

Manipulating Topological Domain Boundaries in the Single-layer Quantum Spin Hall Insulator $1T'-WSe_2$

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Abstract

We report the creation and manipulation of structural phase boundaries in the single-layer quantum spin Hall insulator $1T'-WSe_2$ by means of scanning tunneling microscope tip pulses. We observe the formation of one-dimensional interfaces between topologically non-trivial $1T'$ domains having different rotational orientations, as well as induced interfaces between topologically non-trivial $1T'$ and topologically trivial $1H$ phases. Scanning tunneling spectroscopy measurements show that $1T'/1T'$ interface states are localized at domain

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3 boundaries, consistent with theoretically predicted unprotected interface modes that form
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5 dispersive bands in and around the energy gap of this quantum spin Hall insulator. We observe a
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7 qualitative difference in the experimental spectral lineshape between topologically “unprotected”
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9 states at 1T’/1T’ domain boundaries and protected states at 1T’/1H and 1T’/vacuum boundaries
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11 in single-layer WSe₂.
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17 Keywords: Scanning tunneling microscopy, transition metal dichalcogenides, quantum spin hall
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19 insulators, domain boundary, ferroelasticity.
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24 Recent experimental studies have reported the observation of the quantum spin Hall (QSH)
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26 effect in single layers of the transition metal dichalcogenides (TMDs) WTe₂ and WSe₂ in the 1T’
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28 structural phase.¹⁻⁵ Evidence of the QSH state include inverted bandgaps,¹ topologically
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30 protected edge states,^{1,2,5} as well as quantized edge conduction of e^2/h per edge.^{3,4} QSH edge
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32 states have been observed to reside at 1T’/1H and 1T’/vacuum boundaries, both of which are
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34 interfaces between non-trivial (1T’–TMD) and trivial (1H–TMD phase or vacuum) media.^{1,2,5}
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36 Such interfaces are expected to host topologically protected edge states.⁶⁻¹¹ A less well-studied
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38 type of boundary in quantum spin Hall insulator (QSHI) materials is the interface between
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40 different non-trivial domains where the Z_2 topological invariant of the bulk does *not* change
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42 across the interface. We refer to a domain boundary as “topological” when there is a change in
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44 the topological invariant across the interface, and “trivial” when the invariant is the same on
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46 either side of the domain wall. A recent theoretical study of charge transport in quantum Hall
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48 insulators with trivial interfaces predicted that conduction through otherwise dissipationless
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50 quantum Hall edge states can be controllably deflected into trivial interface states, thus enabling
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52 gate-tunable charge and spin transport.¹² Single-layer TMD materials provide a new strategy for
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3 constructing such coexisting topological and trivial interfaces by switching the layer structural
4 phase via some local stimuli.¹³⁻¹⁷ For example, the 1T' phase has recently been predicted to be
5 ferroelastic in single layers, suggesting that it can be switched between different dimerization
6 orientations by applied stress. This provides a mechanism to induce topologically trivial
7 interfaces between QSHI domains with different crystallographic orientations. A similar strategy
8 could allow generation of topologically non-trivial interfaces by inducing local phase switching
9 between structures that have different Z_2 indices.^{18,19}

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11 Here we report the local phase manipulation of single-layer 1T'-WSe₂ for the purpose of
12 creating two kinds of one-dimensional interfaces: (1) trivial interfaces between two 1T' domains
13 and (2) topological interfaces between 1T' and 1H domains. By using scanning tunneling
14 microscope (STM) tip pulses we are able to locally switch from the 1T' phase to the 1H phase of
15 WSe₂, as well as between different orientations of the 1T' phase. 1T'/1T' domain-boundary
16 formation is observed to be reversible, supporting the conjecture that single-layer 1T'-WSe₂ is a
17 ferroelastic material.¹⁸ Our STM measurements show that 1T'/1T' domain boundaries are well-
18 ordered interfaces that exhibit several different structures. By combining scanning tunneling
19 spectroscopy (STS) measurements and first-principles calculations we have determined that
20 1T'/1T' domain boundaries exhibit topologically unprotected one-dimensional (1D) modes that
21 are dispersive near the Fermi level and that exhibit energy dependent decay lengths. These
22 modes reside both inside and outside of the 1T'-bulk bandgap and, unlike 1T'/1H interface
23 modes, do not directly connect bulk valence and conduction bands.

