Quasi-two-dimensional superconductivity from dimerization of atomically ordered AuTe$_2$Se$_{4/3}$ cubes

J.G. Guo 1, X. Chen 1,2, X.Y. Jia 3, Q.H. Zhang 1, N. Liu 1,2, H.C. Lei 4, S.Y. Li 3,5, L. Gu 1,6,7, S.F. Jin 1,6 & X.L. Chen 1,6,7

The emergent phenomena such as superconductivity and topological phase transitions can be observed in strict two-dimensional (2D) crystalline matters. Artificial interfaces and one atomic thickness layers are typical 2D materials of this kind. Although having 2D characters, most bulky layered compounds, however, do not possess these striking properties. Here, we report quasi-2D superconductivity in bulky AuTe$_2$Se$_{4/3}$, where the reduction in dimensionality is achieved through inducing the elongated covalent Te–Te bonds. The atomic-resolution images reveal that the Au, Te, and Se are atomically ordered in a cube, among which are Te–Te bonds of 3.18 and 3.28 Å. The superconductivity at 2.85 K is discovered, which is unraveled to be the quasi-2D nature owing to the Berezinsky–Kosterlitz–Thouless topological transition. The nesting of nearly parallel Fermi sheets could give rise to strong electron-phonon coupling. It is proposed that further depleting the thickness could result in more topologically-related phenomena.
he dimensional reduction or degeneracy usually induces the significant change of electronic structure and unexpected properties. The monolayer, interface and a few layers of bulky compounds are typical resultant forms of low dimensionality. The two dimensional (2D) material, for instance, graphene, is found to have a linear energy dispersion near Fermi energy ($E_F$) and possess a number of novel properties \(^1\). The monolayer MoS$_2$ exhibits a direct energy gap of 1.8 eV \(^4\) and pronounced photoluminescence \(^6\), in contrast to trivial photoresponse in bulky MoS$_2$ with an indirect band-gap.

2D superconductivity (SC), a property closely related to dimensional reduction, has been observed in a variety of crystalline materials like ZrNCl$_6$, NbSe$_2$, and MoS$_2$ recently through the electric-double layer transistor (EDLT) \(^9\) technique. Many emerging properties, i.e., the well-defined superconducting dome, metallic ground state and high upper critical field \(^8, 10\), significantly differ from those of intercalated counterparts. Besides, the lack of in-plane inversion symmetry in the outmost layer of MoS$_2$/NbSe$_2$ with strong Ising spin-orbital coupling induces a valley polarization \(^7\). In the scenario of low-dimensional interface, the unexpected 2D SC \(^10, 11\), the remarkable domed-shaped superconducting critical temperature ($T_c$) \(^12\), pseudo-gap state \(^13\), and quantum criticality \(^14\) have been demonstrated in La(Al,Ti)O$_3$/SrTiO$_3$(001) \(^15\). Very recently, the interface between Bi$_2$Te$_3$ and FeTe thin films displayed 2D SC evidenced by Berezinsky–Kosterlitz–Thouless (BKT) transition at 10.1 K \(^15\). The tentative explanations are related to the strong Rashba-type spin–orbit interactions in the 2D limit.

At the moment, the way to fabricating low-dimensional materials generally involves molecule beam epitaxy and exfoliation from the layered compounds. The top-down reduction processes usually are sophisticated and time consuming for realizing scalable and controllable crystalline samples. There are other chemical routes to tuning dimensionality by means of either changing the size of intercalated cations or incorporating additional anions. It is reported that increasing the size of alkaline-earth metals (Ae) (Ae=Mg, Ca, and Ba) between [NiGe] ribbons can reduce three dimensional (3D) structure to quasi-1 dimensional one \(^16\). In addition, the ternary CaNiGe can be converted into ZrCuSiAs-type CaNiGeH by forming additional Ca–H bonds, which exhibits different properties owing to the emergence of 2D electronic states \(^17\). The metastable Au$_{1-x}$Te$_x$ (0.6 < x < 0.85) show an a-type polonium structure \(^18\), \(^19\), in which the Au and Te disorderly locate at the eight corners of a simple cubic unit cell. The $T_c$ fluctuates in the range of 1.5–3.0 K, but the mechanism of SC has been barely understood \(^20\). Besides, the equilibrium phase Au$_{x}$Te$_{2-x}$ known as calaverite, is a non-superconducting compound, in which distorted AuTe$_6$ octahedra are connected by Te–Te dimers \(^21\).

