Atomic-Scale \textit{in Situ} Observations of Crystallization and Restructuring Processes in Two-Dimensional MoS$_2$ Films

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**ABSTRACT:** We employ atomically resolved and element-specific scanning transmission electron microscopy (STEM) to visualize \textit{in situ} and at the atomic scale the crystallization and restructuring processes of two-dimensional (2D) molybdenum disulfide (MoS$_2$) films. To this end, we deposit a model heterostructure of thin amorphous MoS$_2$ films onto freestanding graphene membranes used as high-resolution STEM supports. Notably, during STEM imaging the energy input from the scanning electron beam leads to beam-induced crystallization and restructuring of the amorphous MoS$_2$ into crystalline MoS$_2$ domains, thereby emulating widely used elevated temperature MoS$_2$ synthesis and processing conditions. We thereby directly observe nucleation, growth, crystallization, and restructuring events in the evolving MoS$_2$ films \textit{in situ} and at the atomic scale. Our observations suggest that during MoS$_2$ processing, various MoS$_2$ polymorphs co-evolve in parallel and that these can dynamically transform into each other. We further highlight transitions from in-plane to out-of-plane crystallization of MoS$_2$ layers, give indication of Mo and S diffusion species, and suggest that, in our system and depending on conditions, MoS$_2$ crystallization can be influenced by a weak MoS$_2$/graphene support epitaxy. Our atomic-scale \textit{in situ} approach thereby visualizes multiple fundamental processes that underlie the varied MoS$_2$ morphologies observed in previous \textit{ex situ} growth and processing work. Our work introduces a general approach to \textit{in situ} visualize at the atomic scale the growth and restructuring mechanisms of 2D transition-metal dichalcogenides and other 2D materials.

**KEYWORDS:** MoS$_2$, graphene, aberration-corrected scanning transmission electron microscopy, \textit{in situ}, physical vapor deposition, crystallization, two-dimensional heterostructures
films crystallize and restructure to nanocrystalline MoS2 (nc-MoS2) domains and thereby explore in situ and at the atomic scale the richness of MoS2’s structural evolution via multiple polymorphs.

The importance of MoS2 stems from the current interest to use this material as a device active layer in low-dimensional (opto-)electronics as well as a potent catalyst in (photo-)electrochemical energy applications, such as the hydrogen evolution reaction (HER).

All of these application fields share the key prerequisite of scalable synthesis of MoS2 with controlled properties. The desired structural characteristics of MoS2 in electronic and catalytic applications vary however drastically: For electronics, semiconducting MoS2 with precisely controlled layer number, large crystals, and a low defect density is desired in order to achieve, for example, high current on/off ratios and high carrier mobilities in field effect transistor (FET) MoS2 devices. In stark contrast, for (electro)catalytic applications such as HER typically finely nanostructured or even amorphous MoS2 with good electrical conductivity, a large specific surface area and a large number of pronounced defects and edge sites are desired, since these imperfections rather than a highly crystalline basal plane are considered as electrocatalytically active sites.

Important in this context, MoS2 occurs in multiple polymorphs: First on the monolayer level, the arrangement of the three covalently bonded atomic sublayers (S−Mo−S) within a MoS2 monolayer can principally show trigonal prismatic (commonly termed “2H monolayer”, also often referred to as “1H monolayer”) or octahedral (termed “1T monolayer”) symmetries. Importantly, the more commonly found 2H monolayers are semiconducting, while the comparatively metastable 1T monolayers are metallic, implying a key influence of MoS2 monolayer symmetry on the material’s application profile. Second, when individual monolayers of a given symmetry type are stacked upon each other by van der Waals interactions, multiple stacking arrangements are possible, which in turn impact on optoelectronic properties. For instance, for the 2H monolayer type, several stacking arrangements are possible, where the most commonly occurring equilibrium types are 2H (AA’ stacking) and 3R (ABC stacking). Important beyond the equilibrium 2H and 3R stacking, also more complex nonequilibrium stacking sequences including homonuclear stacking (e.g., AA) have been reported. With increasing layer number, the possible complexity of these layer arrangements generally increases, since different stacking types can also co-exist within

Figure 1. (a) HAADF STEM image series of ~2 nm MoS2 on graphene during continuous e-beam exposure (time stamps indicated) which leads to e-beam-induced crystallization from a-MoS2 to nc-MoS2 domains. (b) FTs of selected frames in (a) with corresponding time stamps indicated. (c) Schematic illustration summarizing the observations deduced from (a).
multilayers, let alone given the further additionally complex possibility of different monolayer types (2H/1T) stacking onto each other. This polymorphism of MoS$_2$ in both monolayer type and multilayer stacking opens a complex parameter space of possible layered MoS$_2$ structures which require control in any synthesis route.