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25 Mixed-phase single layers of WSe₂ were grown using molecular beam epitaxy (MBE) on
26 bilayer graphene (BLG) supported by SiC. These samples exhibit islands that are single domains
27 of either 1T' or 1H phase, as well as mixed-phase islands with coexisting 1T' and 1H domains.²

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3 Voltage pulses applied between the STM tip and monolayer $1T'$ -WSe₂ islands were used to
4 manipulate the WSe₂ structural phase. Fig. 1 shows STM topographic images of a $1T'$ -WSe₂
5 island before and after application of STM tip pulses. The “before” image (Fig. 1a) shows a
6 single-domain region of the $1T'$ phase with a uniform orientation of atomic rows running from
7 top to bottom (each row contains a zigzag chain of W atoms^{1,2,9,11}). Fig. 1b shows the same
8 region after a voltage pulse of 10 V was applied for 100 ms between the tip and surface at a
9 constant tip-surface separation of ~ 6 Å. After application of the pulse the island exhibits multiple
10 domains having different orientations that are connected by ordered 1D domain boundaries (tip
11 pulses can also cause the formation of adsorbate clusters near domain boundaries, as shown in
12 Fig. S7).

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14 Because the $1T'$ phase can be formed in three equivalent orientations (via a Peierls
15 distortion of its C_3 -symmetric $1T$ parent phase) several possible $1T'/1T'$ domain boundaries are
16 expected.¹⁸ The most common $1T'/1T'$ interface observed in our samples is the 120° domain
17 boundary which occurs for 85% of all observed boundaries and which connects neighboring
18 domains rotated with respect to each other by 120° (Fig. 2a). Other observed domain boundaries
19 are the 60° domain boundary (observed 13% of the time) and the 0° domain boundary (observed
20 2% of the time), as shown in Figs. 2b, c (structural models are shown in Figs. 2d-f). These well-
21 ordered interfaces are straight-line defects that extend up to 20 nm in length in our samples. The
22 formation of $1T'/1T'$ domain boundaries is reversible through the application of a high current
23 raster scan by the STM (300mV, 1nA). Such scans remove the adsorbate clusters that form
24 during the generation of $1T'/1T'$ domains, likely changing local strain distributions. This
25 provides additional evidence of the ferroelastic nature of $1T'$ -WSe₂ (further details are discussed
26 in section 6 of the SI).

