Chirality is essential for various phenomena in life and matter. However, chirality and its switching in electronic superlattices, such as charge density wave (CDW) superlattices, remain elusive. In this study, we characterize the chirality switching with atom-resolution imaging in a single-layer NbSe₂ CDW superlattice by the technique of scanning tunneling microscopy. The atomic arrangement of the CDW superlattice is found continuous and intact although its chirality is switched. Several intermediate states are tracked by time-resolved imaging, revealing the fast and dynamic chirality transition. Importantly, the switching is reversibly realized with an external electric field. Our findings unveil the delicate switching process of chiral CDW superlattice in a two-dimensional (2D) crystal down to the atomic scale.
Chirality refers to the character of an object or state that cannot be superimposed on its mirror image, which can be used to tune various intriguing properties in condensed matter, such as solitons, Luttinger liquids, and gyrotropic order. Due to the insulator–metal transition accompanied by phase change, charge density wave (CDW) with its chirality switching have potential applications in ultrafast switches and memories, especially when combining with the laser beam. Although the optical approach is useful in the time scale, the different interlocking manners of the triangular-shaped SOD clusters can be used to distinguish their chiralities from each other, even if the STM images do not have the atomic resolution. Such chirality can be intuitively understood because the SOD superlattice has a \((\sqrt{13} \times \sqrt{13})R13.9^\circ\) relationship with respect to the top-layer Se atom lattice, and this relationship intrinsically corresponds to two mirror domains.

Visualisation of atomic lattice before and after chiral CDW chirality switching. Next, we concentrate on the chirality change, and we find that the atomic arrangement of the CDW superlattice is still continuous and intact after the chirality change. Figure 2a–c shows the movement of a domain boundary (DB) in the chiral CDW superlattices. By setting a large current (2.0 nA) for the rightward scan, the DB can be moved rightward as denoted by the yellow dashed line (Fig. 2a) and the red one (Fig. 2b), corresponding to the expansion of the L domain and shrinking of the R domain. It further transforms into one domain with a single L chirality, as shown in Fig. 2c. To better illustrate this chiral CDW switching process at the atomic scale, we superpose the models of both the SOD superlattice and atomic lattice onto the STM images, as shown in Fig. 2d–f. The green and blue balls represent the top-layer Se atoms within the SOD clusters with left and right chiralities, respectively, while the gray balls represent the Se atoms out of the SOD clusters. The STM images of Fig. 2a–c without color-coding are also shown in Supplementary Fig. S1 for comparison.

Importantly, two features remain unchanged after the DB movement of the chiral CDW superlattice, as exemplified in Fig. 2a–f: (1) The well-ordered hexagonal lattice of balls which describes the positions of the top Se lattice, is continuous not only at each single chiral CDW domain but also across the DB. Namely, the top-layer Se atoms keep the same periodicity with the continuous arrangement. (2) The orientations of all triangular SOD at the two chiral CDW domains keep the same. These two features together suggest that the whole Se–Nb–Se structure is preserved over the island, with the continuous 1T structure in the two chiral CDW domains. These features are also illustrated in the side view in Supplementary Fig. S1.

To the best of our knowledge, this is the pioneering in-situ atomic-resolution analysis of the DB movement in 2D chiral CDW materials. The intact and robust atomic structure is an important virtue for future chiral CDW-based devices, as it avoids the atomic defect at the DB and will be favorable for the reversible switch application. More examples of controlling the DB movement direction by using different current setpoint values are shown in Supplementary Fig. S2 and Supplementary Movie. Notably, although there are some “imperfect SOD” at the DB (labeled by both green and blue balls), the intact and robust atomic structure is still preserved.

These geometric features obtained by the STM images are also reflected in the fast Fourier transform (FFT) images (Fig. 2g–i). All FFT images reveal a single set of reciprocal spots (as marked by the outer brown circles in Fig. 2g–i, labeled as \(a_{1g}^*\) and \(a_{2g}^*\)), which originated from the well-ordered Se lattice. In contrast, another two sets of spots (marked by green and blue circles, labeled \(b_{1g}^*\) and \(b_{2g}^*\)) with mirror symmetry are shown in Fig. 2g, h, which can be ascribed to the chiral CDW superlattice. The intensities of the two sets of spots show a change following the shift of the chiral domain boundary. The spots marked by green circles have larger intensity, corresponding to expanding of the L domain in the real space (Fig. 2a, b). The spots
in Fig. 2i are reduced to one set, corresponding to the single chiral domain (Fig. 2f). This kind of intensity change is also illustrated clearly by the magnified FFT image (Supplementary Fig. S3).

Tracking the dynamic process in chiral CDW superlattice. After resolving the atomic arrangement with the ultrahigh spatial resolution, we further investigated the dynamic process of chiral change in CDW superlattice with high time resolution. Two metastable statuses at the chiral DB are tracked by STM observations, as shown by status I and II in Fig. 3a, b, respectively. The fuzzy SOD sites (denoted by dashed triangles) at the DB belong to either (or both) L or R domain. The switching rate is so high that instant SODs are difficult to capture by normal STM scanning (500 ms/scan line) and thus fuzzy features are present. These fuzzy SOD sites look like touching or overlapping, different from the ideal SOD superlattice with hexagonal lattice. Taking the region around one fuzzy SOD cluster in the 3D STM images (Fig. 3d, e) as an example, the SOD cluster has a change of the residence time at the L and R domains, respectively.