The key methods to realize MoS$_2$ layers are CVD (including solid metal/vapor sulfurization methods), PVD (e.g. sputter deposition or evaporation), and wet chemical synthesis. Important in this context is that, unlike other key 2D materials (such as graphene or hexagonal boron nitride), MoS$_2$ does not require a metallic process catalyst to grow and crystallize. Also MoS$_2$ growth can be achieved already at comparatively low temperatures ($\sim$400 °C). Based on this comparatively facile crystallization of MoS$_2$, CVD-type synthesis is most promising for electronic-grade MoS$_2$ while PVD and wet chemical synthesis offer a high degree of control over nanostructured electrocatalytically active MoS$_2$. While the various MoS$_2$ synthesis techniques comprise completely different formation environments, precursors, constituent species fluxes, and significantly different growth kinetics, all techniques nevertheless routinely employ elevated temperature treatments ($\sim$400 °C) at some stage during growth or postprocessing in order to stabilize a certain MoS$_2$ structure. Therefore, in all synthesis routes of MoS$_2$ the structural mechanisms proceeding at elevated temperatures such as nucleation, sustained growth, crystallization, and restructuring are of fundamental importance. Yet, at present little work has been done to elucidate these mechanisms. In particular, in situ observations of growth, crystallization, and restructuring processes in MoS$_2$ and other TMDCs at the atomic scale are critically missing. This results in a limited understanding of the fundamental mechanisms underlying synthesis and processing, thereby hindering rational synthesis and postgrowth process development for MoS$_2$.

Our here presented approach for atomically resolved in situ STEM imaging of MoS$_2$ crystallization and restructuring therefore contributes to such much needed understanding by directly identifying various mechanistic growth and restructuring steps: In particular we observe, depending on the initial thickness of a-MoS$_2$ deposited, in-plane crystallization toward few-layer nc-MoS$_2$ with layers parallel to the support for thin a-MoS$_2$ films, while comparatively thicker a-MoS$_2$ films evolve into a two-segment nc-MoS$_2$ film morphology with interfacial in-plane MoS$_2$ layer crystallization parallel to the support and with perpendicular MoS$_2$ layering farther away from the support. Our data reveal that during this crystallization and restructuring various MoS$_2$ polymorphs co-evolve in parallel. Importantly, we find that these polymorphs dynamically transform into each other during processing, driven via diffusion of Mo and S species and, depending on conditions, influenced by a weak MoS$_2$/graphene heterostructure support epitaxy. Contextualizing these in situ observations with recent ex situ MoS$_2$ growth and processing literature, our work visualizes in situ and at the atomic scale the multiple fundamental structural processes occurring in parallel under widely used ex situ MoS$_2$ processing conditions. Notably, our insights are based on a model system and a STEM environment that is readily extendable to in situ studies of other TMDCs and 2D materials.

RESULTS AND DISCUSSION

We fabricate a-MoS$_2$ samples for high-resolution STEM by sputter deposition of ultrathin PVD MoS$_2$ films directly onto free-standing monolayer CVD graphene membranes. The graphene membranes, which act as an ultrathin and light support for STEM, were suspended across the holes of a holey carbon support foil of a TEM grid by a polymer-free transfer process which ensures an as clean as possible MoS$_2$-graphene interface. During PVD of MoS$_2$ onto the graphene-covered TEM grids, the samples were not intentionally heated leading to deposition of a-MoS$_2$. MoS$_2$ film with nominal thicknesses ranging from $\sim$2 nm to $\sim$10 nm were deposited. For further details on experimental methods, see the Methods section.

Figure 1a shows a high-angle annular dark-field (HAADF) STEM image series (60 kV electron acceleration voltage) of a nominally $\sim$2 nm-thick MoS$_2$ film on a graphene monolayer during its structural evolution as a function of continuous e-beam scanning time (time stamps indicated in Figure 1; sample was not intentionally heated during STEM imaging). As apparent from the initial image at 0 min, the $\sim$2 nm MoS$_2$ in its as-deposited state does not homogeneously cover the graphene support but shows a morphology of interconnected islands (bright regions in Figure 1) with bare graphene areas in between (dark regions in Figure 1). The STEM image at 0 min also gives the visual impression of an amorphous structure in the MoS$_2$ deposit. This is corroborated by the Fourier transform (FT) data in Figure 1b for 0 min, which only shows a broad halo indicative of amorphicity. Upon continued scanning of the e-beam over the field of view of Figure 1a, the appearance of the MoS$_2$ islands gradually changes: The visual impression suggests gradual island restructuring resulting in (i) crystallization of the amorphous MoS$_2$ islands toward nc-MoS$_2$ with the MoS$_2$ layers parallel to the graphene support as well as (ii) slight "dewetting" of the MoS$_2$ from its support. Corroborating a-MoS$_2$ crystallization, after 8 min e-beam exposure not only the visual appearance in the STEM data (Figure 1a) but also the FT data in Figure 1b suggest some e-beam-induced crystallization of the MoS$_2$ as a more pronounced ring in the FT pattern has emerged. The emerging ring corresponds well to the 2H MoS$_2$ (010) reflection ($\sim$0.26 nm), consistent with 2H MoS$_2$ crystallization with layers parallel to the support. After 17 min of continuous e-beam scanning, the FT in Figure 1b has even developed signs of one discrete hexagonal spot pattern (indicated by white arrows). This suggests an emerging dominant crystalline 2H MoS$_2$ layer orientation across the entire field of view in Figure 1a (17 min). Interestingly, we find that the e-beam-induced crystallization is a phenomenon highly localized to the e-beam in STEM with a sharp boundary between exposed crystallized and nonexposed amorphous material (Figure 2). We note that such a good spatial definition of the beam-driven crystallization implies that e-beam exposure could potentially be used to spatially selectively transform a-MoS$_2$ to nc-MoS$_2$ in a fabrication scenario. Given the lower chemical stability of a-MoS$_2$ compared to nc-MoS$_2$, this may be useful for direct resist-free patterning of crystalline MoS$_2$ devices where nonexposed a-MoS$_2$ could be chemically etched away with the more stable crystalline MoS$_2$ remaining.