Through incorporating more electronegative Se anions, we fabricate a new layered compound AuTe$_x$Se$_{4/3}$ by conventional high temperature solid-state reaction. In a basic cube subunit, the Se anions attract electrons from Te and lead to the ordered arrangement of Au, Te and Se atoms. The cubes stack into strip through Te–Te dimers at 3.18 Å and 3.28 Å along the a- and b-axis, respectively, which composes 2D layers due to the existence of weak Te–Te interaction (4 Å) along the c-axis. Electrical and magnetic measurements demonstrate that the SC occurs at 2.85 K. Furthermore, this SC exhibits 2D nature evidenced by the BKT transition in the thin crystals. The observed results are interpreted according to the crystallographic and electronic structure in reduced-dimensionality.

Results

Structural characterization. Figure 1a shows the scanning electron microscope (SEM) image of AuTe$_x$Se$_{4/3}$ crystal, and the typical shape is like a thin rectangle. Its anisotropic morphology implies that there exist weak bonds and the weakest one determines the most cleavable face in AuTe$_x$Se$_{4/3}$. The elemental mapping shows homogeneous distribution of Au, Te, and Se atoms. The atomic ratio is Au:Te:Se = 25:46:29, as EDS pattern in Fig. 1b. Most reflections except a few weak peaks in the experimental powder X-ray diffraction (PXRD) pattern (Supplementary Fig. 1a) of as-grown sample can be indexed based a triclinic unit cell. Efforts for determining the crystal structure by solving PXRD pattern have failed.

We exfoliated the crystals by Scotch tape and obtained thin enough samples for observing the HAADF images. As the contrast of HAADF image exhibits a $Z^{1/2}$-dependent relation, where Z is the atomic number, different kinds of atom columns can be directly distinguished (i.e., in HAADF images, the largest contrast indicates Au columns). Figure 1e delineates the atomic arrangement of the ab-plane of AuTe$_x$Se$_{4/3}$. It can be seen that elongated atoms locate equispaced at 2.69 Å along two orthogonal directions based on the peak interval in Fig. 1c. The corresponding selected area electron diffraction (SAED) pattern along [1−3] zone axis in Fig. 1d confirms the tetragonality of the ab-plane. It is noted that two kinds of superstructure spots are present, one being along a* with a new a*/3 periodicity corresponding to a triple in real space and other plane spacing is a*/3 + b*.

After tilting the ab-plane by ~13° along diagonal Kikuchi line (Supplementary Fig. 2), clear planar 2×2 blocks appear as the image in Fig. 1h. Closer examination of a block reveals that two separated atoms exist at the center spots and hence elongated feature shows up, which actually comes from the tilted top and bottom atoms in a block (Supplementary Fig. 3). Each block is separated by 3.18 Å and 3.20 Å along transversal and longitudinal directions as indicated by the red and blue dash lines in Fig. 1f and i, respectively. The bond lengths are periodically modulated by two short bonds and one long bond (~2.69 Å–3.20 Å) and (~2.69 Å–2.69 Å–3.20 Å···), respectively. Figure 1g, j and m show the SAED patterns along the [00−1], [−10−3], and [−100] zone axis. Again, one can see that a modulation of a*/3 exists. The HAADF images of Fig. 1k and i present that the bright spots are sandwiched by two weak spots separated by 2.69 and 3.20 Å, indicating the atomic distributions of front/back face differ from those of middle layer in each block. To observe how the blocks stack along the weakly combined direction, namely the c-axis, we prepare narrow samples (~50 nm) by focused ion beam (FIB) milling. The HAADF image in Fig. 1n reveals that the building blocks are 3.20 Å and 3.47 Å apart along two directions, as shown in Fig.1l, respectively. So the periodic bond lengths along c-axis can be described as ~2.69 Å–2.69 Å–3.47 Å···.