To track the dynamic process of the chiral switching, we performed current–time (I–t) spectroscopy (0.05 ms/point) by holding the STM tip over a fuzzy region (marked by the crosses in Fig. 3a, b and Fig. 3d, e) with a constant height mode. Such I–t measurement is a reliable STM method described in the literature for studying the dynamic process of on-surface adsorbates, such as tautomerization27,28, hopping or rotation29–32, bond breaking33, and even orbital dynamics34. In our cases, when the fluctuating SOD site is just beneath the tip (e.g., at the L domain in Fig. 3a, d), the tip–sample distance is smaller and the local density of states is enhanced, thus the tunneling current is higher. When the fluctuating SOD is away from the tip (e.g., in the R domain in Fig. 3b, e), the tip–sample distance enlarges and the local density of states reduces; thus, the current will be lower.

In this way, the low and high current values are obtained following the SOD occupation change in two chiral domains, which provide direct evidence of the fast-dynamic process of the chirality switching. The current–time (I–t) curve in Fig. 3f shows the evolution of the current in the duration of 40 ms. The low state is at 100 pA, and it changes to a high state at 240 pA for ~2.5 ms, and then changes back.

By increasing the tunneling current, the switching events become more frequent (Fig. 3g for the I–t curve with a setpoint of

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**Fig. 1 STM topography and schematic model of chiral CDW superlattice.** a (upper panel) Top view of SOD model showing thirteen Nb atoms, six top Se atoms, and six bottom Se atoms, with three-fold symmetry; (lower panel) side view of SL 1T-NbSe2 on a BLG/SiC(0001) substrate. The top- and bottom-layer Se atoms are projected from the upper to the lower panel to show the relative positions. b CDW-resolved STM topography around a domain boundary of SL 1T-NbSe2. The top-Se close-packed direction is marked by the black dashed-arrow. c Left (right): Zoomed-in atomic-resolved image in the L (R) domain marked by the green (blue) frame in (b). The SOD superlattices in these two domains are labeled by green and blue triangles, respectively. The SODs arrangements are represented by the clockwise (CW) and counterclockwise (CCW) arrows. d Atomic model of SL 1T-NbSe2 with chiral CDW domains. The atomic lattice of the top layer Se is indicated by the lattice of orange balls, and the SOD clusters in these two domains are labeled by green and blue triangles encircled by hexagram stars, respectively. Green and blue arrows crossing the adjacent SOD triangles represent the chiral CDW close-packed directions, which are rotated by ±14° with respect to the top-layer Se atom close-packed direction (bL1 and bL2 for the L domain, and bR1 and bR2 for the R domain). Scanning parameters: b, bias voltage $V_b = -1.5$ V, tunneling current $I_t = 1$ nA; c, $V_b = -1.5$ V, $I_t = 2$ nA.
400 pA). The statistical analysis of the residence time distribution (displayed in Supplementary Fig. S4) reveals that it can be fitted by an exponential curve \( y = y_0 + Ae^{Rx} \), where the switching rate \( R \) is determined by the fitting, \( y_0 \) and \( A \) are constants. By plotting the switching rate against the tunneling current with a log scale, a linear relationship with a slope of 1.042 ± 0.055 is obtained (Fig. 3h), indicating a one-electron process\(^3\) during the chiral CDW switching. Thus, the switching rate can be tuned by the tunneling current.

Based on these experimental findings, the mechanism of the chiral CDW switching process is illustrated by the schematic in Fig. 3i. Regarding the intermediate CDW statues I and II, they jump fast over the low energy barrier, back and forth. When the intermediate state obtains sufficient energy to jump over the higher barrier, it falls into more stable status without fuzzy features (status III). In short, using the \( I-t \) curve to monitor the dynamic switching process of chiral CDW superlattice, we observed several intermediate states at the chiral DB. As far as we know, this is the pioneering dynamic study together with the atomic characterization on a chiral 2D CDW system till now and it may be extended to the bulk non-chiral CDW systems with various DBs\(^3\) in future.

Reversible switch of chiral CDW superlattice induced by an external electric field. After visualizing the switching process of the chirality at the atomic scale, we further managed to switch the chiral CDW islands as a whole reversibly, as shown in Fig. 4a–c. A schematic of the experimental setup is shown in Fig. 4g. The switching process from L to R chirality was realized by applying voltage pulses of \( \approx -5.5 \) V at a lateral distance of \( \approx 5 \) nm from the STM tip to the edge of the 1T-NbSe\(_2\) island. The reversible switching process from R to L chirality was realized by applying voltage pulses of \( \approx -6.6 \) V. Taking advantage of our simultaneous imaging of chiral CDW superlattice with atomic resolution (Supplementary Fig. S5), we demonstrate that the atomic arrangement is still continuous and intact although the chirality of CDW is switched.