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3 The local conversion of single-layer WSe₂ from the 1T' phase to the 1H phase can be
4 induced using the same voltage pulse method as described above. Fig. 1c shows a different
5 single-phase 1T' island where the dimer chains run from top to bottom before applying a tip
6 pulse, while Fig. 1d shows the same region after applying a voltage pulse of 10 V for 100 ms.
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8 The tip pulse causes an extended region of the island to convert into a new phase that exhibits
9 reduced apparent height. We identified this region as the 1H phase of single-layer WSe₂ (see Fig.
10 S6 for details). Such 1T' to 1H phase conversion was only observed in “confined” regions as
11 seen here (i.e. tip-induced 1H domains were always surrounded by other material).
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22 While 1T'/1T' domain boundaries could be created by applying tip pulses with $V_{\text{pulse}} \geq 6$
23 V, stronger tip pulses ($V_{\text{pulse}} \geq 10$ V) were required to locally induce the 1T' to 1H phase
24 transition. These observations agree with predictions that the transition barrier between different
25 orientations of the 1T' phase should be lower than the barrier for a 1T' → 1H transition.¹⁸ Tip
26 pulses with $V_{\text{pulse}} > 6$ V often caused damage to 1T' islands, either by creating holes or by
27 breaking apart the island. However, once 1T'/1T' and 1T'/1H domain structures are successfully
28 induced then they remain stable under normal scan conditions.
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38 In order to explore the electronic structure of topologically trivial interfaces in a QSHI we
39 performed STS at the sites of the 120°, 60°, and 0° boundary structures shown in Figs. 2a-c. The
40 dI/dV spectra obtained in the 1T' bulk (green curves in Figs. 2g-i) reflect the 1T'-WSe₂ bulk
41 bandgap which has an average full width at half maximum (FWHM) of 85 ± 21 mV centered at -
42 130 ± 5 mV (determination of the bulk bandgap was performed as described in Supplementary
43 Note 3 of ref. 2). The spectral weight observed inside the bulk bandgap is explained by lifetime
44 broadening and the -130 mV offset is due to n-doping induced by the bilayer graphene/SiC
45 substrate, consistent with previous studies.^{2,20,21} The narrow dip at $V=0$ seen in Figs. 2g-i likely
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3 arises from an interplay between disorder and long-range electron-electron interactions as has
4 been suggested previously.^{2,5,20,21,27,28} This feature is more pronounced at 1D domain boundaries
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6 which is consistent with predictions regarding disorder-induced behavior in low dimensions.²⁸
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10 The spectra for the 120° and 60° domain boundaries (Figs. 2g, h) are similar in that they
11 both have a minimum at $V = 0$ and exhibit broad, sloping features in the filled state regime over
12 the range $-300 \text{ mV} < V_s < 0$. Neither of these domain boundaries show any significant signatures
13 of the bulk bandgap. The 0° domain boundary (Fig. 2i) also has a minimum at $V = 0$, but it
14 shows a pronounced dip right in the bulk bandgap energy range. These experimental features are
15 qualitatively different from dI/dV spectra observed at topologically-protected 1T'/vacuum and
16 1T'/1H boundaries where a clearly defined edge-state peak is seen at the bulk bandgap energy² (a
17 reference dI/dV spectrum taken at the topological 1T'/vacuum edge is shown in Fig. S8).
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28 Because the 120° domain boundaries are the dominant defect feature, we performed a more
29 in-depth study of their spatially-dependent electronic structure. Fig. 3 shows dI/dV maps over the
30 energy range $-400 \text{ mV} < V_s < 150 \text{ mV}$ for a 120° 1T'/1T' domain boundary that intersects a
31 1T'/1H domain boundary. The first panel (Fig. 3a) shows an STM topograph of the region and
32 includes the 120° 1T'/1T' domain boundary (dashed oval) as well as the 1T'/1H boundary
33 (marked by a vertical dashed line with the 1T' phase to the right). This area allows us to
34 simultaneously compare the electronic structure of “topological” and “trivial” domain
35 boundaries.
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47 Fig. 3b shows a dI/dV map measured at -400 mV , which corresponds to an energy below
48 the lower edge of the bulk 1T'-WSe₂ bandgap shown in Fig. 2g (i.e., the bulk valence states).
49 Bright intensity corresponding to high LDOS is observed at the site of the 120° domain boundary
50 while the LDOS near the 1T'/1H interface remains low. Fig. 3c shows a dI/dV map at -120 mV ,
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3 which lies inside the $1T'$ bulk bandgap. At this energy high LDOS intensity is observed at both
4 the 120° domain boundary and the $1T'/1H$ interface region (intensity near the $1T'/1H$ boundary
5 originates from the topological edge state.² Fig. 3d shows a dI/dV map measured at -60 mV,
6 which is near the upper edge of the bulk bandgap. Here high-intensity LDOS is observed near
7 the $1T'/1H$ interface (from the topological edge state), while the LDOS at the 120° domain
8 boundary shows low intensity. Fig. 3e shows a dI/dV map measured at $+150$ mV, which
9 corresponds to an energy well into the bulk conduction band. At this energy neither the
10 topological $1T'/1H$ interface state nor the trivial 120° domain boundary show high intensity
11 features. The high intensity LDOS localized at the $1T'/1T'$ boundary in the dI/dV maps indicates
12 the existence of defect states in the energy range -400 mV $< V_s < -60$ mV. The broad
13 spectroscopic feature measured in the dI/dV point spectrum over this range for 120° domain
14 boundaries (Fig. 2g) can thus be attributed to confined dispersive defect modes. The 120°
15 domain boundary mode is seen to have a more strongly energy-dependent decay length than the
16 $1T'/1H$ interface state and to have more intensity at lower energies.