On the basis of above observations, the crystal structure can be regarded as the stacking of cube subunits in real space. The stereotyope picture of a block is drawn in Fig. 2a. Inside the block, all of the atoms are orderly arranged and connected by bonds with identical length 2.69 Å. Each Au atom is four planar-coordinated by Te, and each Se atom is connected by three Te atoms. The body center is vacant, which could account for the elongated spots in Fig. 1d. The atomic distribution of front/back face of the cube is AuTe$_x$Se$_{4/3}$, while the middle layer is Au$_{3}$Te$_4$. This ordered structure is intimately related to the disorder phase Au$_{1−x}$Te$_x$ (0.6 < x < 0.85) with simple cubic structure. The arrays of cubes in the ab-plane are drawn in Fig. 2b. There are one Te–Te dimer (3.18 Å) and three Te–Te dimers (two at 3.18 Å and one at 3.28 Å) in the front/back face and middle layer, respectively. Figure 2c and e shows the front/back and middle strips that are constructed by two kinds of layers. The corresponding electron density difference (EDD) slices from DFT calculations are shown in Fig. 2d and 2f, respectively. One can see that although covalent bonds dominate the whole cube,
the Au atoms lose electrons in Au–Te bonds and Se atoms attract electrons to some extent. It is noted that the Te–Te bonds of middle layer accumulate more electrons than that of front/back face. These covalent Te–Te bonds among cubes suggest that the electrons can mainly populate in the middle layers along the a-axis and are perturbed by the coordinating Se atoms. Besides, the longer Te–Te bond lengths (3.28 Å) link horizontal-shifted strips, forming 2D stacking of cubes in the ab-plane.

The determined crystal structure of AuTe2Se4/3 is shown in Fig. 2g (see detail crystallographic parameters in Supplementary Table 1). It possesses a triclinic unit cell with P-1 (No. 2) symmetry and the lattice parameters are \(a = 8.85\, \text{ Å}, \quad b = 8.43\, \text{ Å}, \quad c = 9.28\, \text{ Å}, \quad \alpha = 77.24^{\circ}, \quad \beta = 95.53^{\circ}, \quad \gamma = 72.36^{\circ}\). The atomic ratio is Au:Te:Se = 3:6:4 deduced from crystallographic symmetry, which shows a little discrepancy from the measured value owing to the uncertainty in the EDS method. One can see that the spacings among cubes are 3.18 Å, 3.20 Å and 3.47 Å along the a-, b- and c- axis, respectively. The longest Te–Te distance between the ab-plane is \(\sim 4\, \text{ Å}\), which is larger than the Te–Te bond length in element Te22, the Van der Waals bonds of MoS2 (3.48 Å)23 and graphite (3.37 Å)1. The simulated PXRD pattern based on the present crystallographic parameters is plotted in Supplementary Fig. 1b. All the indices of high-intensity peaks of experimental PXRD can match the simulated ones very well except minor impurity peaks. Furthermore, the atomic arrangements viewed from [1–39], [00–1], [−10–3], and [−100] zone axes are highly consistent with experimental HAADF images (Supplementary Fig. 3), demonstrating the validation of present structure within the accuracy of electron microscopy.

