

Heterocontact-Triggered 1H to 1T' Phase Transition in CVD-Grown Monolayer MoTe₂: Implications for Low Contact Resistance Electronic Devices

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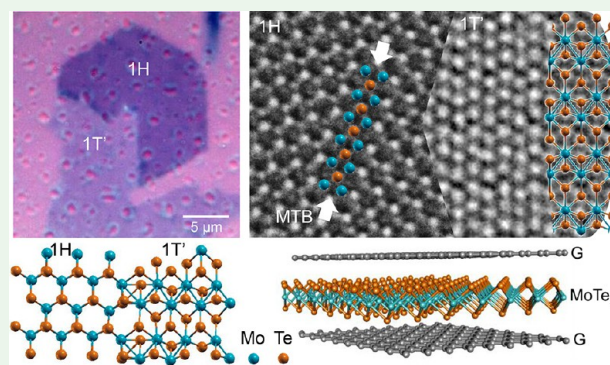
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ABSTRACT: Single-layer molybdenum ditelluride (MoTe₂) has attracted attention due to the smaller energy difference between the semiconducting (1H) and semimetallic (1T') phases with respect to other two-dimensional transition metal dichalcogenides (TMDs). Understanding the phenomenon of polymorphism between these structural phases is of great fundamental and practical importance. In this paper, we report a 1H to 1T' phase transition occurring during the chemical vapor deposition (CVD) synthesis of single-layer MoTe₂ at 730 °C. The transformation originates at the heterocontact between monoclinic and hexagonal crystals and progresses to either yield a partial or complete 1H to 1T' phase transition. Microscopic and spectroscopic analyses of the MoTe₂ crystals reveal the presence of Te vacancies and mirror twin boundaries (MTB) domains in the hexagonal phase. The experimental observations and theoretical simulations indicate that the combination of heterocontact formation and Te vacancies are relevant triggering mechanisms in the observed transformation. By advancing in the understanding and controlling of the direct synthesis of lateral 1T'/1H heterostructures, this work contributes to the development of MoTe₂-based electronic and optoelectronic devices with low contact resistance.

KEYWORDS: MoTe₂, monolayer, phase transition, heterocontact, CVD, quantum materials, HRTEM



INTRODUCTION

Molybdenum ditelluride (MoTe₂) displays structural polymorphism with numerous phases such as 2Ha, 2Hc, 2Hd, 1T', Td, and 3R existing in the bulk material and the 1H and 1T' phases stable in the monolayer limit. Because of the lower energy difference (i.e., 30–60 meV/f.u.) between the semiconducting-1H and the semimetallic-1T' phases^{1,2} with respect to other two-dimensional transition metal dichalcogenides (TMDs), monolayer MoTe₂ is an extremely attractive candidate for the development of phase change devices³ and low-resistance contacts.⁴ Additional interest in these two phases arises as 1H-MoTe₂ has a direct optical gap of 1.10 eV⁵ and strong spin–orbit coupling,⁶ while 1T' (Td) exhibits superconductivity in the monolayer limit,⁷ and is predicted to be a 2D topological and large-gap quantum spin Hall (QSH) insulator,⁸ with important implications for the development of spintronic,⁹ valleytronic, near-infrared optoelectronic, and quantum devices. While in the past most of the experiments have been carried out on mechanically exfoliated flakes,^{10–14} nowadays, it has become possible to synthesize 1T'¹⁵ and 1H¹⁵ as well as several different polymorphs of MoTe₂, i.e. 2H,¹⁵

Td,¹⁶ 3R¹⁷ and also new forms such as 2D-Mo₅Te₈¹⁸ and 1D-Mo₆Te₆ nanowires.¹⁹ However, while 1H-MoTe₂ is a well-studied material, 1T'-MoTe₂ experimental research lags behind due to the extreme air instability of this material, which rapidly degrades upon air exposure, with a lifetime in the minutes range.^{20,21}

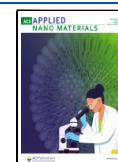
To date, significant attention has been focused on understanding and achieving controllable 1H/2H to 1T' structural transformation. Theoretically, different methods have been proposed such as electron/hole injection,^{22,3,23,24} electronic excitation,^{25–27} strain,^{28–30} annealing,^{1,22} chalcogen³¹ or metal³ atom substitution, Li²³ or H^{23,32} doping, and Te vacancies creation.^{2,23,27} Phase transition (PT) has been achieved on few-layers or bulk MoTe₂ via annealing,^{33,10} ion

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liquid gating,³⁴ electric field in vertical RRAM devices,¹¹ Te vacancies creation,³⁵ and Li intercalation.³⁶ In the monolayer limit, investigations on 1H/1T' phase transition are highly complicated by the air instability of monolayer 1T', although promising initial results have been reported by adopting annealing,¹⁰ ionic liquid gating,^{34,37} THz laser irradiation³⁸ and Te vacancies creation.³⁹ While developing approaches for triggering 1H–1T' phase transition with external stimuli remains of undoubted interest for the realization of phase change devices, progressing in understanding and controlling the direct synthesis of lateral 1T'/1H heterostructures is the base for the development of low resistance contacts for electronics and optoelectronics and for devising novel device architectures. Nowadays, growth of 1H/1T' lateral heterostructures has been demonstrated only in UHV conditions via Mo and Te evaporation on conductive HOPG¹⁸ or graphene substrates.⁴⁰ The 2H–1T' phase transition critical temperature, a relevant aspect in the controlled synthesis of the two phases, is still an open question. For bulk MoTe₂ crystals, the pioneering work of Velinga et al.³³ reported a critical temperature of 850 °C, while Keum et al.⁴¹ demonstrated critical temperatures between 500 and 700 °C, with the latter result being debated in the literature.⁴² Remarkably, measuring the transition temperature is even more challenging for monolayer MoTe₂ films due to the low material stability, as Te vacancies and 1D-Mo₆Te₆ chains formation were observed already at 200 °C.¹² Recently, the transition temperature of monolayer MoTe₂ encapsulated with hexagonal boron nitride (hBN) has been measured to be 1075 °C.¹⁰

In this paper, we demonstrate that monolayer 1H/1T' MoTe₂ lateral heterostructures can be obtained via chemical vapor deposition (CVD) directly on SiO₂/Si, thanks to a heterocontact-triggered structural phase transition taking place at 730 °C during the growth process. The resulting 1H/1T' crystals are stabilized with a scalable encapsulation approach recently reported by our group²⁰ and their structural and chemical properties are investigated by Raman spectroscopy, high-resolution transmission electron microscopy (HRTEM) and X-ray photoemission spectroscopy (XPS). Experimental data indicate that the phase transition is triggered independently from the contact angle and reveal the presence of Te vacancies and mirror twin boundary (MTB) domains in the hexagonal phase. Ab initio calculations support the observed phenomenology by indicating that defects such as Te-vacancies favor phase transition, while the presence of MTB and excess Mo are possibly responsible for its termination. The final 1H/1T' contact fronts can have a lateral size in the range of tens of micrometers, which is relevant for device fabrication such as field effect transistors (FETs) and photodetectors. The results presented are of use to devise and engineer tailored 1H/1T' lateral heterostructures for the development of novel phase-change, spintronic, and quantum devices with reduced contact resistance compared to fabricated metallic contacts.