Note that even if the SOD protrusions are not atomically resolved in the large-scale STM images, the chiralities and chiral change of the CDW domains can still be identified in another easy way. As shown in Fig. 4a–c, the SOD close-packed directions have rotation angles of \( \pm 14^\circ \) compared to the direction of the island edge, with \( +14^\circ \) corresponding to L and \( -14^\circ \) corresponding to R, similar to those defined in Fig. 1c, d. The magnified images of the triangular SOD superlattice of the R and L domains...
are shown in Fig. 4d–f, also demonstrating the chirality change well. By moving the tip in the lateral direction and repeatedly testing the threshold voltage \( (V_{th}) \) to switch the chirality of CDW, we obtained the plot shown in Fig. 4h. By fitting \( V_{th} \) against the lateral distance, a linear relationship was obtained, demonstrating that the switching process was controlled by the electric field effect. As far as we tested, at a lateral distance of ~20 nm, the chirality can still be switched with a voltage-pulse of 10 V. In addition, the chirality of the island size of 50 nm can be switched as a whole, suggesting a domino-like effect in changing the chirality. This effect should be associated with the collective electronic nature of the CDW. Notably, the CDW chirality of the 1T-NbSe\(_2\) islands is very robust can be stable even above room temperature (RT). This advantage further facilitates the application of chiral CDW-based nanodevices above RT.

**Discussion**

To induce the chiral domain switch, we have used electron tunneling and voltage pulse methods, and also realized it by dipping the STM tip into the sample with a controlled depth. Thus, various techniques (including mechanical force, heat, doping, electric field or voltage, and laser) to switch the chirality of the CDW superlattice are expected as long as some form of excitation energy is input to the system (Fig. 3i). Our study opens new
In conclusion, with high-spatial-resolution imaging, fast-tracking I-t spectroscopy, and voltage-pulse manipulation, we have comprehensively investigated the atomic structure, dynamic process, and reversible manipulation of the chiral CDW superlattice, in a 2D crystal of SL 1T-NbSe₂. In our study, three important features are identified: (1) The top-layer Se atom lattice of NbSe₂ is unchanged although the CDW chirality changed. (2) Chirality switching of CDW superlattice is fast and intermediate states exist in the dynamic switching process. (3) The chiral switching can be reversibly manipulated in the CDW domains, induced by an external electric field. Our findings provide in-depth understanding of the structure and manipulation of chiral CDW systems down to the atomic level. The 2D chiral CDW systems may lay a versatile avenue for triggering the chirality switch by different methods and may be applicable for sensors or electronic devices based on 2D chiral CDW materials.

Data availability
The main data supporting the findings of this study are available within the article, and Supplementary Information. Extra data are available from the corresponding author upon reasonable request.

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Methods
Sample preparation and STM measurements. MBE and STM experiments were performed by a Unisoku ultrahigh-vacuum system (base pressure ~1 × 10⁻¹⁰ mbar) comprising an MBE chamber and LT-STM chamber. Epitaxial graphene was grown by thermal decomposition of 4H-SiC(0001) at 1200 °C for 40 min. NbSe₂ islands were prepared using the MBE method at a deposition rate of ~0.002 ML/min. An elemental Nb rod was used as the metal source in the e-beam evaporators (Focus Ltd.), whereas an elemental Se source in a Knudsen cell was heated to 120 °C.

The deposition ratio of Se to Nb was larger than 20:1. During the film deposition, the background pressure in the chamber was ~3.0 × 10⁻⁹ mbar. The BLG/SiC substrate was stored at ~550 °C during the deposition and post-annealing processes to improve the diffusion and desorb excess Se. STM imaging was performed using mechanically cut Pt–Ir tips at the liquid helium temperature (4.2 K). The current-time (I-t) curves were acquired by positioning the tip at the fuzzy site at the domain boundary with a setpoint (I₀ = 100 pA), then turn off the feedback loop and record the current with time.

Fig. 4 Reversible switching of chiral CDW superlattice induced by an external electric field. a–c Large-scale STM images of the reversible switching of chiral CDW superlattice of a 1T-NbSe₂ island, realized by applying voltage pulses. Two CDW close-packed directions are represented by green and blue arrows, respectively, which are rotated around ±14° with respect to the island edges (labeled by the white dashed double arrows). d–f Magnified images of the areas in (a–c), showing the chirality identified by the relative arrangement of neighboring triangular SOD clusters. The top-layer Se close-packed direction is marked by the black dashed-arrow. g Schematic of the experimental setup. The bias voltage is applied between the tip and the substrate to form an external electric field. h Vth (magnitude of threshold voltage to switch the chirality of CDW) as a function of the lateral distance. The linear fit indicates an electric field effect (0.380 ± 0.016 V/nm) during the chirality switching. The error bar on each point gives the deviation from multiple measurements, which is too small to display. Scanning parameters: h, f, –1.5 V, 20 pA.
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