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19 In order to clarify the origin of these electronic features, we performed first-principles DFT
20 simulations. The atomic structure of the interfaces was first relaxed using periodic boundary
21 conditions in a ribbon geometry with a plane-wave basis set²² (the relaxed structures are
22 presented in Figs. 2d-f). We then used the non-equilibrium Green's function method (NEGF) to
23 model the line defects with semi-infinite boundary conditions.^{23,24} The resulting spin-dependent
24 electronic band structures for the three different interfaces are presented in Figs. 4a-c. The
25 number of bands, the dispersion, and the spin character of the defect modes changes dramatically
26 for the different boundary types. Defect modes belonging to the 60° domain boundary (which
27 has the least amount of symmetry) span the entire bandgap energy region while the localized
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3 states of the 120° and 0° domain boundaries do not completely close the bulk energy gap (the
4 bulk bandgap of the 60° domain boundary model is somewhat affected by the strain that arises
5 from matching bulk lattice constants with the defect periodicity). We see that while the defect
6 modes mostly close the overall energy gap for the 60° and 120° domain boundaries, the gap
7 remains bulk-like for the 0° defect, consistent with our STM spectroscopy observations (Figs.
8 2g-i). The simulated 120° domain boundary states are observed to be spin-polarized out of the
9 plane while for 60° domain boundaries the direction of spin polarization rotates for different
10 states within the same band (colors in Figs. 4a, b). 0° domain boundaries possess inversion
11 symmetry and so defect states associated with this boundary show no spin polarization (Fig. 4c).