EXPERIMENTAL SECTION

Growth of Monolayer MoTe₂. The MoTe₂ samples were grown via liquid precursor CVD, as we previously reported in Pace et al.²⁰ In this method, the molybdenum precursor is obtained from an aqueous solution and directly spun on the SiO₂ substrate. First, three mother solutions were prepared, namely, solutions A, B, and C. Solution A was obtained by dissolving 0.11 g of ammonium heptamolybdate (AHM, Sigma-Aldrich), weighted by analytical balance ABJ 80-4NM, in 40 mL of DI water. Solution B was obtained by dissolving 0.1 g of NaOH (Sigma-Aldrich) in 40 mL of DI water. Solution C consists of

OptiPrep (Sigma-Aldrich) used as purchased. The growth solution was obtained by mixing the mother solutions with volume ratio A:B:C = 0.8:0.5:0.3. The growth solution was then spin-coated on a clean SiO₂/Si substrate at 300 rpm for 10 s, plus 20 s at 3000 rpm. Before spin-coating, all SiO₂ substrates were cleaned via sonicated cleaning in acetone and isopropanol for 5 min and oxygen plasma (power = 100 W, process pressure = 80 mTorr) for 5 min to increase the hydrophilicity of the SiO₂ surface. The spin-coated substrate and metallic tellurium were then loaded in a Lenton hot-wall horizontal CVD reactor (see Figure S1 for furnace schematic). The growth was carried out at near-atmospheric pressure for 2 to 15 min, under a constant flow of Ar/H₂ (H₂ 3%) gas at 100 sccm. Growth temperature, molybdenum concentration, and cooling process were optimized to increase the density of spontaneously contacting hexagonal and elongated MoTe₂ flakes as well as material thickness uniformity. Best results were obtained at a growth temperature of 730 °C, with increased molybdenum precursor concentration, and by opening the CVD reactor at 700 °C. After the growth, the reactor was allowed to cool rapidly under constant flux of Ar by opening the lid at 700 °C. We note that in our process the 1H to 1T' transformation is of a rapid “explosive” nature; i.e., the percentage of the transformed area of hexagonal flakes undergoing phase transition is independent of the growth time, which can be reduced down to 2 min (see Figure S2). In our experiments, for consistency, we adopt a growth time of 15 min. We report that growth time reduction below 2 min leads to the synthesis of incomplete crystals.

Semidry Encapsulation. Semidry top-encapsulation of as-grown MoTe₂ was obtained via the delamination of CVD hBN or graphene films grown on copper foil. Nominally, single-layer hBN (15 × 15 cm²) from Graphene Supermarket was used for encapsulation. Graphene samples were grown in-house using a 4" Aixtron BM Pro CVD reactor.⁴³ The hBN (graphene) was cut into squares with dimensions larger than the target SiO₂ sample. Then, it was covered with a double-layer polymeric membrane of PMMA AR-P 679.02 (Allresist) and 15% PPC in anisole (Sigma-Aldrich) by spin coating for 60 s at 2000 rpm and baked at 90 °C for 2 min. A PDMS frame with few millimeters of thickness was placed on the top of the sample as a supporting frame. The sample was then floated in a NaOH electrolyte solution (1 M), where an electrochemical reaction takes place: here the hBN (graphene) on copper foil acts as anode and a platinum foil is used as cathode. A constant 2.42 V voltage is applied to the Pt electrode until the PPC/PMMA/hBN(graphene) is completely delaminated from copper foil. Then the PPC/PMMA/hBN(graphene) was allowed to float in DI water for 3 min to remove the NaOH residuals. The detached PPC/PMMA/Graphene membrane was then laminated on top of the as-grown sample and heated at 90 °C. Finally, a double cleaning in acetone and isopropanol was used to remove the supporting polymer.

MoTe₂ Transfer. Graphene or hBN top-encapsulated MoTe₂ was transferred using HF (Sigma-Aldrich) as SiO₂ etchant, as previously reported in.²⁰ In particular, the sample (i.e., encapsulated MoTe₂ on SiO₂/Si) was covered with a double-layer of 679.02 PMMA (Allresist), which was sequentially spun (60 s, 2000 rpm) and baked (90 °C, 2 min). Then, the sample was left floating in concentrated hydrofluoric acid (HF 48%, Sigma-Aldrich) for few seconds until the complete detachment of the membrane. After detachment, the floating membrane was fished and rinsed in DI water for a few seconds to minimize the exposure of MoTe₂ to water. Finally, the membrane was rapidly fished with the target substrate (silicon for XPEEM measurements and 100 nm SiO₂/Si for TEM sample preparation) and backed at 90 °C for a few minutes. The polymeric membrane was then removed via a standard cleaning in acetone. Indeed, often the formation of a parasitic Te-like phase was observed in the monolayer 1T' regions upon the transfer process,^{44,45} hence in this work we limited our PEEM/XPS analysis to the 1H and 2H regions.

Graphene top-encapsulated MoTe₂ transferred on a 100 nm SiO₂ substrate was used for TEM sample preparation. Optical microscopy was used to identify suitable MoTe₂ monolayers. Such flakes were then transferred to a Quantifoil R 1.2/1.3 TEM grid with the help of a

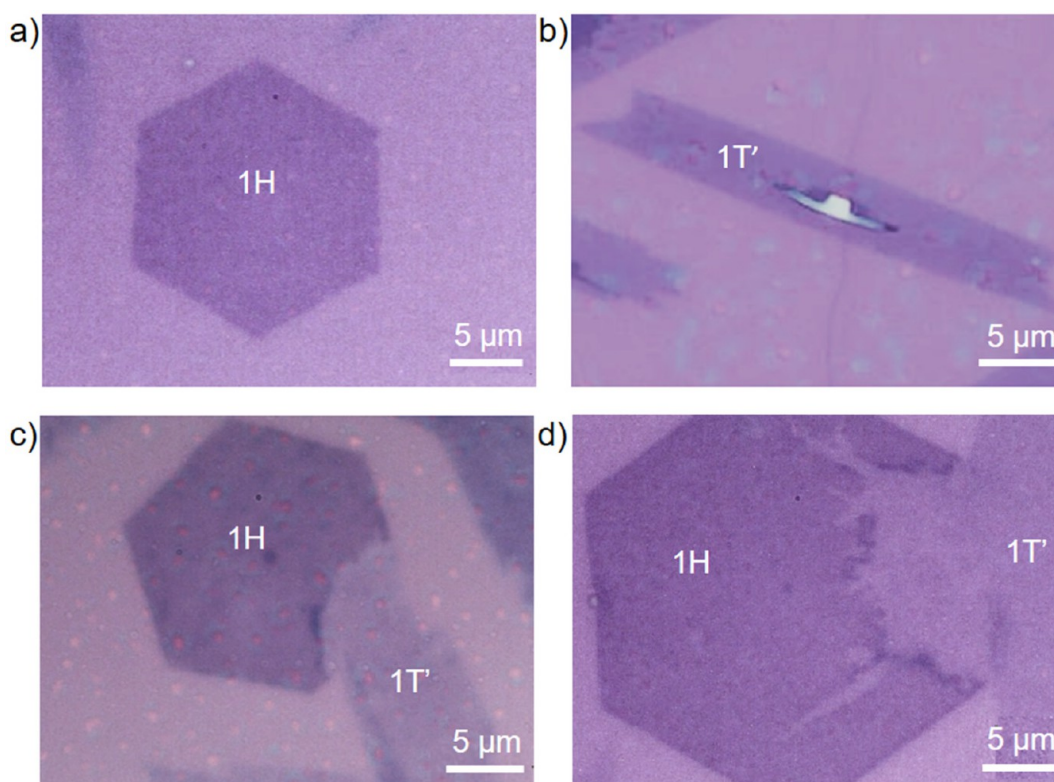


Figure 1. Optical images of pristine hexagonal 1H single crystal (a), pristine monoclinic 1T' single crystal (b), heterocontact between 1H and 1T' single crystals with uniform (c), and nonuniform 1H to 1T' transformation (d).

drop of isopropyl alcohol (IPA) to bring the grid into contact with the flake. Evaporation of the IPA attached the grid to the flake. With potassium hydroxide (KOH), the underlying SiO₂ substrate was etched away, releasing the grid with the MoTe₂ flake. Afterward, the residues of the preparation are removed with double-distilled water.