To complement our STEM measurements, additional time-resolved bright-field (BF) transmission electron microscopy (TEM) and selected area electron diffraction (SAED)
measurements at electron acceleration voltages from 60 kV to 200 kV are presented in Supporting Information Figures S1–S3. Figure S1 (80 kV) corroborates at a wider field of view in the TEM (up to 900 nm) the e-beam-induced restructuring and crystallization of our ∼2 nm a-MoS2 films to nc-MoS2 with 2H MoS2 layers parallel to the graphene support, consistent with our STEM data. Figure S2 (60 kV, same electron acceleration voltage as used in STEM) shows similar e-beam-induced crystallization at 60 kV and reveals via time-dependent energy dispersive X-ray spectroscopy (EDX) measurements that the S/Mo ratio in the films during their e-beam-induced transition from a-MoS2 to nc-MoS2 only slightly drops from S/Mo_{0 min} = 2.1 ± 0.03 to S/Mo_{18 min} = 2.0 ± 0.03. This suggests that the loss of S via e-beam-induced sputtering processes in our MoS2-graphene heterostructures in particular at 60 kV can remain limited, consistent with previous literature.44,45 Figure S3 (80 kV vs 200 kV) finally confirms that a-MoS2 crystallization is also observed for 200 kV electron acceleration, whereby we find that the rate of a-MoS2 crystallization for 80 kV and 200 kV appears roughly similar, while in contrast the degradation rate of the graphene support is much more pronounced at 200 kV due to much increased electron knock-on damage to the graphene.46 Overall and most importantly, our TEM data in Figures S1–S3 confirm that the observed e-beam-induced a-MoS2 crystallization is a generic processes independent of our particular employed microscope type (i.e., STEM or TEM); note that one STEM and two different TEM systems were found to give consistent results, see Methods section) and is working over a wide range of typical (S)TEM electron acceleration voltages and imaging parameters, making our model heterostructures an easily implemented in situ imaging platform.

Figure 1c schematically illustrates our observations of this e-beam-induced crystallization and restructuring of initial a-MoS2 clusters to nc-MoS2 of a few layers thickness with MoS2 planes parallel to its graphene support. Atomic-scale in situ work on MoS2 has to date primarily concentrated on the formation of defects in and amorphization of initially fully crystalline MoS2 monolayers,44,47–50 that is, the reverse process of the a-MoS2 crystallization observed here and on phase transitions (e.g., 2H to 1T) in fully crystalline MoS2.13 In contrast, crystallization of a-MoS2, as followed here at the atomic scale, has previously been studied only at comparatively large fields of view, insufficient to discern details on the single atom level, be it in or ex situ from thermal activation11,37–40,51,52 or e-beam irradiation.39,52–54 In contrast to previous work, our high-resolution STEM data now allow us to discuss atomic-scale details of the crystallization and restructuring processes based on direct in situ information.

To quantify the HAADF STEM intensity data from Figure 1a, we show in Figure 3a the central region from Figure 1a at higher magnification after 18 min e-beam exposure. Taking a HAADF intensity line profile (Figure 3b) along the yellow line indicated in Figure 3a allows to identify the nature of the atoms in the image based on the element-specific intensity of HAADF data of ultrathin films which has a dependence55 on atomic number Z of Z^{−1.64}. We thereby identify the thinnest region in Figure 3a (across which the line profile is drawn) to be a MoS2 monolayer of 2H monolayer structure12,13 (Figure 3b inset). Consistently this region displays a 6-fold FT (inset of Figure 3a) with distances of ∼0.26 nm and ∼0.15 nm corresponding to the (010) and (110) reflections of 2H MoS2.
Figure 4. False colored recalculation of the *in situ* crystallization time series in Figure 1 (time stamps indicated) for which the HAADF intensity has been normalized to the intensity of a single S atom.55 The labeled spots (a) to (g) point to salient structural features and evolutions discussed in the main text.