Figure 3a presents the XPS pattern of Au 4f, Te 3d, and Se 3d core-level measured at 300 K. The binding energy of Te 3d5/2, 573.2 eV, is close to the value in element Te (573.0 eV), indicating the 5p orbitals of Te are not fully occupied and responsible for the conducting electrons. The binding energy of Au 4f7/2 is 84.6 eV, which is close to 84.4 eV (Au+) and smaller than 85.2 eV (Au3+)24. For the Se 3d5/2, the binding energy is 54.2 eV, which is close to the value in CdSe (54.4 eV) and obviously smaller than that in Se element (55.1 eV). Therefore, the charge balanced formula of AuTe2Se4/3 can be written as \([\text{Au}(1+)][\text{Te}^{2\times}(5/3^2-\delta)+][\text{Se}^{2\times}]_{\mu/3}\). Figure 3b shows the band structure of AuTe2Se4/3. The flat bands lead to the multiple Van Hove singularities in the plots of density of states. It can be seen that there are four bands, denoted as I, II, III, and IV bands, crossing the \(E_F\). The I, III, and IV bands cross the \(E_F\) along G–F, G–B, and Z–G in first Brillouin zone. The II band only crosses the \(E_F\) along G–F, which exhibits a nearly flat dispersion along F–Q with quasi-1D character. Figure 3c shows the projected density of states (PDOS) around the \(E_F\). The total DOS at the \(E_F\) is estimated to be 6.5 states eV\(^{-1}\) per unit cell, and the contribution of Te 5p states is \(\sim 70\%\), which are the primary source of conductive electrons. The whole Fermi surface of AuTe2Se4/3 and four separated sheets are individually plotted in Fig. 3d. One can find that the Fermi surface consists of three 3D...
Fig. 2 Crystal structure and electron density difference of AuTe₂Se₄/₃. a The structure of single block of atomically ordered Au–Te–Se. b The stereotype structure of the ab-plane with two kinds of Te–Te bond lengths of 3.18 and 3.28 Å. c The strips of front/back face of cubic arrays. The violet dash lines denote the Te–Te bonds of neighbor cubes along the a-axis. d The calculated EDD of front/back face, which is calculated by subtracting free atom electron density from the total electron density. The plus and minus scale represent the accumulated and depleted electrons. e The strips of middle layer of cubic arrays. The two violet and one blue dash lines denote the Te–Te bonds of neighbor cubes. f The corresponding EDD images of middle layer. g The whole crystal structure of AuTe₂Se₄/₃. The dash lines represent the triclinic unit cell (P̅1).
sheets from the I, III, and IV bands and one quasi-1D Fermi sheets from the II band. The last sheets are nearly perpendicular to the a-axis in real space, which means that the conductive electrons mainly flow along this direction. In addition, it should be emphasized that the strong nesting between quasi-parallel Fermi sheets will enhance the electron–phonon coupling, which possibly favors the SC.

**Superconductivity.** Figure 4 shows the electrical transport and magnetic properties for AuTe2Se4/3. The electrical resistivity exhibits metallic conductivity, as the plot in Fig. 4a, which is consistent with the finite DOS value at the E_F. The residual resistivity ratio (RRR) is ~20, suggesting the good quality of sample. The resistivity in low-temperature range can be well fitted using \( \rho(2) = \rho(0) + AT^2 \) equation, which indicates the sample is a typical Fermi-liquid system. The superconducting transition occurs at \( T_{\text{onset}} = 2.85 \) K. The drop of resistivity can be described by Aslamazov–Larkin25 and Maki–Thompson26,27 formulas that account for superconducting fluctuations. Our fitting results indicate that the finite scale of 2D order parameters emerge below \( T_{c0} = 2.81 \) K, see Supplementary Fig. 4. We find that temperature dependent upper critical fields obey 2D SC characters below \( T_c \). As seen from Fig. 4b and 4c, \( T_c \) monotonically decreases under in-plane (H//ab) and out-of-plane (H//c) external magnetic fields. The two \( \mu_0H_c(0) \), as shown in Supplementary Fig. 5a, can be determined by the Ginzburg–Landau (GL) expressions for 2D SC

\[
\mu_0H_c^{/c} = \frac{\Phi_0}{2\pi^2\xi (0)c^2(0)} (1 - T/T_c) \\
\mu_0H_c^{/ab} = \frac{\Phi_0\sqrt{12}}{2\pi^2\xi (0)c^2} \sqrt{1 - T/T_c}
\]

where \( \Phi_0 \) is the flux quantum, \( \xi (0)c^2 \) the GL coherence length at \( T=0 \) K, and \( d_c \) the SC thickness. Fitting the data of \( \mu_0H_c^{/c} \) against temperature yields \( \xi (0)c^2 = 28.7 \) nm. From the second equation, we can extract the value of \( d_c \). The drop of resistivity can be described by Thompson26,27 formulas that account for superconducting fluctuations. Our fitting results indicate that the finite scale of 2D order parameters emerge below \( T_{c0} = 2.81 \) K, see Supplementary Fig. 4. We find that temperature dependent upper critical fields obey 2D SC characters below \( T_c \). As seen from Fig. 4b and 4c, \( T_c \) monotonically decreases under in-plane (H//ab) and out-of-plane (H//c) external magnetic fields. The two \( \mu_0H_c(0) \), as shown in Supplementary Fig. 5a, can be determined by the Ginzburg–Landau (GL) expressions for 2D SC