Characterization Techniques. The optical images of the samples were obtained with a ZEISS AxioScope 7. Amplitude modulated Kelvin probe force microscopy (AM-KPFM) was performed with a Bruker Dimension Icon microscope equipped with an FMV-PT probe without applying bias to the tip. Raman characterization was carried out with a Renishaw InVia spectrometer. The laser wavelength used was 532 nm with a fwhm of the Gaussian beam \sim 1600 or 800 nm for 50 \times or 100 \times lenses, respectively, in long exposure time mode (60/180 s) and low power (1.16/0.58 mW) and in short exposure time (1–2 s) and high power (5.8 mW) mode. Polarized Raman measurements were implemented in $z(x,-)z$ configuration using a $\lambda/2$ waveplate for the incoming light.

X-ray photoemission electron microscopy (XPEEM) measurements were performed by using the ELMITEC-LEEM III instrument at the MAXPEEM beamline of the MAX-Lab synchrotron radiation facility in Lund, Sweden. The measured photocurrent is 20 mA at 100 eV and 70 mA for 350 eV. The work function of the analyzer is 4.8 eV. High-resolution (HR)TEM images were acquired with the C_c/C_s -corrected “Sub-Ångström Low-Voltage Electron microscope” (SALVE)⁴⁶ at an acceleration voltage of 80 kV. Measured values for the chromatic aberration C_c and spherical aberration C_s were in the range of $-10 \mu\text{m}$ to $-20 \mu\text{m}$. The vacuum pressure in the column of the TEM was on the order of 1×10^{-5} Pa.

Scanning transmission electron microscopy (STEM) and energy-dispersive X-ray spectroscopy (EDX) were carried out to determine the local elemental composition. For the STEM-EDX measurements, a Thermo Fisher Talos 200X (S)TEM was operated at 80 kV. The system is equipped with a SuperX EDX detector for spectroscopy and elemental mapping.

Computational Methods. Density Functional Theory (DFT) calculations were performed in the open-source Quantum ESPRESSO

(QE) package.⁴⁷ The exchange-correlation functional was described by the modified Perdew–Burke–Ernzerhof (PBEsol) version of the generalized gradient approximation (GGA) with scalar-relativistic Optimized Norm-Conserving Vanderbilt and Ultrasoft pseudopotentials used for Mo and Te atoms, respectively.^{48–50} The kinetic energy cutoff for wave functions was set to 45 Ry. We used Marzari–Vanderbilt–Devita–Payne cold smearing approach for electronic occupations with 0.1 eV.⁵¹ A $12 \times 7 \times 1$ Monkhorst–Pack wavevector grid was used for both rectangular 1H and 1T' cells with an 18.729 Å vacuum thickness for both 1H and 1T' phases. Vc-relaxed lattice parameters were calculated to be $a = 3.505 \text{ \AA}$, $b = 6.07 \text{ \AA}$ for 1H and $a = 3.398 \text{ \AA}$, $b = 6.307 \text{ \AA}$ for 1T' pristine cells containing 6 atoms (2 Mo + 4 Te). We used $3 \times 2 \times 1$, $7 \times 1 \times 1$, and $1 \times 7 \times 1$ supercells with $4 \times 4 \times 1$, $2 \times 7 \times 1$, and $12 \times 1 \times 1$ k -space grids, respectively, to simulate Mo diffusion and heterocontact-induced phase transitions in monolayer MoTe₂. For the orthogonal $b_{1T'}/2a_{1H}$ heterocontact case, we used a combination of $3 \times 1 \times 1$ and $2 \times 2.5 \times 1$ supercells of 1T' and 1H phases, respectively, with $2 \times 7 \times 1$ k -space grid. To minimize the periodic effect of the replicas on the studied heterointerface, the atoms of the first and last “subcells” of the supercells were kept fixed in the positions obtained from the pristine 1H and 1T' relaxed supercells at equilibrium cell size. In the case of the ZZ supercell, 1 Te atom from the supercell edge was removed because of the small Te–Te distance at the supercell boundary. Te-deficient heterocontact cells were relaxed using “epitaxial_bc/ac” method to avoid additional out of plane deflections. The climbing image nudged elastic band method (CI-NEB) was performed to calculate kinetic barriers between the states. Phase diagrams calculations were performed using linear elasticity theory.⁵² Elastic energy difference was calculated using elastic constants C_{ij} calculated by applying deformations reported in the work.⁵³

RESULTS AND DISCUSSION

In Figure 1a,b, we show the optical micrograph of typical crystals of MoTe₂ grown on SiO₂/Si via liquid precursor CVD.

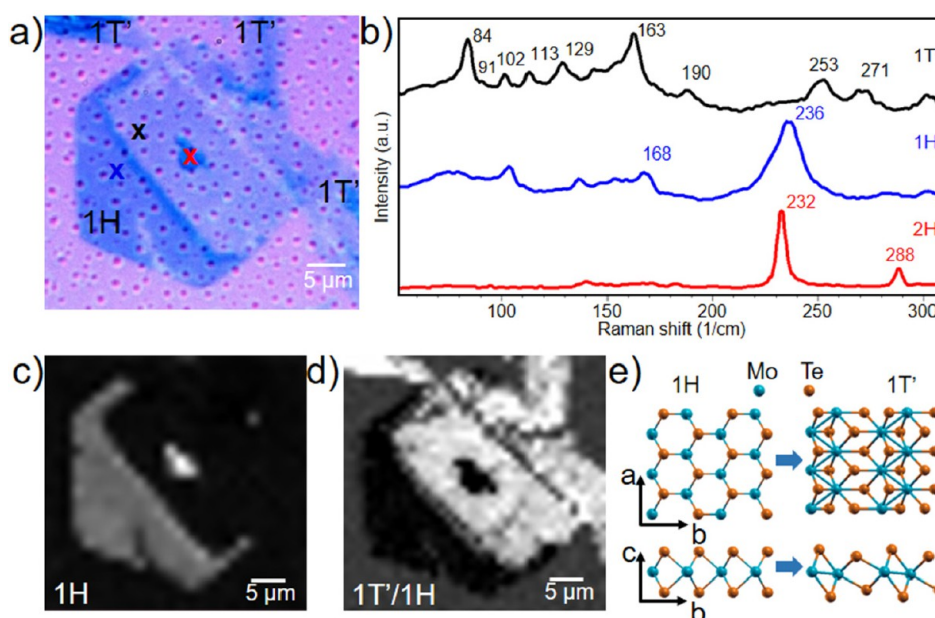


Figure 2. (a) Optical image of a flake presenting a heterocontact-triggered phase transition. (b) Raman spectra recorded on the blue, red, and black crosses in part a. (c) Map of the intensity of the 1H 236 cm^{-1} E_{2g}^1 Raman peak. (d) Map of the intensity ratio of the Raman 84 cm^{-1} A_g^1 ($1T'$) and 236 cm^{-1} E_{2g}^1 (1H) peaks. (e) Top and side views of 1H and $1T'$ - MoTe_2 ball-and-stick models; blue (orange) colors indicate the Mo (Te) atoms. The well-visible crystal break in Figure 2a,d running from side to side of the hexagonal flake originated during hBN encapsulation, and it does not influence the crystallographic orientation of the $1T'$ transformed hexagonal crystal.