Figure 5. (a,b) HAADF STEM image series of other locations from a ∼2 nm MoS2 on graphene during continuous e-beam exposure (relative time stamps indicated). The corresponding HAADF STEM *in situ* videos taken during the continuous e-beam exposure (temporal resolution ∼2.7 s per frame) for (a) and (b) are given in Video S1 and Video S2, respectively (time lapsed to 4 frames per second, time stamps indicated for salient frames in the videos).
respectively. Based on this identification of a 2H monolayer MoS\(_2\) region, we recalculate the HAADF intensity counts in Figure 3a to a relative intensity with respect to the HAADF intensity from a single S atom as shown in the false color coded image in Figure 3c, in which a single S atom (\(Z_s = 16\)) has relative intensity 1 and a single Mo atom (\(Z_{Mo} = 42\)) has a relative intensity of \(\sim 4.9\). In doing so we establish a straightforward way of identifying the structure of further MoS\(_2\) regions in our in situ STEM data. For increasing layer numbers, the spatial average intensity over a region scales approximately linearly with number of layers. The atomic stacking type in such multilayers can then in turn be discerned by further analyzing the spatially resolved intensities as a function of atomic positions. For instance, the region left of the monolayer patch in Figure 3c is thereby consistent with a 2H bilayer, as indicated in the image. Furthermore, in Figure 3c several isolated Mo atoms can be identified on the graphene support (see labeled examples) as well as one Mo adatom on the 2H bilayer patch (correspondingly labeled).

Following this method, we present in Figure 4 the recalculated data from the time series in Figure 1 and identify via the spots (a) to (g) labeled in Figure 4 three salient structural evolution processes that we find to occur in parallel in this image series. Additional HAADF STEM time series data in Figure 5 for two other regions on a \(\sim 2\) nm MoS\(_2\) on graphene sample show a matching evolution. Importantly, for Figure 5a,b we also provide the corresponding HAADF STEM data. For increasing layer numbers in the temporal resolution \(\sim 2.7\) s per frame as Video S1 and Video S2, respectively.

In Figure 4 spots (a) and (b) we follow the structural evolution that we most commonly observe upon e-beam-induced crystallization: An initially amorphous region crystallizes into bilayer patches of 2H MoS\(_2\).\(^{30}\) In particular, for spot (a) we observe after 3 min nucleation of a 2H bilayer patch in the upper right. This region has expanded after 8 min, whereupon at 14 min, two more nonconnected 2H bilayer regions appeared in the lower left and lower right of spot (a). From 8 to 15 min, these regions restructure, including some intermittent shrinking, and before 17 min, the 2H bilayer patches have expanded into one connected single crystalline grain. This grain in spot (a) at 17 min covers \(\sim 5.7\) nm\(^2\), which is the largest connected single crystalline grain imaged in Figures 1 and 4. A similar evolution is also found in Figure 5 and Video S1 and Video S2: In Figure 5a/Video S1, a small 2H bilayer nucleus near the center of the image grows in lateral size at the expense of surrounding amorphous MoS\(_2\) deposits on the graphene. In Figure 5b/Video S2, a 2H monolayer is observed in the center of the frame with an adjacent largely amorphous bilayer region to the upper right. Upon continued e-beam exposure, this amorphous bilayer region crystallizes into a larger 2H bilayer grain.

This generally observed preferential formation of the 2H phase from a-MoS\(_2\) confirms previous formation energy calculations of various MoS\(_2\) bulk polymorphs that predicted 2H to be the energetically most favored structure.\(^{11,18}\) Since in our ultrathin MoS\(_2\)-graphene heterostructures thermodynamic bulk properties may be modified by effects from heterogeneous interfaces and free surfaces etc., we model in Figure S4 nonbulk representations of heterostructures of a-MoS\(_2\) on graphene in comparison to a crystalline 2H MoS\(_2\) bilayer patch on graphene and calculate their formation energies using density functional theory (DFT). From our calculations, we find the crystalline 2H bilayer MoS\(_2\) patch on graphene to be between \(\sim 0.26\) eV/atom and \(\sim 0.34\) eV/atom lower in formation energy than the corresponding a-MoS\(_2\) on graphene. This suggests (in general agreement with previous bulk calculations)\(^{11,18}\) also for our ultrathin MoS\(_2\)-graphene heterostructures that a thermodynamic driving force is behind the experimentally observed crystallization of a-MoS\(_2\) to 2H MoS\(_2\), whereby we hypothesize that the energy input from the scanning e-beam is helping to overcome kinetic barriers\(^{27}\) to crystallization.