\[
\mu_0H_c^{/c} = \frac{\Phi_0}{2\pi^2\xi (0)c^2(0)} (1 - T/T_c) \\
\mu_0H_c^{/ab} = \frac{\Phi_0\sqrt{12}}{2\pi^2\xi (0)c^2} \sqrt{1 - T/T_c}
\]

In comparison, the experimental data clearly deviates the curve of 3D anisotropic mass model \( H_c(\theta) = H_c^{/ab}/(\sin^2 \theta + \gamma^2 \cos^2 \theta) \); \( \gamma = H_c^{/ab}/H_c^{/c} \) around \( \theta = 90^\circ \), as shown in the upper inset of Fig. 4e, unambiguously showing the SC of AuTe2Se4/3 is quasi-2D nature.

The SC of AuTe2Se4/3 was examined by the magnetic susceptibility \( \chi \) in the low-temperature range. Figure 4f plots the temperature dependent \( \chi \) of AuTe2Se4/3 in zero-field cooling (ZFC) and field-cooling (FC) mode under 10 Oe. The diamagnetic signals are observed below 2.78 K. The superconducting volume fraction is estimated to be 90% at 1.8 K. The different magnitudes of \( \chi \) under H//ab and H//c modes indicate a strong anisotropy of AuTe2Se4/3 single crystal. The magnetization curves

---

**Fig. 3** Electronic structure of AuTe2Se4/3. **a** The XPS pattern of AuTe2Se4/3 at 300 K. The olive and orange dots and lines are the measured data and fitted curves. **b** The scalar relativistic band structure of AuTe2Se4/3. The four bands crossing the Fermi level are colored as blue, yellow, green, and red. **c** The projected density of states of AuTe2Se4/3. **d** The Fermi sheets of AuTe2Se4/3 at 3D view, side view and four separated bands in the first Brillouin zone. The G point is the zone center. The coordination a, b, and c are the real space notions.
around $T_c$ are shown in Fig. 4g, in which the type-II SC is observed. One can obtain the lower critical field $\mu_0 H_{c1}(0)$, 8.2 mT, from fitted curve of the minima of all the $M$–$H$ curves in Supplementary Fig. 5b. The penetration depth $\lambda(0)$ is estimated as 283.5 nm, which is comparable to the value in tellurides.31

2D BKT transition. It is known that the BKT transition can be detected from the slope evolutions of standard $I$–$V$ curves around $T_c$ in 2D superconductors. Figure 5a plots the $I$–$V$ curves in log–log scale near $T_c$, in which the critical $I_c$ increases as the temperature decreases and finally reaches 5 mA at 1.6 K. The $I$–$V$ curve at 3.0 K shows a typical ohmic conductivity. The exponent, $\eta$, from power law $V$–$I^n$ increases from 1 and then rapidly as temperature approaching $T_c$ from the high temperature side in Fig. 5b. The $\eta$ reaches 3 at $T_{BKT} = 2.78$ K, which is the signature of BKT transition. One thing should be noted that the fitting should be done at the low-current limit. Moreover, in a narrow temperature range just below $T_c$, it is theoretically proposed that the resistance would follow the equation $R = R_0 \exp[-b(T-T_{BKT})^{1/2}]$, where $R_0$ and $b$ are material dependent parameters. Figure 5c plots $(dn(R)/dT)^{-2/3}$ against temperature from 2.80 to 2.68 K. $T_{BKT}$ is extrapolated to be 2.69 K. The self-consistent $T_{BKT}$ could further confirm the 2D SC nature in AuTe$_2$Se$_{4/3}$. The BKT transition is believed to be related to topological elementary excitations, in which planar vortices and anti-vortices are starting to bound together without the symmetry breaking of order parameters below the $T_{BKT}$.33, 34