Crystals of different shape are visible within the same sample: hexagonal (a) and elongated (b) ones, which are normally attributed to the 1H and $1T'$ phase, respectively. In fact, 1H- MoTe_2 is stable in a hexagonal phase (space group $P\bar{6}m2^{22}$) and hence grows with hexagonal symmetry, while $1T'$ - MoTe_2 is stable in a monoclinic structure (space group $P2_1/M^{22}$) and grows in an elongated shape. We optimized the growth parameters to maximize the density of MoTe_2 flakes in each sample, which in turn led to the observation of a significant number of heterocontacts between hexagonal and elongated flakes (like those shown in Figures 1c,d and 2a). The MoTe_2 samples were encapsulated immediately after growth with monolayer CVD hBN (or graphene) to increase the lifetime of $1T'$ - MoTe_2 and allow further characterization, similar to what reported in.²⁰

In Figure 2a, one can observe three elongated flakes contacting a hexagonal crystal, which interestingly presents different optical contrasts. A precise assignment of the structural phase of the material, carried out by Raman spectroscopy indicates that only part of the hexagonal flake displays a 1H phase, with the remnants being $1T'$ (see Figure 2b–d). Indeed, Figure 2b reports the typical Raman spectra measured in different regions (marked with crosses) of panel a. The blue and black spectra were recorded on the regions with dark and light contrast, respectively, while the red spectrum was recorded at the center of the hexagonal crystal. The blue spectrum shows typical features of a monolayer 1H phase, such as the in-plane E_{2g}^1 mode at 236 cm^{-1} and the out-of-plane A_{1g} mode at 168 cm^{-1} ,⁵⁴ while few-layer peaks E_{1g} at 116 cm^{-1} and B_{2g}^1 at 288 cm^{-1} are absent.^{13,55} The portion of the hexagon with darker contrast in the left side of panel a is hence assigned to 1H- MoTe_2 . The triangular feature with darker contrast placed at the center of the hexagon presents a Raman spectrum that is indicative of few-layer 2H- MoTe_2 , as typically observed in seeding areas. The portion of the hexagon with lighter

contrast presents a Raman spectrum with six A_g peaks located at $84, 113, 129, 163, 253,$ and 271 cm^{-1} and three B_g peaks positioned at $91, 102,$ and 190 cm^{-1} , which are indicative of monolayer $1T'$.⁵⁶ Indeed, the spatial distribution of the two different phases is very visible with spatially resolved Raman mapping. Panels c and d report Raman maps of the intensity of the 236 cm^{-1} E_{2g}^1 peak and of the intensity ratio of the 84 cm^{-1} A_g^1 and 236 cm^{-1} E_{2g}^1 peaks for a straightforward visualization of the 1H and $1T'$ phases, respectively (ball-and-stick models of the two phases are sketched in panel e). The presence of a $1T'$ region within the hexagonal crystal is indicative of a phase transition taking place during the growth process itself, which is performed at a maximum temperature of $730\text{ }^\circ\text{C}$. As also supported by the data presented in the rest of the paper, the transformation is triggered by the presence of a direct contact between a 1H and a $1T'$ crystal. Indeed, we confirm this phenomenology in many samples (>100), where hexagonally shaped flakes contacted by elongated ones present partial or total transformation to $1T'$ (see Figure S3). The observed PT mechanism is enticing as it leads: (i) to the generation of large lateral 1H/ $1T'$ heterostructures (i.e., tens of micrometers, significantly larger than those typically found between randomly touching flakes); (ii) to full transformation of large $1T'$ - MoTe_2 hexagonal crystals.

To better understand the crystal orientation of the contacting and transformed $1T'$ flakes, we adopted a combination of high-resolution transmission electron microscopy (HR-TEM) and polarized Raman spectroscopy measurements. Indeed, polarized Raman spectroscopy has been intensively used in the study of bulk,^{57–59} few-layer^{59,60} and monolayer⁵⁶ MoTe_2 optical and crystallographic properties, as the A_g^5 (253 cm^{-1}) and A_g^6 (271 cm^{-1}) modes allow defining the $1T'$ crystal direction, if excited with linearly polarized light.⁶¹ The maximum value of the intensity ratio $I(A_g^5)/I(A_g^6)$ —which we refer to as R -value in the following text—

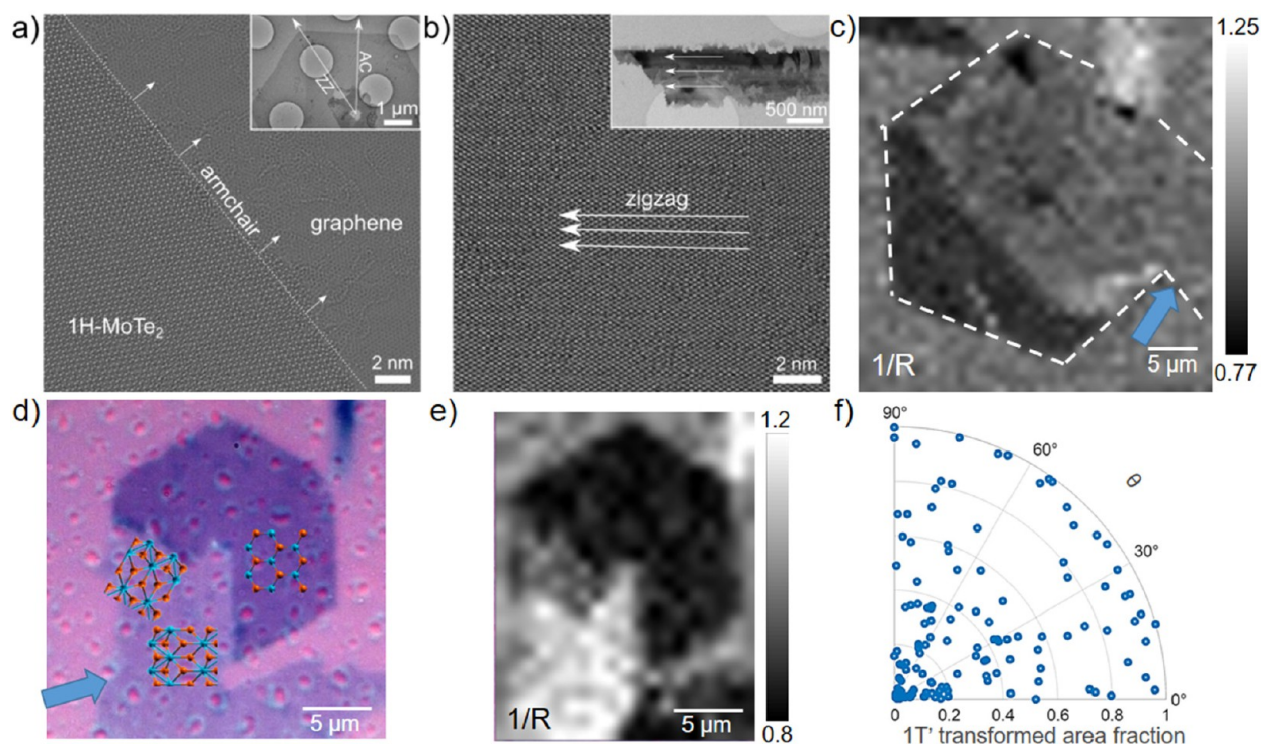


Figure 3. (a) 80 kV C_c/C_s -corrected HRTEM image of an untransformed 1H flake edge with identified AC direction. In the inset: TEM overview image of the 1H hexagon. (b) 80 kV C_c/C_s -corrected HRTEM image of a 1T' flake with identified ZZ direction. Inset: TEM overview image of the monoclinic 1T' crystal. (c) 1/R Raman mapping of the flake studied in Figure 1. (d) Optical image of a partially transformed flake in the case of collinear contact with ball-and-stick models of the identified crystal orientations. (e) 1/R Raman mapping of the flake in panel d. (f) Collected statistical data of the percentage of hexagonal flakes transformed into 1T' phase with respect to the heterocontact angle (θ), i.e., the angle between 1H and 1T' ZZ directions.