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24 Figs. 4d-f compare the calculated LDOS of the 120° , 60° and 0° domain boundaries (blue
25 curves) with the LDOS of the 1T' bulk (grey curves) as a function of energy (the Fermi level has
26 been shifted to match the experimentally observed n-doping). Overall we find reasonable
27 agreement between the simulated LDOS in Fig. 4 and the corresponding STM spectroscopy
28 measurements of Fig. 2. For example, while the experimental bulk bandgap feature vanishes for
29 the 60° defect (Fig. 2h), an energy gap remains for the 0° defect (Fig. 2i), similar to the theory
30 plots of Figs. 4e, f. In the 120° case the predicted small 10 meV energy gap is likely smeared out
31 by level broadening effects that are observed experimentally but not accounted for in
32 conventional DFT simulations.²⁵ When we add Gaussian broadening to our calculation then the
33 120° gap feature is smeared out (Fig. 4d) similar to what is seen experimentally (Fig. 2g). A
34 significant discrepancy between the theory and the data is the pronounced LDOS peak seen near
35 $E = 0$ for all three domain boundary types. By contrast, all three domain boundaries show a
36 broad dip at $V = 0$ in the STM spectroscopy rather than the predicted peak. This is explained by
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3 the coexistence of disorder and electron-electron interactions in these materials which opens a
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5 pseudogap at $V = 0$ eV but which is not accounted for in DFT simulations.
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8 In conclusion, we have successfully manipulated the local electronic and structural
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10 properties of single-layer $1T'$ -WSe₂, thereby inducing a local phase transition from the $1T'$ to
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12 the $1H$ phase, as well as creating $1T'/1T'$ domain boundaries. The induced $1T'/1T'$ domain
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14 boundaries exhibit different rotational configurations, with a 120° domain boundary being the
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16 most common structure. Our combined STS measurements and first-principles calculations show
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18 that these new $1T'/1T'$ domain boundaries yield topologically unprotected 1D states that are
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20 dispersive in energy near the Fermi level and exhibit energy-dependent decay lengths. These
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22 results create new opportunities for exploring electron- and spin-based devices where charge
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24 carriers traveling along QSH edges might be deflected into trivial domain boundary modes in a
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26 controllable fashion.¹²
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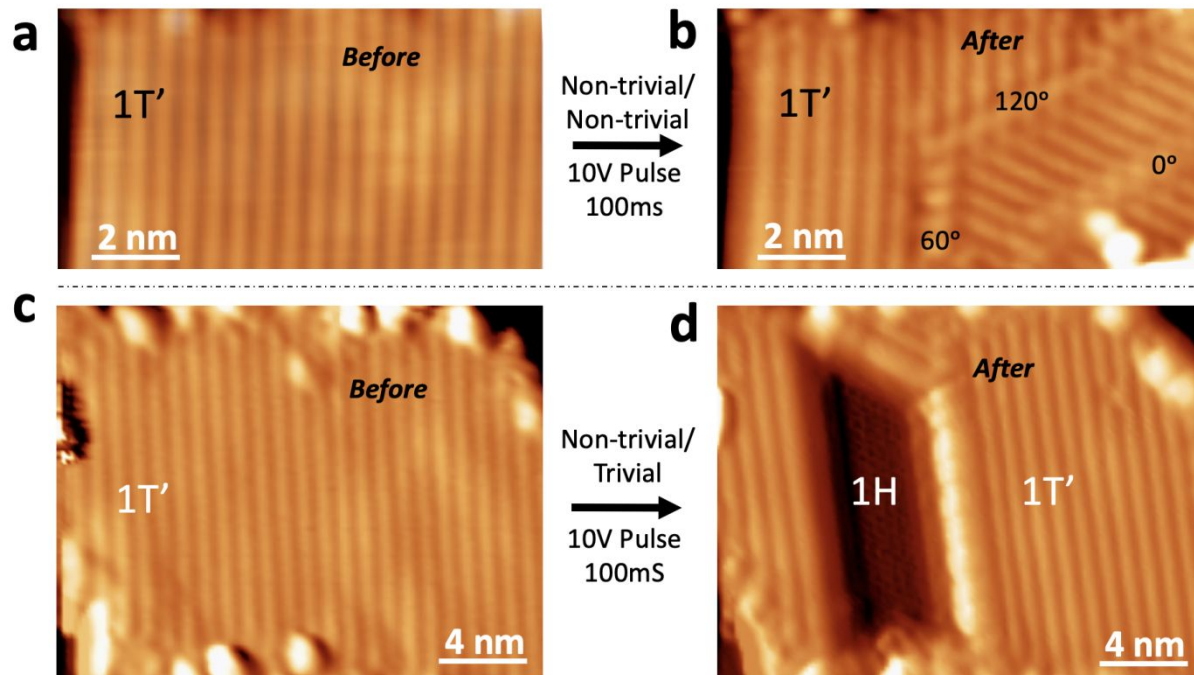


Figure 1: STM tip-induced structural change in monolayer 1T'-WSe₂. STM topographic images of a monolayer 1T'-WSe₂ island (a) before and (b) after applying a tip voltage pulse ($V_{\text{pulse}} = 10 \text{ V}$, $\Delta t = 100 \text{ ms}$, tip-surface separation = 6 \AA). The tip pulse creates 1T'/1T' domain boundaries having different rotational orientations. STM topographic images show a different island (c) before and (d) after an applied tip voltage pulse ($V_{\text{pulse}} = 10 \text{ V}$, $\Delta t = 100 \text{ ms}$, tip-surface separation = 6 \AA) induces a 1T' to 1H structural phase transition near the center of the island. $V_s = 1 \text{ V}$, $I_t = 10 \text{ pA}$, $T = 4.5 \text{ K}$ for all images. (Image intensity here is proportional to dz/dx (where z is height) in order to enhance contrast between regions having different structural phases.)

