magnitude too slow to produce $\nu$-weakening, except perhaps at temperatures of 500° to 700°C reached in experiments at coseismic slip velocities (4, 5, 14, 19).

The internal polycrystalline substructure of the nanospherules and nanofibers that we observed bears a striking similarity to microstructures found in shocked ductile metals (24). As in metals, the well-known ductility of calcite (25) may allow the $\sim$5- to 20-nm substructure to form by progressive development of nano-cell walls from dense dislocation networks generated by crystal plasticity. Plastic deformation, fracturing, and abrasion presumably generated the observed nanospherules from the starting “gouge.” To explain the chaining of nanospherules, producing the observed fiber structure and CPO, we note that oriented attachment at coherent nanoparticle interfaces is widely reported (26).

In principle, the mechanism of frictional slip that we propose (Fig. 3, A to C, and fig. S1) is similar to the Ashby-Verral model for superplasticity by diffusion-accommodated grain boundary sliding (GBS) (Fig. 3D) (30), but allows for frictional GBS and for intergranular cavitation (porosity generation by dilatation) when diffusive mass transport is too slow to accommodate GBS. Our findings imply that nanocrystalline PSZs developed in calcite faults can produce $\nu$-weakening, and hence seismogenic fault friction, by a mechanism of cooperative nanogranular or nanofiber flow plus diffusive mass transfer (Fig. 3, A to C), even in the upper crust where temperatures are generally considered too low to support diffusion or superplasticity at active fault slip rates. The reason that these processes are observed in our experiments is because diffusive mass transfer is dramatically accelerated by the nanogranular nature of the slip-zone rock that forms, and by water-enhanced grain boundary diffusion. A similar mechanism can also be envisaged to operate at coseismic slip rates, where the high temperatures generated will promote solid-state diffusion. Given the abundant recent observations of nanogranular fault surfaces in tectonically active terrains (17–19), and the anomalously high rates of diffusion found in nanomaterials (35, 36), the proposed mechanism may be relevant not only to faults cutting calcite-rich rocks such as limestones, but to crustal seismogenesis in general.

SOLID STATE THEORY

Quantum spin Hall effect in two-dimensional transition metal dichalcogenides

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Quantum spin Hall (QSH) effect materials feature edge states that are topologically protected from backscattering. However, the small band gap in materials that have been identified as QSH insulators limits applications. We use first-principles calculations to predict a class of large-gap QSH insulators in two-dimensional transition metal dichalcogenides with 1T' structure, namely, 1T'-MX2 with M = (tungsten or molybdenum) and X = (tellurium, selenium, or sulfur). A structural distortion causes an intrinsic band inversion between chalcogenide-$p$ and metal-$d$ bands. Additionally, spin-orbit coupling opens a gap that is tunable by vertical electric field and strain. We propose a topological field effect transistor made of van der Waals heterostructures of 1T'-MX2 and two-dimensional dielectric layers that can be rapidly switched off by electric field through a topological phase transition instead of carrier depletion.

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The discovery of graphene (1) has fueled vigorous investigation of two-dimensional (2D) materials (2), revealing a wide range of extraordinary properties (3–5) and functionalities (6, 7). Owing to their atomic thickness, 2D materials can be horizontally patterned through chemical and mechanical techniques (8). Moreover, the weak van der Waals (vdW) interaction between adjacent layers enables vertical stacking of different 2D materials, forming vdW heterostructures (9), which offer unprecedented opportunities for exploring quantum electronics at the nanoscale.
Quantum spin Hall (QSH) insulators (10–16) have an insulating bulk but conducting edge states that are topologically protected from backscattering by time-reversal symmetry. Quantized conductance through QSH edge states have been experimentally demonstrated in HgTe/CdTe (13, 14) and InAs/GaSb (17, 18) quantum wells. This could in principle provide an alternative route to quantum electronic devices with low dissipation. However, the realization of such QSH-based devices for practical applications is impeded by three critical factors: (i) band gaps of existing QSH insulators are too small, which limits the operating regime to low temperatures. This has motivated efforts to search for large-gap QSH insulators (19–26); (ii) the small number of conducting channels (e²/h per edge, where e is the elementary charge and h is Planck’s constant) results in a small signal-to-noise ratio; and (iii) efficient methods of fast on/off switching are lacking.