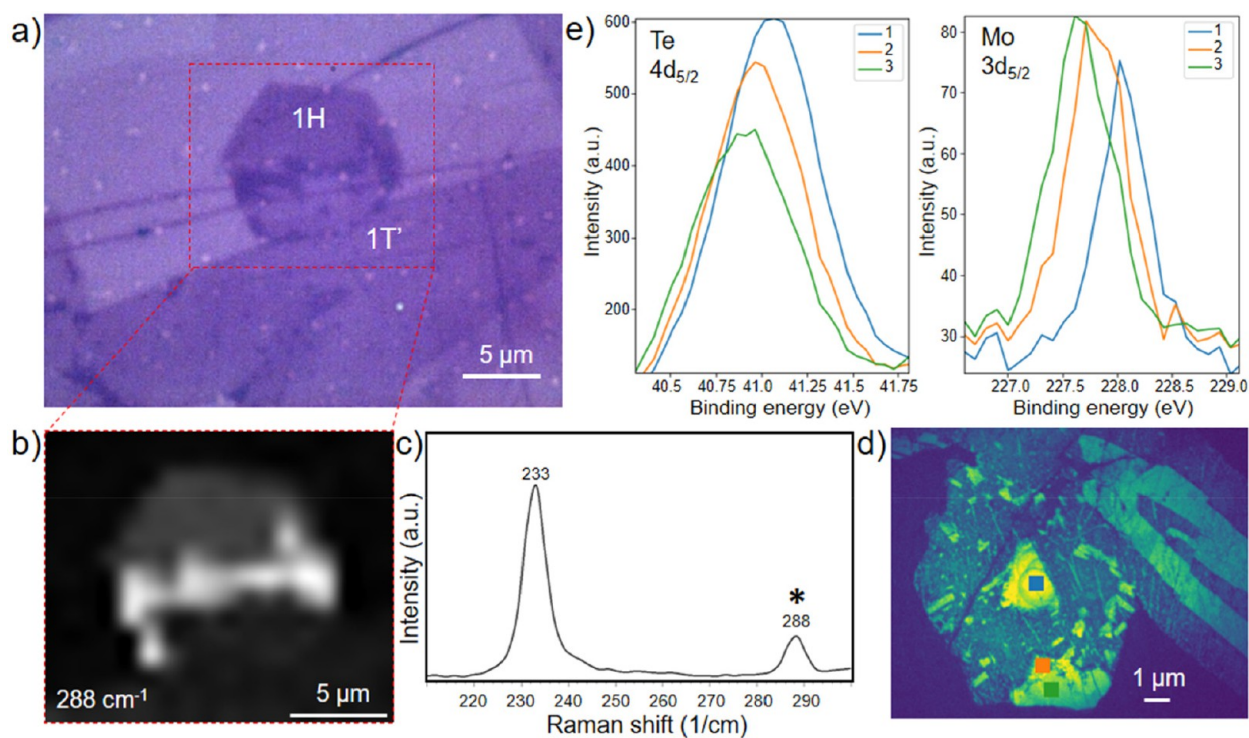


Figure 4. (a) Optical image and (b) B_{2g}^1 mode Raman mapping of a partially converted MoTe_2 crystal. (c) Raman single spectrum obtained from the heterocontact region of the flake in parts a and b confirming the presence of few-layers 2H. (d) Te 4d μ XPEEM image and (e) relative Te $4d_{5/2}$ and Mo $3d_{5/2}$ intensity profiles.

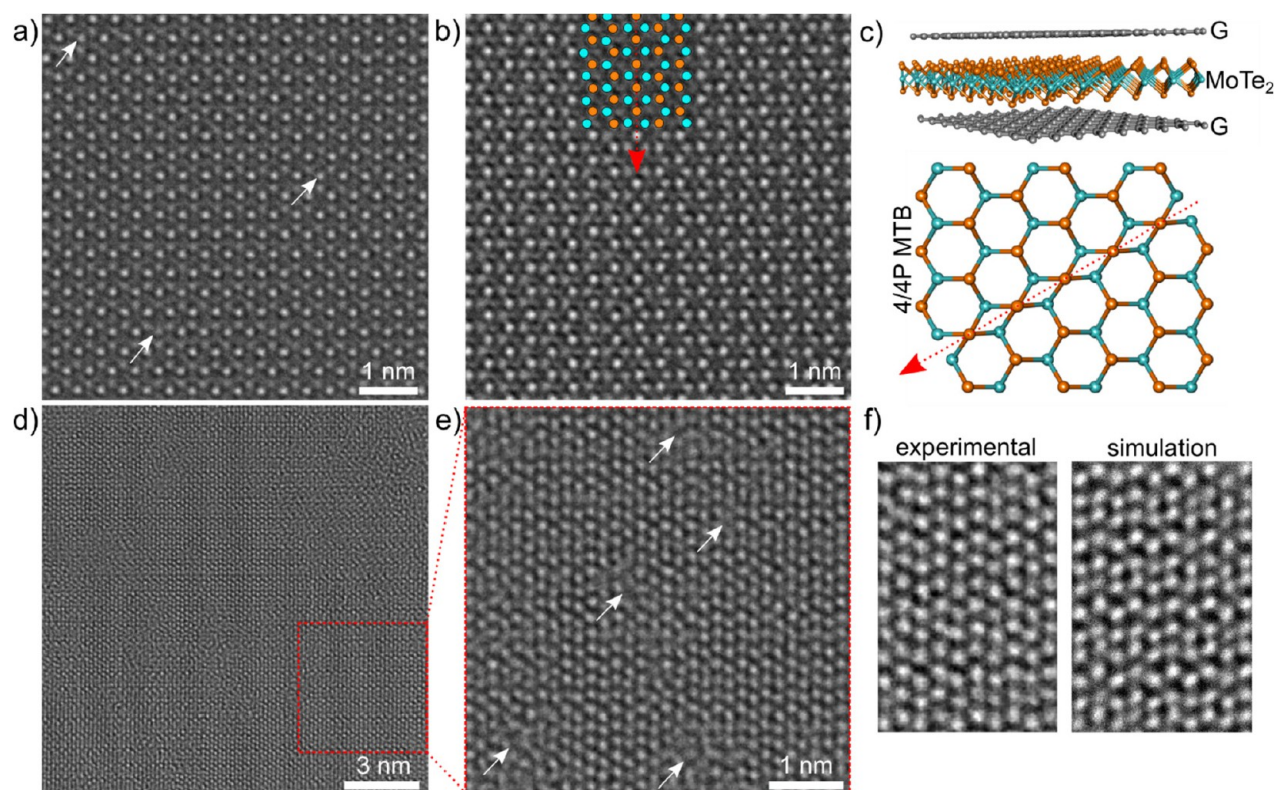


Figure 5. 80 kV C_c/C_s -corrected HRTEM analysis of monolayer 1H and 1T' MoTe₂. Images of (a) Te single vacancies and (b) 4/4P MTB in the nontransformed 1H area. (c) Schematic of the graphene-encapsulated MoTe₂ and ball-and-stick model of the 4/4P MTB in part b. Blue (orange) balls correspond to the Mo (Te) atoms. (d) Atomically resolved image of monolayer 1T' crystal showing the presence of defects. (e) Magnified region from part d. Defective regions are marked with white arrows. (f) Experimental and simulated HRTEM images of graphene-encapsulated monolayer 1T' phase (see Figures S11 and S12 for corresponding selected area electron diffraction patterns and additional image simulations, respectively).

