}{6\alpha_{111}}. \quad (1.14)$$

If the parameters of Tab. ?? are filled in, we find $P = \pm 0.311$ and $\varepsilon = 5 \cdot 10^{-3}$. This is higher than $P = 0.265$ [Marton2010] without any electrostriction included, because the system can lower its energy through $-q_{11} P_z^2 \varepsilon_{zz}$ by increasing P_z and having tensile strain $\varepsilon_{zz} > 0$. Similarly, when all terms are included in the full numerical simulation, we find for the domains that $P_z = \pm 0.363$ and $\varepsilon_{zz} = 1.4 \cdot 10^{-2}$.

The effective stiffness \tilde{C}_{ijkl} can be found in general by taking the double derivative of the free energy with respect to ε :

$$\tilde{C}_{ijkl} = \frac{\partial^2 f}{\partial \varepsilon_{ij} \partial \varepsilon_{kl}}, \quad (1.15)$$

or, making the same assumptions as above (only P_z and ε_{zz}), we arrive at a stiffness of

$$\tilde{C} = C - \frac{q_{11}^2}{2 \left(\alpha_{11}^2 - 3\alpha_{111} \left(\alpha_1 - \frac{q^2 P^2}{C_{11}} \right) \right)}. \quad (1.16)$$

Again filling in all parameters for the homogeneous case leads to $\tilde{C}_d = 0.778 C$ in the domain. If there would be no penalty for shear strain, we find that, at the all, $P = 0$ and $\varepsilon_{zz} = 0.0$, leading to a stiffness of $\tilde{C}_w = 0.642 C = 0.82 \tilde{C}_d$. While this does look like a triumph for the theory (since we're trying to find a 19% softer wall), in reality the shear elastic constants C_{44} will lead to a nonzero strain ε_{zz} at the surface of the wall of around 40% that in the domain, leading to $\tilde{C}_w = 0.79 C = 1.01 \tilde{C}_w$. This is confirmed by simulations with a very narrow tip.

This is in contradiction with the observations of the experiments, so a different explanation has to be sought.

The first possible source for the mechanical softening originates from the electrostriction term, and the strain texture it results in. As mentioned before, electrostriction stretches the domains in the direction of the polarization, i.e. $\varepsilon_{zz} \neq 0$ in the domains. In the domain wall, however, P_z^2 is diminished and

even zero at the center, causing ε_{zz} to be diminished, but never reduced to zero due to compatibility relations and the elastic coupling to neighboring unit cells. Nonetheless, this results in an indentation that forms at the location of the domain wall, as shown pictorially in Fig. ??(b), and more realistically in (c-d). As it turns out, the strain texture of this indentation stretches out relatively far [actual derivation and formula for this?] from the domain wall. This long-rangedness of strain is a general phenomenon, and depends on the morphology of the strain defect [more indepth on this?]. When the tip is then applied in an area where this strain texture is present, the wall will try to bend rigidly (i.e. the strain texture shifts but doesn't change shape) towards the tip in order to gain on the displacement. This leads to a relatively big displacement when the tip is applied, making the material appear soft.

Even though the interaction between the pinning potential, Peierls-Nabarro barriers, and electrostatics, with the force applied by the tip is hard to analytically describe, we can make statements about two extremes of the behavior: i) If the force of the tip is large enough, the wall slide towards it, maximizing the possible energy gain from the interaction with the tip. ii) A bending of the wall, where it remains inside the original Peierls-Nabarro potential, but deviates from the equilibrium position. [The situation that happens in the real material is more like a mix between the two, the top part of the wall bends almost completely towards teh tip, but it's not moved as a whole because the bottom/bending electrostatics pins it. Can we say that these things are causing the potential for the entire wall to behave like the one we describe below?]

The first case can be ignored because this would mean that in the experiments, the wall would be dragged along the tip since the tip moves at a relatively slow rate, which would lose any contrast between wall and domain during the full measurement. We therefore try to formulate a simple free energy expansion for the second situation, where we assume that the wall at x_{DW} is pinned by a parabolic potential, and perturbed by a tip applying a force F_z at x_{tip} ,

$$E = E_0 - F_z u_z(x_{tip} - x_{DW}) + \frac{m\omega^2 x_{DW}^2}{2}. \quad (1.17)$$

We can expand this equation under the assumption of a small x_{DW} , i.e. that the wall doesn't move far from the $x_{DW} = 0$ equilibrium situation. Together with minimizing the energy we obtain $x_{DW} = -Fu'(x_{tip})/m\omega^2$, with a compliance correction $\Delta c = u'(x_{tip})^2/(m\omega^2)^2$. Thus, we can conclude that to maximize the softening, the tip should be applied where $u'(x_{tip})$ is large, i.e. within the strain variation caused by the above discussed electrostrictive coupling. This part of the effect is pictorially depicted in panel (b) of Fig. ??.