In contrast to this theoretically predicted evolution of our a-MoS\(_2\) toward 2H, we however find in Figure 4 in spot (c) in the lower right at 17 min a crystalline MoS\(_2\) region to have evolved from initial a-MoS\(_2\) that has a spatial average intensity consistent with bilayer, but where the intensities as a function of atomic positions indicate that this bilayer patch is not of the 2H type. Instead the measured atomically resolved intensity profile of spot (c) at 17 min is consistent with a bilayer that shows homonuclear stacking (either 2H\(^{\prime}\) or 1H), where Mo atoms of the second layer are placed directly above Mo atoms of the first layer. This observation of homonuclear stacking next to 2H stacking suggests that, besides crystallization toward equilibrium 2H, initial a-MoS\(_2\) can also crystallize into other MoS\(_2\) polymorphs under fixed processing conditions, thereby resulting in co-existence of several MoS\(_2\) polymorphs. While homonuclear stacking is energetically not favored,\(^{18}\) it has been previously observed \(ex situ\) in annealed liquid-phase exfoliated MoS\(_2\) layers,\(^{17}\) where similar to our observation here, equilibrium 2H bilayers and nonequilibrium homonuclearly stacked bilayer regions co-existed. We note that this resemblance between our \(in situ\) and previous \(ex situ\) data implies that our atomic-scale \(in situ\) observations are indeed capturing processes which are relevant to \(ex situ\) MoS\(_2\) processing.

Besides predominant 2H stacking and homonuclear stacking, we find after extended e-beam exposure (17 min) also a third salient stacking type shown in Figure 4 at the spots labeled (d). Compared to the 2H and homonuclear bilayers, this region exhibits no six-fold symmetry but a line appearance of different symmetry. Measuring characteristic distances for spots (d) in Figure 4, we find this structure to exhibit a spacing of \(\sim 0.23\) nm which is comparably shrunk from the typical \(\sim 0.26\) nm distance in 2H MoS\(_2\). This structure is thereby reminiscent of merging line defects in MoS\(_2\) layers that result from loss of S under continued e-beam illumination.\(^{50}\) Such a \(\sim 0.23\) nm fringe spacing is also approaching the spacings expected for metallic Mo phases,\(^{58}\) and the observed line-like symmetry is also evocative of previously reported S-deficient MoS\(_{2-x}\) phases.\(^{59}\) For these reasons we tentatively assign the structure at spots (d) in Figure 4 to locally S-deficient MoS\(_{2-x}\), which is created during our continued e-beam exposure in the STEM by S loss from the initially present MoS\(_2\). This S loss leads to crystallization/restructuring not toward a MoS\(_2\) polymorph but a S-deficient structure, akin to recent results on e-beam-induced S-deficient phase formation in SnS\(_2\).\(^{60}\) We note that controlled \(ex situ\) formation of such S-deficient MoS\(_{2-x}\) has previously been suggested to be beneficial for certain applications requiring 2D Mo–S compounds with increased reactivity.\(^{61}\) We also note however that our EDX measurements in the TEM in Figure S2, discussed above, as well as the observation in Figure 4 that globally the 2H MoS\(_2\) phase is the predominant phase suggests that on a larger scale,
the loss of S is limited at 60 kV for our in situ crystallization conditions.

Having established the three salient Mo–S structures in our data, we note that the observed MoS$_2$ crystallization and restructuring processes under the e-beam are found to be highly dynamic: Notably, in spot (e) in Figure 4 (corresponding also to the region shown in Figure 3), the small bilayer-thick region toward the left in Figure 4/spot (e) evolves from an amorphous island with approximate bilayer thickness (0 min) to a crystalline bilayer with 2H stacking (14 min). This 2H bilayer then intermittently evolves to homonuclear bilayer stacking (15 min) only to then return to 2H-type stacking (17 min and 18 min in Figure 3). This time-dependent appearance/disappearance of MoS$_2$ polymorphs indicates that various polymorphs cannot only co-evolve but also dynamically transform into each other during processing. Similarly, the in situ e-beam exposure videos (Video S1 and Video S2, corresponding to Figure 5) indicate a highly dynamic local evolution during the overall a-MoS$_2$ to nc-MoS$_2$ crystallization, where in particular the emerging 2H bilayer grains are far from static but exhibit alternating growth and shrinkage periods. A key question behind such dynamics is the underlying mechanism of atomic movement. In this context, currently little is known about the diffusing moieties in MoS$_2$ during crystallization and restructuring.$^{62−64}$ This results from the difficulty of their direct observation due to their presumably fast diffusion speeds.$^{37}$ While even the best time resolution in our data during the continuous e-beam exposure in situ videos ($\sim$2.7 s per frame, as shown in Video S1 and Video S2, corresponding to Figure 5) is insufficient to directly observe diffusion species, close inspection of our STEM data can give hints of the diffusing species in our e-beam-induced MoS$_2$ restructuring. We note that some of the adventitious carbon resides on the bare graphene areas in Figure 1a, Figure S, Video S1, and Video S2 can act as intermediate traps for species diffusing over the graphene, thus allowing to draw some preliminary conclusions about moieties diffusing between MoS$_2$ clusters on the graphene: Our element-specific HAADF data identify isolated Mo atoms on the graphene support (some examples labeled in Figure 3c and as spots (f) in Figure 4) that change their location and attach/detach from larger MoS$_2$ structures during the time series in Figures 3 and 4. Such suspected diffusion of Mo atoms between MoS$_2$ clusters on graphene is also consistent with Video 5a/Video S1 where we also observe at better temporal resolution in the in situ video multiple instances of positional changes of Mo atoms during e-beam exposure that lead to overall mass transport from one MoS$_2$ cluster to another across the graphene. An example of this is the evolution of a “neck” between two eventual MoS$_2$ clusters visible left of the image center in Video S1 (location of forming neck indicated in Figure 5a/180 s by a white arrow). Another example is the appearance and diffusional movement of several isolated Mo atoms in Video S1 below the 2H bilayer cluster (location indicated in Figure 5a/743 s by a white arrow). As such our data indicate that some Mo mass transport is occurring between MoS$_2$ clusters across the graphene support during a-MoS$_2$ crystallization and restructuring. Given the lower atomic number of S atoms, clear identification of isolated S on the basal plane of the graphene support next to adventitious carbon adsorbates is more challenging in our data. We have however labeled as spots (g) in Figure 4 some candidates that may be attributed to single S atoms on the graphene basal plane, which would suggest that also isolated S atoms are diffusing over the graphene during the restructuring. Besides longer range mass transport between adjacent grains, a second type of diffusion during the restructuring is short-range diffusion of atoms within a given grain. An example of such diffusion events within a grain is found in Figure 3c where a Mo adatom is intermittently located on a MoS$_2$ bilayer patch, consistent with a recently identified$^{65}$ metastable adatom configuration on a MoS$_2$ lattice. The in situ data in Video S2 further shows multiple instances of diffusional steps and positional changes between adjacent atoms within a given bilayer MoS$_2$ grain during its crystallization from a-MoS$_2$ to 2H MoS$_2$ (location indicated by white arrow in Figure Sb/190 s). Thereby our data show that such short-range diffusion events within a given grain are another major mechanism of crystallization and restructuring of a-MoS$_2$ to nc-MoS$_2$.