Here, we use first-principles calculations to show that 2D materials can provide a practical platform for developing topological electronic devices that may potentially overcome the above hurdles. Specifically, we predict a class of large-gap (~0.1 eV) QSH insulators in 2D transition metal dichalcogenides (TMDCs) MX₂ with M = (W, Mo) and X = (Te, Se, S). We demonstrate the feasibility of a vdW-heterostructured topological field-effect transistor (vdWHFET) made of 2D atomic layer materials. The proposed device exhibits a parametrically enhanced conductance through QSH edge channels in the “on” state and can be rapidly switched off via a topological phase transition by applying a vertical electric field.

Our findings have potential applications in low-power quantum electronics and spintronics and may enable topological quantum computing based on Majorana fermions (27–30).

Monolayer TMDCs (31)–MX₂ with M = (W, Mo) and X = (Te, Se, S) possess a variety of polytypic structures such as 1H, 1T, and 1T’–32, 33 shown in Fig. 1. The most-studied 1H structure is a sandwich of three planes of 2D hexagonally packed atoms, X-M-X, forming Bernal (ABA) stacking. In contrast, the three atomic planes in the 1T structure form rhombohedral (ABC) stacking. It has been known that the 1T structure in MX₂ is typically unstable in free-standing condition and undergoes a spontaneous lattice distortion in the x direction to form a period-doubling 2 × 1 distorted structure—i.e., the 1T’ structure, consisting of 1D zigzag chains along the y direction (Fig. 1C) (33); this structure is the subject of our work.

The electronic structures of various 1T’–MX₂ materials were obtained by many-body perturbation theory calculations (34). Figure 2A shows a typical band structure of 1T’-MX₂ using 1T’-MoS₂ as an example, and the results of the other five compounds are shown in fig. S1. Unlike its 1H or 1T counterparts, 1T’-MoS₂ is a semiconductor with a fundamental gap (Eg) of about 0.08 eV, located at Λ = ±(0.146 Å⁻¹)
The conduction and valence bands display a camelback shape near \( \Gamma \) in the 2D Brillouin zone (BZ) (see Fig. 2B), suggestive of band inversion with a large inverted gap at \( \Gamma \) of about 0.6 eV. The fundamental gap \( (E_g^\text{B}) \) and inverted gap \( (E_g^\text{d}) \) of all six 1T\(^{-}\)-MX\(_2\) are shown in Fig. 2C and listed in table S1. The inverted gap \( (E_g^\text{d}) \) was later used in the fitting of a low-energy \( \mathbf{k} \cdot \mathbf{p} \) model \( \langle 34 \rangle \). Because the 1T\(^{-}\) structure has inversion symmetry, we calculated the \( Z_2 \) index \((0 \text{ or } 1, \text{ indicating trivial or nontrivial topology}) \) based on the parity criterion \( \langle 35 \rangle \) of valence bands \( \langle \text{fig. S3} \rangle \) and found \( Z_2 = 1 \), indicating that 2D 1T\(^{-}\)-MX\(_2\) materials are in the QSH insulator phase.

To understand the origin of the inverted band structure near \( \Gamma \), we analyzed the orbital character of the bands and found that the valence band mainly consists of metal \( d \)-orbitals and the conduction band mainly consists of chalcogenide \( p \) orbitals, as expected. The unexpected band inversion arises from the period doubling of the metal chain in the 1T\(^{-}\) structure, which lowers the metal \( d \) orbital below chalcogenide \( p \) orbital \( \langle \text{Fig. 2A} \rangle \) around \( \Gamma \). The large band inversion at \( \Gamma \) \((\sim0.6 \text{ eV}) \) is an intrinsic characteristic of the 1T\(^{-}\) structure and already takes place without including spin-orbit interaction in the first-principles calculations. In the absence of spin-orbit coupling, this band inversion leads to the appearance of two Dirac cones centered at finite momenta on Y-Y in 2D BZ. Spin-orbit coupling then opens up a fundamental gap of 0.08 eV at the Dirac points \( \langle \text{Fig. 2A}, \text{inset, and fig. S3} \rangle \), leading to a QSH insulator through a similar mechanism as the Kane-Mele model \( \langle 10 \rangle \); we show that this is the case by using a low-energy \( \mathbf{k} \cdot \mathbf{p} \) Hamiltonian \( \langle 34 \rangle \) \( \langle \text{fig. S4} \rangle \). In addition to MoS\(_2\), we found that all other five 1T\(^{-}\)-MX\(_2\) have \( Z_2 \) nontrivial band topology.