In reality, depending on where the tip is pressed, a combination of the two above mentioned extremes will happen. When the center of the tip is close enough to the wall, the upper part of it will completely bend to shift the indentation underneath the tip. The lower portion of the wall will remain at the center of its original Peierls Nabarro potential, since it experiences less effect of the tip. It is therefore impossible to solve analytically the realistic situation, and we proceed with simulations to make semi-quantitative statements on the behavior.

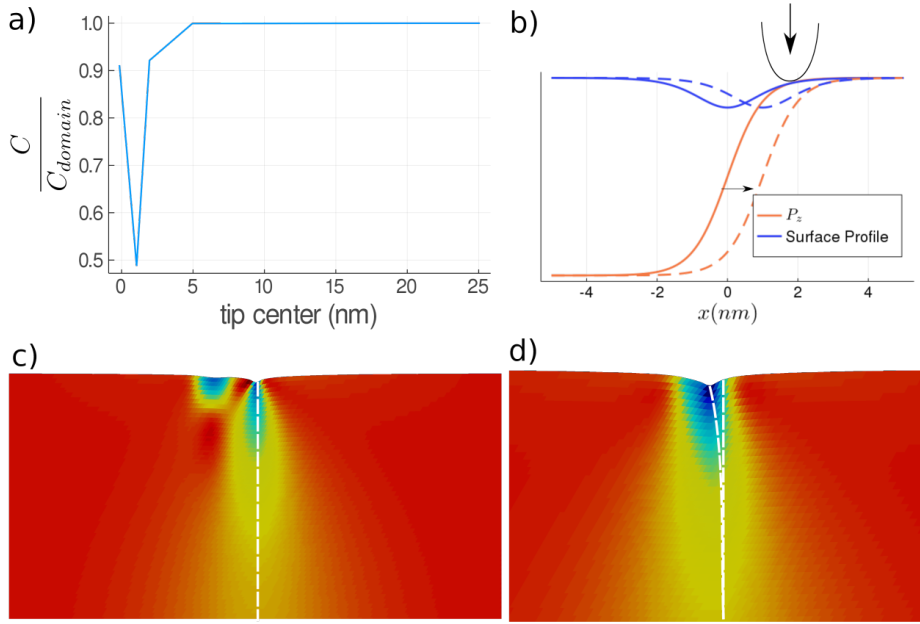


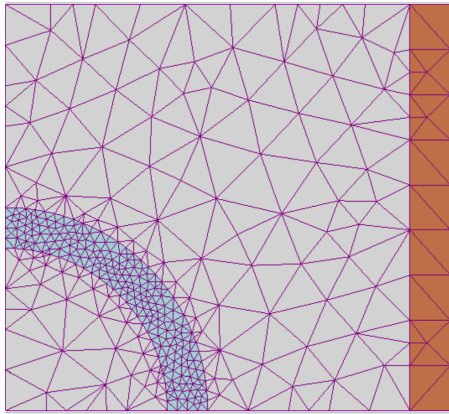
Figure 1.2

1.4 Methods

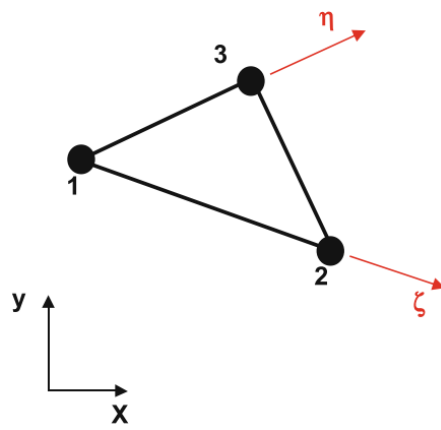
The Finite Elements method (FEM) provides a natural framework to solve continuum problems, sometimes also referred to as phase-field problems, that involve gradients such as strain (gradient of the displacement u) and the gradient of polarization [Biner].

Similar to the well-known finite-difference methods, FEM is a local method, dividing the problem into separate domains (the elements) for which the contribution to the total energy can be calculated locally. A fundamental building block of this method is the isoparametric representation. In this representation, the geometry is divided into a set of elements that obey certain connectivity requirements, e.g. in Fig. ??(a) a 2D geometry is divided into triangular elements. The values of the the displacement field $\mathbf{u}(\mathbf{x})$ and ferroelectric polarization $\mathbf{P}(\mathbf{x})$ are defined on the nodal points that link the different elements shown by the black dots of Fig. ??(b). Inside each element a local coordinate system with axes η and ζ can then be defined as shown by the red arrows in Fig. ??(b). Together with the shape functions that are defined on each local element [Should I give some more info on what they are], this allows for the values of the fields to be interpolated at all points inside the elements from their values at the nodes. Since the element represents a given volume of the total geometry, the contribution of each element to the total integral can thus be evaluated. A big bonus of this framework is that gradients of the fields can be evaluated purely locally, inside each element, through the use of this interpolation method. It also allows for a great flexibility through the density of the chosen grid and the order of interpolating functions used, although we do not exploit these.