corresponds to the $[0 \pm \pi]$ angle between the incident light polarization and the crystal's zigzag (ZZ) direction.^{56,61} HRTEM allowed us to identify the edges of our untransformed hexagonal 1H crystals to be armchair (AC) terminated, while 1T' single crystals were found to elongate along the ZZ-direction (see Figure 3a,b). Via polarized Raman, we measured a maximum R -value in 1T' flakes in horizontal configuration (i.e., long side/ZZ direction parallel to the light polarization), with the minimum measured for the orthogonal alignment configuration (see Figure S4). For the flakes displayed in Figure 2a, the spatial distribution of the R -value allowed us to determine that the dominating crystallographic direction of the transformed hexagonal crystal corresponds to that of the lower contacting 1T' flake (see Figure 3c and Figure S5). This finding suggests that the 1H–1T' phase transition was originally triggered by the flake indicated in Figure 3c by the light blue arrow.

We verified with a large number of flakes and found that in all instances the transformed 1T' crystals have a direction that is either collinear or 60° rotated with respect to that of the original 1T' contacting flake. Figure 3 reports the case of transformation with collinear orientations of initial 1H and 1T' crystals (i.e., ZZ-directions originally parallel in both crystals), where we see the formation of a 60° rotated 1T' domain and the incompleteness of the transition process. We argue that such polycrystallinity observed in some of the flakes (see also Figures S6 and S7) is a consequence of the 1H 3-fold rotational symmetry that might lead to three different 1T' crystal orientations rotated by 60°.⁶²

Also, we note that not all contacts between elongated 1T' and hexagonal 1H crystals result in structural phase transitions, while in some instances a limited local recrystallization near the contact points is observed. To better understand the cause of the heterocontact-induced phase transformation phenomenon, we have investigated whether there is any preferential contact angle between the 1T' flake and the hexagonal crystal that leads to the transition and whether this can be related to the percentage of the transformed flake. By inspecting 99 heterocontacts, we can conclude that there is no such indication: neither the occurrence of the phase transition nor the percentage of the transformed 1H flake is obviously related to the initial contact angle (see Figure 3f).

The optical micrographs in Figures 2a and 3d present darker contrast at the 1H–1T' interface, compatible with few-layer MoTe₂. This is also visible in Figure 4a, where we show an optical image of a multiphase MoTe₂ flake, where Raman analysis of the optically darker region reveals few-layer 2H features. Indeed, the B_{2g}^1 mode at 288 cm^{-1} is observed (panel b) while the E_{2g}^1 mode shifts from 236 to 233 cm^{-1} (panel c).¹³ Such few-layers regions observed at the 1H–1T' interface suggest the presence of excess molybdenum in those areas during the CVD process, which transforms into few-layer MoTe₂ as tellurium is fluxed in the tube. Similar evidence of excess Mo diffusion during the CVD process is found in several multiphase MoTe₂ hexagonal flakes presenting elongated lobes at the edge of 1T' transformed portions (see Figure S8). While the stoichiometry during the growth process cannot be verified but only indirectly inferred, more information on the chemical

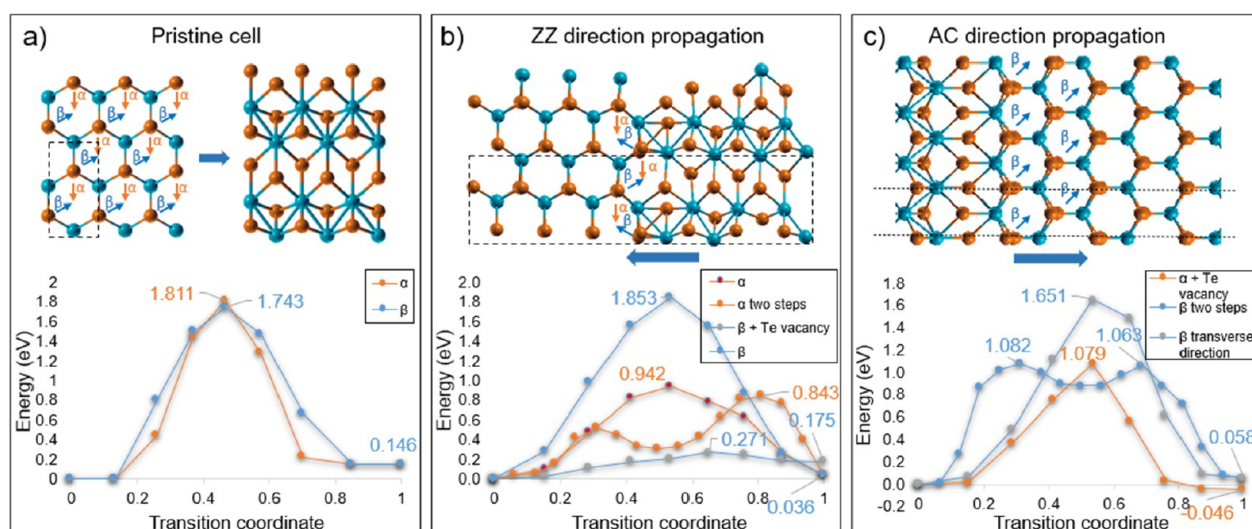


Figure 6. Phase transition energy barriers calculated for pristine case (a) and heterocontact cases with propagation in ZZ (b) and AC (c) directions. Unit cells are indicated by black dashed lines. The total number of atoms used is 6, 41, and 42 for pristine cell, ZZ supercell and AC supercells, respectively.

composition of the grown flakes can be extracted via X-ray photoemission spectroscopy (XPS) measurements, which were performed at the X-ray photoemission electron microscopy (XPEEM) beamline of the MAX-Lab synchrotron. To this end, graphene-encapsulated MoTe_2 samples transferred on silicon were analyzed. In Figure 4e, we show the Te 4d and Mo 3d core levels measured on the areas indicated by colored squares in the Te 4d XPEEM map (panel d). We analytically extracted the Mo:Te ratios for three different regions: 2H seed (blue square), $1\text{T}'$ -1H heterocontact boundary (orange square), and 1H region (green square). The ideal Mo:Te stoichiometry of the pristine MoTe_2 is equal to 0.5. We obtain a value of 0.43 ± 0.04 for the 2H seed area, 0.51 ± 0.04 for heterocontact area, and 0.57 ± 0.05 for the monolayer 1H region (see Supporting Information for more information). These values and the Te $4d_{5/2}$ intensity variation suggest the presence of Te vacancies in the untransformed 1H area.