After close inspection of atomically resolved information, we quantitatively analyze the data in Figures 1 and 4 with respect to the visual notion of a-MoS$_2$ dewetting from the graphene support during its crystallization to nc-MoS$_2$ on wider scale: The analysis in Figure S5 shows that for the STEM data in Figure 1a from 0 to 15 min, the bare graphene area notably increases, while, conversely, MoS$_2$ regions with monolayer and submonolayer MoS$_2$ coverage reduce and MoS$_2$ regions with bi- and trilayer coverage slightly increase in area. This confirms the visual impression that the low coverage a-MoS$_2$ clusters dewet from the graphene support and the thus released Mo and S attaches on average to thicker MoS$_2$ regions. Previous theoretical work has predicted (based on considerations of edge energies and interlayer binding in nc-MoS$_2$ clusters) an increasing equilibrium average layer number for MoS$_2$ crystallites with increasing lateral size.$^{29}$ For our data, this would suggest that our MoS$_2$ clusters possibly transform toward their equilibrium thickness/lateral size ratio by the observed dewetting process via the energy input from the e-beam.

Our atomic-scale in situ observations during crystallization and restructuring of MoS$_2$ have far elucidated two key points: First, various MoS$_2$ polymorphs can co-exist and evolve in parallel for fixed processing conditions. This links directly with previous ex situ reports on in-layer polymorphism$^{12−15}$ and co-existence of various stacking types$^{17,18,21,22}$ in ex situ processed MoS$_2$ including chemical synthesis and CVD. As a second and equally important point, our in situ data now clarify that the structural evolution of the MoS$_2$ leading to such polymorphism is not static but highly dynamic, where phases appear/disappear and transform into each other over time. Observation of such dynamics intrinsically requires an in situ approach as employed here.

While in our experiments the monolayer graphene onto which the a-MoS$_2$ is deposited onto is primarily employed as substrate for high-resolution STEM,$^{6}$ the many emerging applications of vertical MoS$_2$/graphene heterostructures in energy, (opto-)electronics, and catalysis$^{10,65−68}$ make also the properties of this MoS$_2$/graphene heterostructure interesting as such. A key drawback toward their elucidation via the data presented in Figures 1–5 is however that the lattice of the supporting graphene is not resolved in these images due to nonoptimal imaging conditions for the lighter carbon ($Z_C = 6$) as well as static residual adventitious carbon contamination which is typical for graphene samples from sample transport and storage in air. This precludes the assessment of orientational relations between the underlying graphene and...
the crystallizing MoS₂ in Figures 1−4, despite the interesting observation that after 17 min electron beam exposure, the FT in Figure 1b shows signs of one discrete hexagonal spot pattern across several nonconnected MoS₂ crystallites. Such a discrete six-fold FT pattern would suggest a dominant orientation of the crystallized MoS₂ that in turn opens the interesting question whether this dominant orientation may be related to a possible epitaxial relationship of the MoS₂ to the underlying graphene support. Previous literature suggested that the nonexistence²²,⁶⁹,⁷⁰ or existence²²,⁷¹,⁷² of MoS₂/graphene epitaxy is highly process parameter dependent, resulting from the rather weak van der Waals interaction between MoS₂ and graphene.⁶⁹ When MoS₂/graphene epitaxy was found in previous work, rotational misalignment distributions peaked at 0° and 30°.²²,⁷¹,⁷²