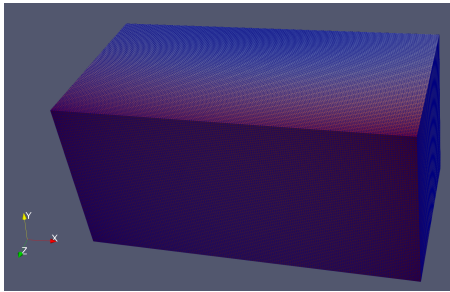
In our case, we use a rectangular geometry uniformly spanned by tetrahedron elements, as shown in Fig. ??(c-d). The uniform mesh is largely motivated by



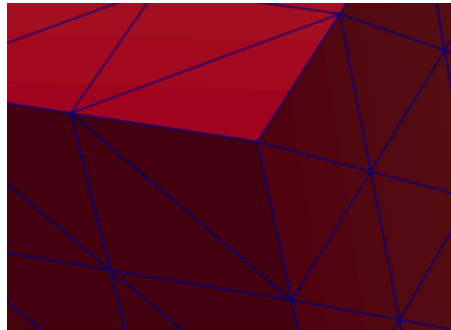
a) 2D grid with triangular elements, [Biner]



b) Triangular element, [wikipedia]



c) Geometry used in simulations



d) Zoom on one element

Figure 1.3: Example of the morphology of a 2D finite element grid.

implementation complexity, since we use a mostly in-house developed code based on the FEM building blocks supplied by the JuAFEM.jl package. It is obvious that the computational cost could be lowered by using a finer meshing close to the domain wall, where the fields vary the most, and a more course one in the domains. One has to, however, consider that when the tip is applied a finer mesh is also required around it, leading to different meshes for different locations of applied tip, greatly increasing the post-processing complexity.

The last bulding block for the simulation is a way to optimize the fields in order to reach the equilibrium condition, both with and without applied tip. To do this we use a Conjugate Gradient method to minimize the total energy in terms of all the degrees of freedom (DOFs), i.e. the values of the fields at each of the nodes on the mesh. This requires us to define the partial derivatives of the total energy in terms of all the DOFs, for which we use a forward automatic differentiation scheme as implemented in ForwardDiff.jl, again for its remarkable simplicity and speed (as an idea, on a 24 core E5-2680 v3 server, the ≈ 3.5 million partial derivatives are computed in around 2.1 seconds).

Using the parameters from previous ab-initio results [Marton2010], we first optimized the equilibrium situation for both a uniform domain and a single domain wall. A bell shaped force field, modeled by function ae^{-x^2/d^2} with $d \approx 20\text{nm}$ representing the diameter of the contact area and $a = 10^{-7} J/m^3$, was then applied at various positions throughout both geometries. The uniform domain simulation is used as a measure of the finite-size effects that are inherently present in our simulations due to the long rangedness of strain and non-infinite geometry. This causes the indentation at the center of the geometry for a single domain to be less than the one when the tip is applied off-center. Since this is unphysical, we compensate for this by adding the differences to the results in the geometry with a wall present [better explanation?].

1.5 Results

We chose to use open boundary conditions[maybe try with periodic too].

When using finite element methods, it's important to verify to what extent the results are influenced by finite size effects and other numerical considerations. Since the effect we are investigating depends crucially on the movement and position of the domain wall, the issue of the numerical Peierls-Nabarro barrier [Marton2018] is quite important since the degree of pinning that the wall experiences depends on the elementsize. We found that when the elementsize is 0.5 nm, the wall is too strongly pinned and never moves towards the tip, even if it's applied very close to it. Using elements of 0.3 nm alleviates this issue and leads to a more realistic behavior.

Furthermore, due to the computational cost, it is desirable to use as small geometries as possible.

In order to get an idea of how robust this behavior is w.r.t. the numerical parameters we show the relative

Multiple geometries and element sizes were tested in order to verify that the results were not too much influenced by these numerical parameters which. The softening is estimated as the ratio of the surface deformation between the domain and the wall.

As seen in Fig. ??(a), and predicted by the above statements, the wall is softest not in the center, but rather at the point where u' or ε_{xz} is highest.