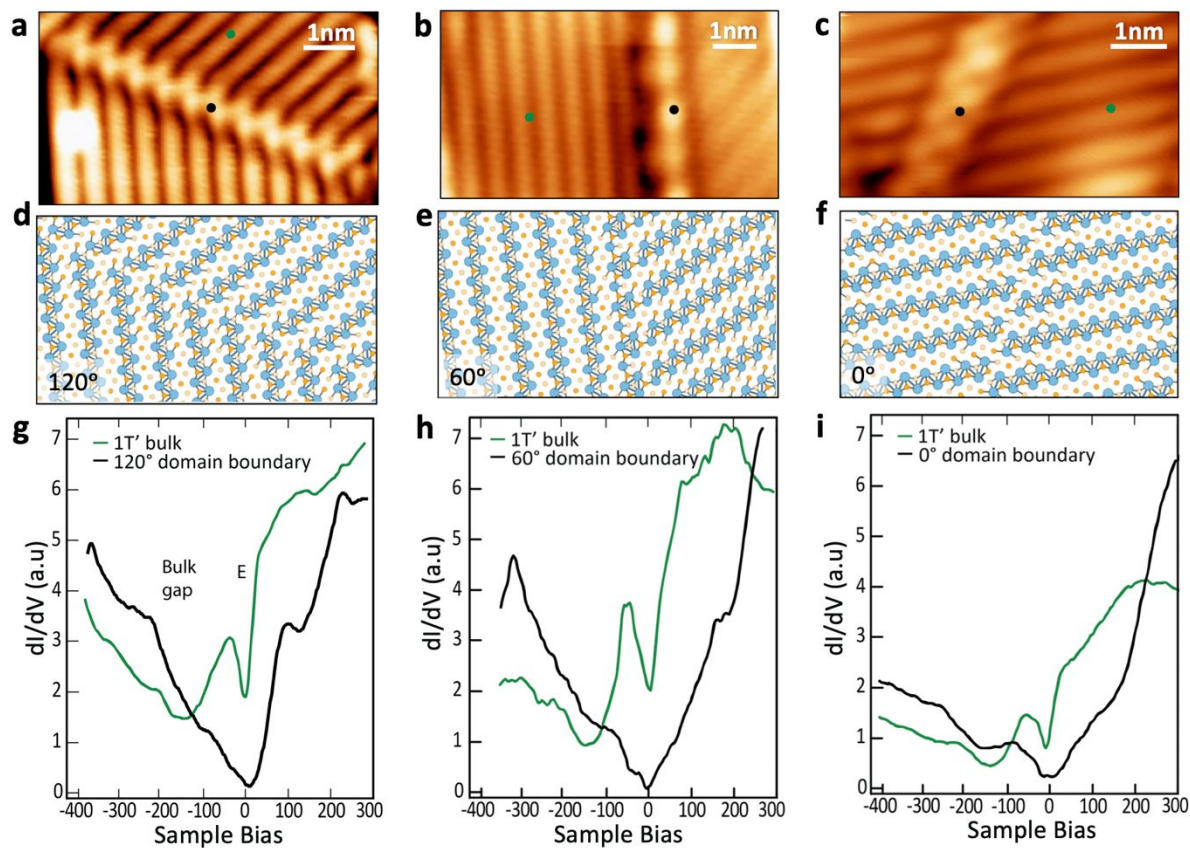


Figure 2: Structural and electronic properties of 1T'/1T' domain boundaries. STM images of (a) 120°, (b) 60°, and (c) 0° 1T'/1T' domain boundaries in 1T'-WSe₂ ($V_s = 1$ V, $I_t = 10$ pA, standard STM topographs). Relaxed structural models of (d) 120°, (e) 60°, and (f) 0° domain boundaries in 1T'-WSe₂ (calculated using DFT). STM dI/dV spectroscopy measured at (g) 120°, (h) 60°, and (i) 0° domain boundaries compared to the bulk for single-layer 1T'-WSe₂ (spectroscopy positions marked by black and green dots in (a)-(c)) ($f = 613.7$ Hz, $V_{ac} = 4$ mV, $T = 4.5$ K. Initial tunneling parameters for spectroscopy measurements: $V_s = -400$ mV, $I = 100$ pA).