To resolve a possible orientational correlation between the graphene support and the crystallizing nc-MoS₂ domains under our conditions, we present the e-beam crystallization sequence in Figure 6a. In this series the graphene support in as-deposited state (0 min e-beam exposure) shows both adventitious carbon covered but also atomically clean graphene areas. The image center of the latter, the six-fold lattice of a single crystalline covered but also atomically clean graphene areas. In the inset of the middle shows a (medium angle annular dark field) close-up of an atomically clean graphene area to resolve the supporting graphene lattice and its orientation by the FT under the inset. (b) HAADF STEM image of the same location as (a) after 35 min continuous e-beam exposure. The FT underneath (b) reveals that the a-MoS₂ has crystallized under the e-beam to a single nc-MoS₂ grain, which is misoriented to the graphene lattice directions seen in (a) by a rotation of ∼30°.

Figure 6. (a) HAADF STEM image of ∼2 nm-thick MoS₂ on graphene before continuous e-beam exposure (0 min). The corresponding FT underneath (a) is consistent with a-MoS₂. The inset in the middle shows a (medium angle annular dark field) close-up of an atomically clean graphene area to resolve the supporting graphene lattice and its orientation by the FT under the inset. (b) HAADF STEM image of the same location as (a) after 35 min continuous e-beam exposure. The FT underneath (b) reveals that the a-MoS₂ has crystallized under the e-beam to a single nc-MoS₂ grain, which is misoriented to the graphene lattice directions seen in (a) by a rotation of ∼30°.

Figure 7 therefore presents time-resolved STEM measurements on comparatively thicker a-MoS₂ films of ∼10 nm nominal thickness. In particular, we are comparing in Figure 7 a region which was partly shadowed during MoS₂ deposition and is therefore of somewhat lower thickness (darker HAADF signal in the central region of Figure 7a) with regions consisting of the full deposited ∼10 nm nominal thickness (bright HAADF signal at the left and right edges of Figure 7a). In keeping with our data for the thinner MoS₂ films above, the thin region in the center of Figure 7a shows no in-plane order for the as-deposited films (0 min e-beam illumination) and is consistent with a-MoS₂. Similarly, the thicker regions toward the left and right edges of Figure 7a are largely amorphous in their visual appearance in the as-deposited state (0 min). This is also corroborated by the corresponding FT in Figure 7b (0 min). We note, however, that on the left side in the thicker region in Figure 7a (0 min), two pronounced lattice fringes with a spacing of ∼0.6 nm are visible. Such ∼0.6 nm layer spacing is indicative of the (002) layer distance in MoS₂, therefore suggesting an image interpretation of MoS₂ planes being parallel to the e-beam and thereby being perpendicular to the graphene support.

Upon continued e-beam exposure we find clear signs of crystallization for the thicker a-MoS₂. As above, the thinner regions of Figure 7a crystallize with MoS₂ layers parallel to the graphene support (22 and 34 min, as also shown at higher magnification in Figure 7c,d). Concurrently and unlike the thinner films above, in the thicker MoS₂ regions, multiple sets of ∼0.6 nm lattice fringes appear upon e-beam exposure. These sets of ∼0.6 nm fringes each consist of ∼3 to ∼9 fringes
parallel to each other, while the individual sets are rotated in plane with respect to each other. The appearance of such sets of ~0.6 nm fringes is a clear sign of crystallization of the initial a-MoS₂ in the thicker regions to nc-MoS₂ with the MoS₂ layers in the direction perpendicular to the graphene support. This is also well reflected in the FT data in Figure 7b (34 min) that shows the corresponding MoS₂ (002) reflections (which are naturally missing in the FTs of the thin MoS₂ which crystallized with the layers parallel to the support in Figure 1b). The emergence of MoS₂ (002) reflections in the ~10 nm MoS₂ films upon e-beam exposure is also corroborated by TEM measurements at a larger field of view, shown in Figure S6. Importantly, further inspection of the STEM data in Figure 7a (34 min) and Figure 7d shows that under the ~0.6 nm fringe sets, an in-plane ordered MoS₂ lattice continues. This leads to the interpretation of the data in Figure 7 that thicker MoS₂ regions (~10 nm nominal thickness) crystallize during e-beam exposure in a two-segment morphology: The first few MoS₂ layers near the support interface crystallize parallel to the graphene support (same as the thinner ~2 nm MoS₂ regions in Figure 1) but then farther away from the graphene support the direction of the evolving MoS₂ layer orientation changes for the thicker films, resulting in further MoS₂ to crystallize with its layers perpendicular to their support (i.e., perpendicular to the graphene support and the first few MoS₂ layers). We note that these layers with overall perpendicular orientation may also partly be curved along their length. Figure 7e schematically illustrates the evolution of this suggested two-segment nc-MoS₂ film structure with in-plane crystallization near the support interface and out-of-plane crystallization beyond for thicker a-MoS₂ films. Previous literature has found both parallel and perpendicular layer growth in thicker MoS₂ films depending on exact synthesis conditions. Importantly we note that the observation of our two-segment morphology is in excellent agreement with previous ex situ studies on annealed PVD MoS₂ films, that is, films that were deposited and processed under similar deposition condition as ours, in which the same two-segment morphology was reported.