High-resolution transmission electron microscopy (HRTEM) was performed to validate the crystallinity of the flakes on the atomic scale and confirm the presence and nature of defects. To minimize beam damage in the monolayers, MoTe_2 was encapsulated with graphene. Monolayer $1\text{T}'$ showed a rather defective crystalline structure (Figure 5d,e) and the heterocontact region could not be imaged (Figure S10d-f), due to damage arising during transfer to the grid because of the high sensitivity of the $1\text{T}'$ phase to oxygen.^{63,64} Conversely, in the 1H region, two types of defects were identified: (i) single Te vacancies (see Figure 5a); (ii) inversion domains, mainly $4\text{I}4\text{P}$ Mirror Twin Boundary (MTB) domains (Figures 5b,c and S10). These MTBs have already been observed in experiments with Mo excess forming “triangular” and “wagon-wheel” shaped domains,⁶⁵ and in the newly reported hexagonal $1\text{H}-\text{Mo}_5\text{Te}_8$ phase,¹⁸ ultimately confirming the Te deficiency already highlighted by XPEEM measurements.

Our experimental data indicate that heterocontact-triggered polymorphism takes place: (i) independent from the $1\text{T}'/1\text{H}$ contact angle; (ii) in the presence of Te vacancies and MTB within the 1H original crystal; (iii) during CVD growth and while Mo is diffusing.⁶⁶ In the following text, we adopt linear

elasticity theory⁵² and ab initio calculations to model and further understand the observed phenomenology.

Before entering into this discussion, we consider the effect of the strain, originating from the $1\text{H}-1\text{T}'$ lattice constant mismatch, on the polymorphism reported in this work. Our density-functional theory (DFT) simulations indicate 1.19% compression along ZZ and 1.56% stretching along AC as typical strain values for a $1\text{H}/1\text{T}'$ HC in the collinear case. The calculated strain values correspond to small variations in the energy difference between the two phases of the order of few meV/f.u. compared to the calculated 60 meV (30 meV/f.u.) ground state energy difference between relaxed 1H and $1\text{T}'$ cells (see Figure S13). This indicates that strain is not likely to play a determinant role in the transformation. This is also confirmed by experimental observations: if the strain was the dominant mechanism triggering the PT, we would expect to measure a maximum of the $1\text{H}-1\text{T}'$ transition probability density as a function of the angle, in correspondence to 60° periodicity, where the calculated energy difference between the two phases is minimum (see Supporting Information for discussion and Figure S13). Considering additional 0.66% biaxial strain from the thermal expansion,⁶⁷ this angle corresponds to the collinear ZZ directions of $1\text{T}'$ and parent 1H crystals, in agreement with previous theoretical predictions for this system.²⁹ In contrast, our experimental data do not show a univocal angle dependence; rather, a uniform probability density for the occurrence of the phase transition is observed (see Figure 3f). Thus, we conclude that the strain effect is not determinant in triggering the phase transition.

With the help of first-principles calculations, we now consider the effects of various structural mechanisms on the energetic landscape of single layer MoTe_2 , specifically aiming at assessing the role of the heterocontact and identifying possible structural mechanisms entailing a reduction of the kinetic barrier between 1H and $1\text{T}'$ phases, thus favoring the phase transition. In the presence of high energy barrier, the 1H to $1\text{T}'$ transition process might not take place or be too lengthy to be observed. Hence, we performed ab initio simulations of heterocontact-triggered transition paths for the collinear contact (i.e., $1\text{H}-\text{ZZ}/1\text{T}'-\text{ZZ}$ and $1\text{H}-\text{AC}/1\text{T}'-$

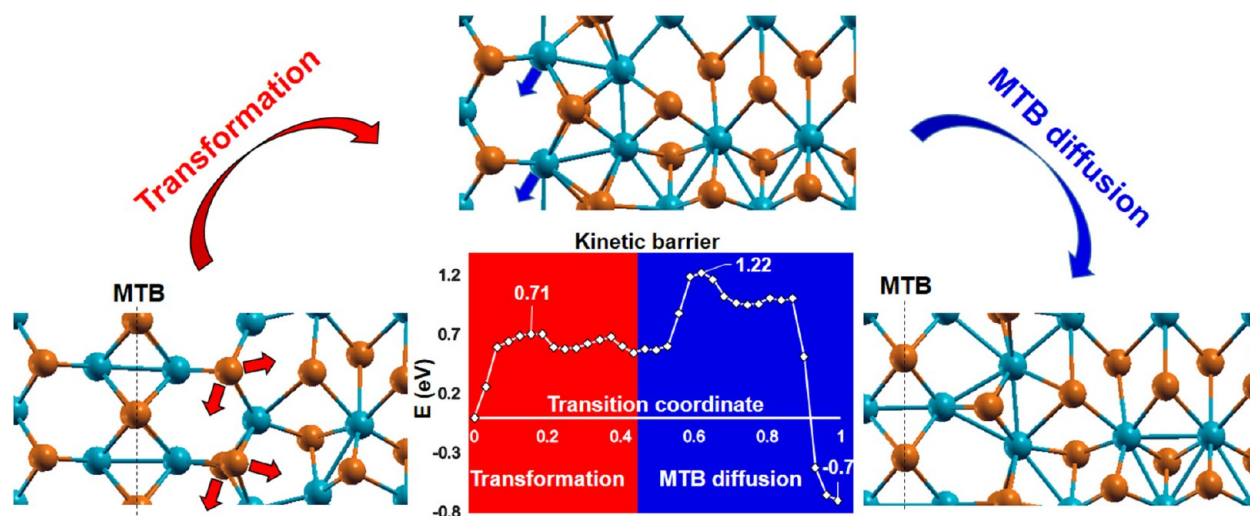


Figure 7. Transformation-diffusion model of the orthogonal $b_{1T'}/2a_{1H}$ interface heterocontact. The total number of atoms is 48.

AC) reported in Figure 3d, considering transition paths involving Te atom displacements. First, we calculated energy barriers for pristine cases, which we refer to as α - and β -paths in Figure 6a. Both paths result in the same final $1T'$ configuration but originate from the different lower Te atom displacement directions in the initial $1H$ cell. For the α -path, Te atoms displacement is parallel to the final $1T'$ -AC direction, while for β it is 45° tilted. We calculated 1.81 (905 meV/f.u.) and 1.74 (870 meV/f.u.) eV energy barriers for α - and β -paths, respectively, with final $1T'$ energy 0.146 eV (73 meV/f.u.) higher than initial $1H$ phase due to the fixed (constant) cell condition. These results are in agreement with previously reported values for pristine cells.^{22,30} Next, we performed simulations for direct $1H$ - $1T'$ contact in AC and ZZ direction considering supercells of 7 pristine unit cells (see Figure 6b,c). We refer to the same α and β notations corresponding to the lower Te atoms displacement. We consider broken periodicity in x and y directions that also results in possible two steps transformations (see Figures S14 and S15) and additional β -path in AC direction corresponding to the transverse direction propagation (see Figure S15) similar to reported in.²⁸ Both ZZ and AC routes show similar behavior of α -ZZ, two step α -ZZ and β -AC paths with 0.94 eV (470 meV/f.u.), 0.84 eV (420 meV/f.u.), and 1.08 eV (540 meV/f.u.) energy barriers, respectively. The energy differences between the two phases are 0.036 (18 meV/f.u.) and 0.056 (28 meV/f.u.) eV for the ZZ and AC cases, respectively. We then extended the analysis to defective paths, considering Te vacancy presence at the heterointerface (see Figures S14 and S15). In general, paths that involve Te vacancies are the most promising: in the ZZ case, the presence of a Te vacancy is found to strongly suppress the kinetic barrier between the two phases from 0.94/0.84 eV (470/420 meV/f.u.) in the pristine case to 0.27 eV (135 meV/f.u.). However, the $1T'$ phase becomes more unfavorable +0.170 eV (+85 meV/f.u. with respect to the $1H$ phase. Conversely, in the AC case the energy barrier is not suppressed but the $1T'$ becomes energetically favorable at -0.046 eV (-23 meV/f.u.) with respect to the reference $1H$ phase. These findings are not surprising and consistent with the well-studied Te vacancy induced phase transition mechanism.^{27,39} Our simulations suggest that Te vacancies work differently in AC and ZZ directions and are not the sole responsible for the observed PT; however, they