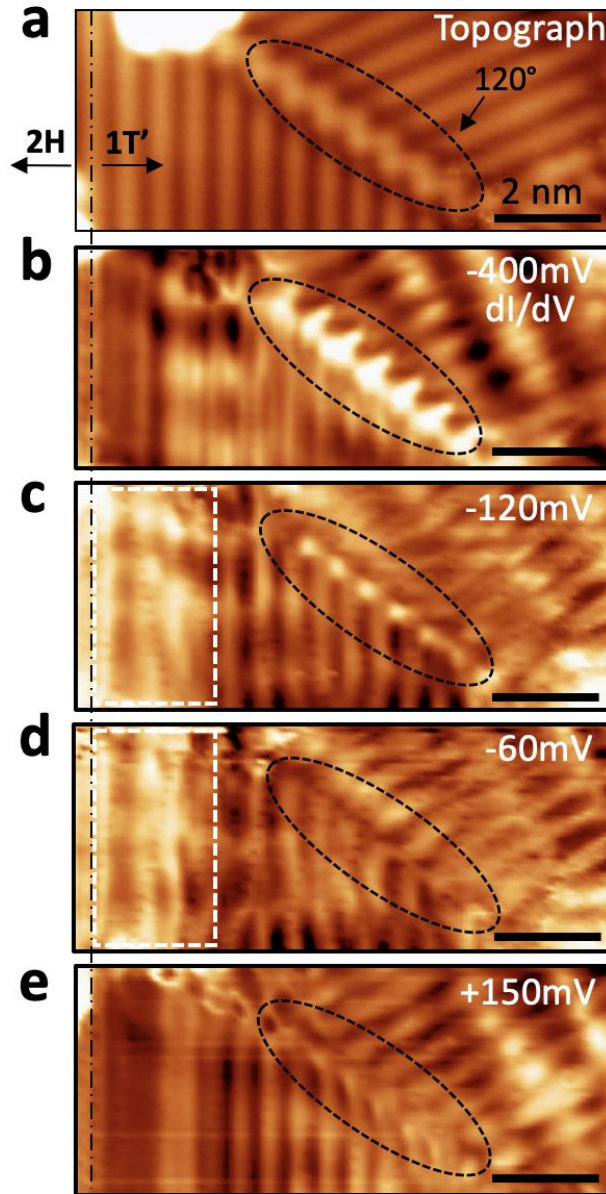


Figure 3: Comparison of electronic properties of 120° $1T'/1T'$ domain boundary and $1T'/1H$ boundary coexisting in single-layer WSe_2 . (a) STM image of a mixed-phase WSe_2 island with a 120° $1T'/1T'$ domain boundary (standard STM topograph). The $1T'/1H$ interface is marked by a vertical dashed line while the $1T'/1T'$ interface is outlined by a dashed oval ($V_s = 1$ V, $I_t = 10$ pA). dI/dV maps of the same area are shown for (b) $V_s = -400$ mV, (c) -120 mV, (d) -60 mV, and (e) $+150$ mV. Spectroscopy parameters: $f = 613.7$ Hz, $V_{ac} = 4$ mV, $I = 100$ pA, $T = 4.5$ K. Dashed white box outlines the topologically protected edge-state.