**Fig. 4. Proposed vdW-TFET.** (A) Topological phase diagram of 1T\(^{-}\)-MoS\(_2\) as a function of vertical electric field. The critical field strength is \( \pm0.142 \text{ V/Å} \), marked by two green dots. (B) Schematic of vdW-TFET. The central component (C) is a vdW heterostructure of alternating monolayer 1T\(^{-}\)-MX\(_2\) and mono-/multilayer wide-gap insulators such as hBN, with horizontal width as narrow as \( \sim20 \text{ nm} \). Carriers (charge or spin) are injected from the source electrode and ejected into the drain electrode. On/off switch is controlled by vertical electric field through the top and bottom gates. Mono-/multilayer wide-gap insulators effectively screen the interaction between adjacent MX\(_2\) layers, preventing them from detrimental topological phase change and parametrically increasing the number of edge channels.

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**Fig. 3. Vertical electric field induced a topological phase transition.** Shown are calculations for monolayer 1T\(^{-}\)-MoS\(_2\) under electric fields of (A) 0, (B) 0.142, and (C) 0.2 V/Å, respectively. The first, second, and third rows show the bulk band structure, edge density of states, and edge spin polarization, respectively.
The QSH insulator phase in 2D 1T-MX₂ leads to helical edge states that are protected from localization and elastic backscattering by time-reversal symmetry. Figure 2D displays the electronic response time scale (46, 49). In addition, the vdW heterostructure and the 2D nature of vdW-TFET make it convenient to both vertically and horizontally pattern devices at a large scale. The short decay length of helical edge states allows the minimum horizontal feature size of vdW-TFET to be ~20 nm, enabling high device density on a chip. Horizontally patterned TFET devices of such nanoscale size have large edge-to-area ratios, which will greatly reduce the contribution of thermally populated bulk carriers to the total electric current and hence enhance the on/off ratio. The predicted QSH insulators and their vdW heterostructures may provide a platform for realizing low-dissipation quantum electronics and spintronics (38–43).

From a materials perspective, the 1T’ structure in monolayer MoS₂ has recently been observed in high-resolution transmission electron microscopy experiments (39). We have checked the structural stability of 1T’ by carrying out first-principles calculations (fig. S7). Although MoS₂, MoSe₂, MoTe₂, WS₂, and WSe₂ are more stable in the 1H structure, we find that there exists a large energy barrier of more than 1 eV/MX₂ between 1T’ and 1H for all MX₂ suggesting that the 1T’ structure can be stabilized under appropriate chemical (47), thermal, or mechanical conditions. Moreover, we found that the 1T’ structure is unstable and will spontaneously relax to 1T, like in a Peierls distortion. We also verified the stability of the 1T’ structure by computing its phonon dispersion (fig. S8). The absence of imaginary frequencies throughout the 2D BZ provides direct evidence of the structural stability in all 1T’-MX₂.

Encouragingly, the desired 1T’ structure with intrinsic band inversion is the natural ground-state structure of WTe₂ (39), also confirmed by our stability calculation. Although monolayer WTe₂ was found to be a semimetal from our calculation, the small negative fundamental gap can be easily lifted by a tensile strain as small as ~1% (fig. S9). We also found that hBN used as a dielectric in the aforementioned vdW-TFET device has little effect on the electronic structure of 1T’-WTe₂ near the Fermi level (fig. S10). These attractive features make WTe₂ potentially the most promising 2D TMDC for realizing the QSH effect and related vdW devices.

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SUPPLEMENTARY MATERIALS
www.sciencemag.org/content/346/6215/1344/suppl/DC1
Supplementary Methods
Figs. 51 to 51D
Tables S1 and S2
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