Discussion
It is known that each Te atom is coordinated by six Te atoms with Te–Te bonds of 2.86 Å (two) and 3.47 Å (four), which make up a series of highly distorted octahedra in hexagonal Te. Stretching the Te–Te bonds and reducing the distortion of octahedral could change the hexagonal ($P3_121$) structure into simple cubic one ($Pm-3m$) through elemental substitutions.22 As previous reports,18, 36, the Au substitution for Te could average the bond lengths and form a regular octahedron in Au$_{1-x}$Te$_x$ (0.6 < $x$ < 0.85) with simple cubic structure. This, however, is not

![Fig. 4 Transport and magnetic properties of AuTe$_2$Se$_{4/3}$](https://www.nature.com/naturecommunications/)

- **a** The electrical resistivity of AuTe$_2$Se$_{4/3}$ from 400 to 2 K. The quadratic fitting in the low temperature range exhibits the Fermi-liquid behavior.
- **b, c** Superconducting transition of AuTe$_2$Se$_{4/3}$ along $H//ab$ and $H//c$ under magnetic field. The magnetic field dependent resistivity of different angles (θ) at 2 K. The angular dependence of the upper critical field $\mu_0 H_{c2}(0)$. The upper inset shows the magnified view of the range near θ = 90°. The θ is the angle between magnetic field and the normal direction of the $ab$-plane. The lower inset shows the schematic diagram of measurement assembly. The wine and cyan solid lines are the fitted curves with 2D and 3D model as discussed in the text.
- **f** Temperature dependence of magnetic susceptibility ($\chi$) with ZFC and FC mode. The magnetization curves of AuTe$_2$Se$_{4/3}$ at different temperatures.
eneutral and the interaction of interlayer is Van der Waals force. On the other hand, inside the ab-plane, the shortest Te–Te bond (3.18 Å) is thought to be the origin of low-dimensional electronic structure, and another Te–Te bond (3.28 Å) might be the origin of the 2D morphology of AuTe2Se4/3.

In equilibrium phase AuTe2, it was reported that structural substitution or application of physical pressure37, 38. The Te bonds could enhance the DOS and hence SC ensues. The Te bond (3.18 Å) is thought to be the origin of c and quantum Griffiths phase are accompanied as well. The AuTe2Se4/3 is easily cleaved to thin crystals, offering a new playground for studying surface or edge states in the ab-plane. Furthermore, the topological transition of SC is demonstrated in AuTe2Se4/3, which would be more significant for examining the topological SC in monolayer limit. At the same time, the Tc and μHc2(0) can be remarkably enhanced through spin–orbital coupling, and a series of quantum phenomena are highly expected due to the large Rashba-type coupling12, 47 in high-Z elements Au and Te.

Methods

Synthesis. Single crystals of AuTe2Se4/3 were grown using the self-flux method. Total 2–4 g starting materials with high purity Au powder (99.99%, Sigma Aldrich), Te powder (99.999%, Sigma Aldrich), and Se powder (99.99%, Sigma Aldrich) were stoichiometrically weighted and sealed in an evacuated silica tube in high vacuum (10−6 mbar) and subsequently mounted into a muffle furnace. The furnace was heated up to 800 °C in 40 h and dwelled 10 h. Afterward, the furnace was slowly cooled down to 450 °C in 4 days and then shut down. Separated crystals from ingot are generally thin with maximum planar size 4 × 1 mm². The single crystals of AuTe2Se4/3 were ribbon shape with shining mirror-like surfaces.