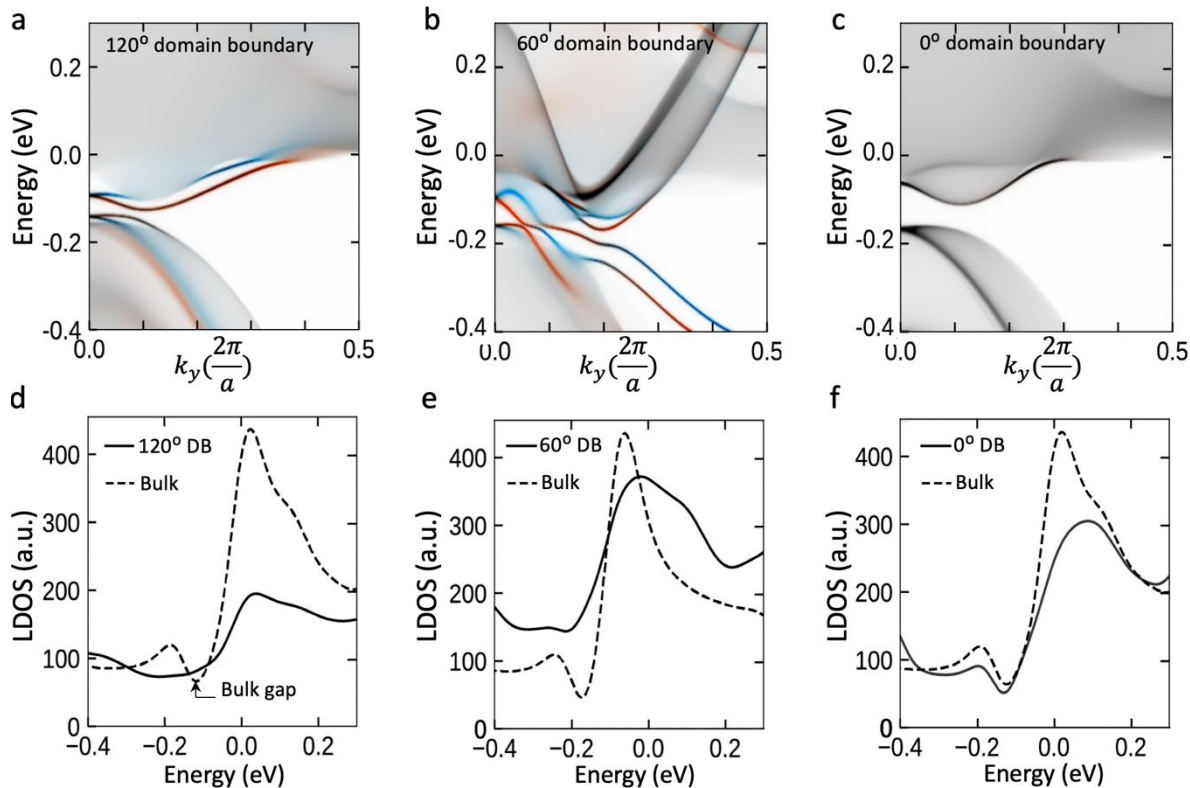


Figure 4: Calculated Band Structure and Local Density of States of Different 1T'/1T'

Domain Boundaries. Calculated band structure of (a) 120°, (b) 60°, and (c) 0° 1T'/1T' domain boundaries for 1T'-WSe₂ monolayer. Bulk states are grey, spin-polarized interface modes are red and blue. (d-f) Calculated LDOS of top-layer Se atoms at a domain boundary (black solid curve) compared to LDOS in the bulk (black dashed curve) for (d) 120°, (e) 60°, and (f) 0° 1T'/1T' domain boundaries. The Fermi level ($E = 0$) has been shifted to match experimental data (plots presented in (d)-(f) have been convolved with a Gaussian having $\sigma = 30$ mV to simulate level broadening effects²⁵).