importantly modify the energetic landscape and can be involved in the stabilization of the $1T'$ phase in combination with other mechanisms. Reported reduced energy barrier values and lowered ground state energy difference between $1H$ and $1T'$ phases indeed can be responsible for the avalanche effect observed experimentally. However, in the ZZ case, Te vacancies, even with a reduced barrier value, lead to an ascending steps energy profile upon increase of the transformed unit cells number and should result in the back-propagation of $1T'$ to $1H$ transformation of the triggering $1T'$ crystal, which we never observe experimentally. In addition, Te vacancies (in both the AC and ZZ directions) tend to accumulate upon phase transition propagation and at some point terminate the transformation (see Figure S14). Therefore, additional mechanisms for $1T'$ phase stabilization should be considered such as temperature, higher strain due to the interaction with SiO_2 substrate or Na and O adatoms⁶⁸ from the initial Na_2MoO_4 precursor⁶⁶ (see Supporting Information for discussion).

Our DFT simulations also confirm that a heterocontact-triggered phase transition is possible for the orthogonal $1T'$ -AC/ $1H$ -ZZ contact, indicated by the arrow in Figure 3c. In that case, we would expect a huge lattice mismatch up to 10% at the $b_{1T'}/2a_{1H}$ interface with approximately 5% $1H$ -ZZ compressive strain taking place. Raman spectroscopy measurements reveal that the R values of both contacting $1T'$ flake and phase transformed flakes are equivalent (or 60° rotated), and hence during this transformation the hexagon's recrystallization of the $1H$ -AC into the $1T'$ -ZZ direction takes place. The possible mechanism of this polymorphism can be explained with a transformation-diffusion model (Figure 7). The initial lowest energy configuration of the heterocontact contains an MTB line and the following Te atom displacement results in the formation of an unstable $1T'$ cell with 0.71 eV (355 meV/f.u.) step-like energy barrier, which would make this transformation unfavorable. However, subsequent Mo atoms diffusion (which was shown experimentally at 250°C ¹² and normally takes place during CVD growth⁶⁶) out of the heterocontact area stabilizes the $1T'$ phase with 0.7 eV (350 meV/f.u.) energy gain and final 1.22 eV (610 meV/f.u.) energy barrier.

Finally, two possibilities are represented by other kinds of defects, namely, MTBs and Mo diffusion, since we have

observed MTBs and the consequences of Mo diffusion in our grown crystals (e.g., material accumulation and 1T' post-transformation growth continuation). Thus, we performed additional DFT simulations to understand how these defects affect the transformation. We find that MTBs and Mo excess in ZZ direction generally lead to energetically unfavorable kinetic paths in all the studied situations (see Figures S16 and S17, respectively). More interesting is the case of excess Mo for phase transition propagation in the AC direction. We find intermediate states with double Mo chains formation, resembling the structures reported in monolayer M_4X_6 TMDs.⁶⁹ These structures are 0.7–0.77 eV (350–385 meV/f.u.) more favorable than the following 1H–1T' cell transformation (see Figure S18). In the end, our calculations show a 0.54 eV diffusion barrier in 1H phase via substitutional mechanism in agreement with,⁶⁵ and 0.66 eV in 1T' phase along the ZZ direction via interstitial diffusion mechanism (see Figure S19). These barriers are significantly lower than those reported for most transition routes. Therefore, we do not exclude that transition kinetics could be limited by Mo diffusion within both the 1H and 1T' domains and can explain why we observe uncompleted PT for different 1H–1T' heterocontact angles. Considering few-layer $MoTe_2$ formation at the 1H–1T' heterointerface and 1T' post-transformation growth continuation, we suppose that the diffusion of excess Mo during CVD growth plays an important role and 1H to 1T' transformation can be stopped at high Mo concentration or in the presence of defects hindering Mo diffusion.

CONCLUSIONS

In conclusion, we report a semiconductor to semimetal phase transition realized in monolayer $MoTe_2$ during CVD growth and triggered by the presence of a direct contact between 1H and 1T' single crystals. The temperature observed for the phase transition is 730 °C, lower than the temperature reported in¹⁰ for the annealing of an 1H- $MoTe_2$ crystal. We systematically studied the heterocontact-triggered transformation and found that its occurrence and size are independent of the original contact angle and can yield sizable 1H/1T' lateral heterostructures. The crystal orientation of the transformed 1T' domains was investigated using linearly polarized Raman spectroscopy and revealed the possibility of two preferential crystal transformation routes: collinear and orthogonal to the initial 1H crystal, both resulting in domains with 60° periodicity. Chemical and structural analyses were performed via XPS and TEM on encapsulated samples to limit their degradation and indicated the presence of MTB and Te vacancies in the 1H crystals. We theoretically modeled transition pathways considering both stoichiometric and nonstoichiometric cases and identified transformation routes with a wide variety of kinetic barrier values. Our simulations demonstrate a significant reduction in the kinetic energy barrier in the heterocontact case compared to the pristine case, which can explain the observed avalanche effect in transformation propagation. Ab initio calculations support the observed phenomenology by indicating that defects such as Te-vacancies favor phase transition, while MTB presence or excess Mo prevent it. For the orthogonal heterocontact case, we propose a transformation–diffusion model involving MTBs to describe the observed polymorphism. The pathways studied in this work can be further investigated in other TMDs, their heterojunctions and in Janus materials.⁵² A recent work has demonstrated that few-layer 2H/1T' $MoTe_2$ heterocontacts

can be fabricated through photolithographic techniques, an approach that has been used to recover damaged 2H- $MoTe_2$.⁷⁰ While material postprocessing offers enticing prospects for heterocontact formation, in monolayer $MoTe_2$ this strategy might be difficult to implement due to the extreme material instability. Defining pathways for bottom-up synthesis of low contact resistance heterojunctions is indeed extremely appealing for fabrication of 2D electronic and optoelectronic devices such as FETs and photodetectors (see Supporting Information for a discussion) and for the development of novel spintronic and quantum devices. By demonstrating heterocontact-induced phase transition and contributing to unveiling the mechanism behind it via theoretical modeling, our work makes a step forward toward the identification of novel strategies for the development of $MoTe_2$ -based technology.

ASSOCIATED CONTENT

Supporting Information

The Supporting Information is available free of charge at <https://pubs.acs.org/doi/10.1021/acsnm.3c01314>.

Discussion on the 1T' crystal orientation analysis and determination of the formed 1T' domains inside 1H hexagons, example of the post-transformation 1T' growth, description of the Mo:Te ratio extraction approach from the μ XPS data, additional KPFM, HRTEM, SAED, and HAADF characterization data, discussion on the transition temperature and chemical stabilization, discussion on the 1H–1T' heterostructures application, theoretical strain analysis by means of DFT and linear strain theory, details on the DFT simulation of the phase transition propagation for pristine and defective cases and Mo diffusion simulations in both 1H and 1T' phases of Mo (PDF)

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Notes

The authors declare no competing financial interest.

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