Characterization. The powder X-ray diffraction (PXRD) pattern of polycrystalline AuTe2Se4/3 were measured by Panalytical X’pert diffractometer with Cu-Kα anode (λ = 1.5408 Å). The scanning electron microscopy (SEM) image of single crystal was captured from Hitachi S-4800 FE-SEM. The element mapping and composition of the sample were determined by Energy Dispersive Spectroscopy (EDS). The real composition was averaged as 10 sets of data. The high-angle annular-dark-field (HAADF) images were obtained using an ARM-200F (JEOL, Tokyo, Japan) scanning transmission electron microscope (STEM) operated at 200 kV with a Gatan Cs corrector (CEOS GmbH, Heidelberg, Germany) to cope with the probe-forming objective spherical aberration. The attainable resolution of the probe defined by the objective pre-field is 78 picometers. The focused ion beam (FIB)
method was used to cut narrow sample (width: ~50 nm and thickness 20–50 μm) for structural analysis. The valences of Au, Te, and Se were examined by X-ray photoelectron spectroscopy (XPS) using a JPS9200 analyzer. Samples were cut into long strips and followed by attaching four silver wires with silver paint. For the resistance measurement, chromium (5 μm) and silver electrodes (150 nm) were deposited first on the freshly cleaned samples by using thermal evaporation through stencil mask. The transport measurements were carried out by using ac lock-in method and performed down to 300 mK in an Oxford Cryostat equipped with Helium-3 and Helium-4 insert. The dc magnetic properties were characterized using a vibrating sample magnetometer (MPMS-3, Quantum Design).

DFT calculation. All the first-principles calculations were performed in the Cambridge Serial Total Energy Package with the plane-wave pseudopotential method. We adopted generalized gradient approximation in the form of Perdew–Burke–Ernzerhof function for exchange-correlation potential. The self-consistent field method was used with a tolerance of 5.0 × 10−7 eV atom−1. We used ultrasoft pseudo-potentials with a plane-wave energy cutoff of 360 eV. The first Brillouin zone is sampled with grid spacing of 0.04 Å−1. Fully optimization of the atomic positions and lattice parameters of compounds were carried out by the Broyden–Fletcher–Goldfarb–Shannon (BFGS) method until the remnant Hellman–Feynman forces on all components are smaller than 0.01 eV Å−1.

Data availability. The data that support the findings of this study has been deposited in the figshare repository (https://doi.org/10.6084/m9.figshare.5248489.v1). All other data are available from the corresponding authors upon reasonable request.

Received: 31 March 2017 Accepted: 7 August 2017
Published online: 11 October 2017

References
1. Novoselov, K. S. et al. Electric field effect in atomically thin carbon films. Science 306, 666–669 (2004).
2. Zhang, Y. B. et al. Experimental observation of the quantum Hall effect and Berry’s phase in graphene. Nature 438, 201–204 (2005).
3. Lee, C. G., Wei, X. D., Kysar, J. W. & Hone, J. Measurement of the elastic properties and intrinsic strength of monolayer graphene. Science 321, 385–388 (2008).
4. Mak, K. F. Lee, C. G., Hone, J., Shan, J. & Heinz, T. F. Atomically Thin MoS2: a new direct-gap semiconductor. Phys. Rev. Lett. 105, 136805 (2010).
5. Splendiani, A. et al. Emerging photoluminescence in monolayer MoS2. Nano Lett. 10, 1271–1275 (2010).
6. Saito, Y. et al. Metallic ground state in an ion-gated two-dimensional superconductor. Science 350, 409–413 (2015).
7. Xi, X. et al. Ising paring in superconducting NbSe2 atomic layers. Nat. Phys. 12, 139–143 (2016).
8. Saito, Y. et al. Superconductivity protected by spin-valley locking in ion-gated MoS2. Nat. Phys. 12, 144–149 (2016).
9. Saito, Y. et al. Highly crystalline 2D superconductors. Nat. Rev. Mater. 2, 16094 (2016).
10. Reynen, N. et al. Superconducting interfaces between insulating oxides. Science 317, 1196–1199 (2007).
11. Biscar...
temperature properties and IV curves. Q.H.Z. and L.G. measured the HAADF images and advised for the structure analysis. J.G.G. and X.L.C. wrote the manuscript based on discussion with all the authors.

Additional information
Supplementary Information accompanies this paper at doi:10.1038/s41467-017-00947-0.

Competing interests: The authors declare no competing financial interests.

Reprints and permission information is available online at http://npg.nature.com/reprintsandpermissions/

Publisher’s note: Springer Nature remains neutral with regard to jurisdictional claims in published maps and institutional affiliations.