CONCLUSIONS

In summary, our work provides in situ atomic-scale observations of the crystallization and restructuring of the important TMDC MoS₂. Our data elucidate the complex evolution of a material with such pronounced and, as we show, dynamic polymorphism. Our observations thereby visualize multiple fundamental processes that are underlying the varied MoS₂ morphologies obtained in previous ex situ MoS₂ processing studies. Our beam-driven in situ imaging and materials modification approach can be expected to be extendable to several other TMDCs and 2D materials that can crystallize equally easily as MoS₂ (i.e., 2D materials that can grow without the requirements for high processing temperatures and for a thick process catalyst). We expect that our here presented in situ methodology will contribute toward an improved fundamental atomic-scale understanding of TMDC and 2D materials synthesis and integration processing.

METHODS

Samples for high-resolution STEM were prepared as follows: First continuous monolayer graphene films were grown by CVD on Cu catalysts in a CH₄/H₂/Ar mixture at 960 °C. The graphene films were then suspended as membranes by transfer onto holey carbon-foil TEM grids with regular hole arrays (Quantifoil) using a polymer-free transfer process, which avoids the detrimental residues typically associated with polymer-based transfers and thus ensures an as clean as possible MoS₂/graphene interface from scalable processing. Onto these graphene-covered TEM grids, PVD MoS₂ (nominal thicknesses from ~2 nm to ~10 nm) was then sputter deposited from a compound MoS₂ target. During PVD, the samples were left at nominal room temperature (i.e., without intentional substrate heating applied). These conditions are known to lead to deposition of a-MoS₂. Throughout and after fabrication, samples were stored and transported in ambient air.

STEM was measured in an aberration corrected Nion UltraSTEM 100 at an electron acceleration voltage of 60 kV, acquiring HAADF STEM images at a wide e-beam was used for imaging and SAED at electron dose rates 34,47 directly under the beam of ~5 × 10⁶ e⁻ Å⁻² s⁻¹, which in turn equate to average dose rates of ~5 × 10⁹ e⁻ Å⁻² s⁻¹ for continuous scanning of a 10 nm × 10 nm area as in Figure 1. For the crystallization series in STEM, continuous e-beam exposure was achieved via continuous STEM scanning. We note that for imaging at a wider field of view/low dose rates in the TEM, the e-beam-induced crystallization correspondingly proceeds less pronounced. In order to minimize reactions with residual gas species during STEM imaging, the employed STEM column leaves the sample in a vacuum of ~10⁻⁹ mbar during imaging. During STEM imaging samples were not intentionally heated. Note that all samples were annealed at ~140 °C in a vacuum of 10⁻⁵ mbar for ~8 h prior to loading into the STEM in order to desorb adventitious hydrocarbons and adsorbed water from sample storage in ambient. We crosscheck by TEM and SAED without preheating that this low-temperature vacuum bake did not result in any significant a-MoS₂ crystallization. BF-TEM and SAED at 80 kV and 200 kV electron acceleration voltage were measured in a Philips CM200 TEM with the sample in a vacuum of ~10⁻⁶ mbar. In the CM200 TEM, a wide e-beam was used for imaging and SAED at electron dose rates (~4 × 10⁷ e⁻ Å⁻² s⁻¹) that did not induce a-MoS₂ crystallization. In order to induce a-MoS₂ crystallization in the CM200 TEM, the e-
beam was focused to achieve electron dose rates of \( 3 \times 10^3 \text{e}^{-} \text{Å}^{-2} \text{s}^{-1} \). SAED at 60 kV electron acceleration voltage was measured in a FEI Tecnai F20 TEM with a vacuum of \( 10^{-6} \text{ mbar} \) and beam current densities of \( 2 \times 10^2 \text{e}^{-} \text{Å}^{-2} \text{s}^{-1} \) for imaging/SAED and of \( 1 \times 10^2 \text{e}^{-} \text{Å}^{-2} \text{s}^{-1} \) to induce \textit{in situ} crystallization. EDX was measured at 60 kV in the F20 TEM with an EDAX Apollo XLTW SDD system. Elemental quantification from thus obtained EDX data of the a-MoS\(_2\) films was crosschecked by additional EDX measurements using an Oxford Instruments X-max system installed in a Zeiss Supra S5VP scanning electron microscope (SEM) operated at 20 kV that was calibrated against mechanically exfoliated MoS\(_2\) reference crystals. Additional SEM-based EDX measurements on blanket a-MoS\(_2\) films deposited at identical conditions as the a-MoS\(_2\)/graphene heterostructures confirmed lateral homogeneity of stoichiometry of our a-MoS\(_2\) films. For details on structural data analysis methodology and our DFT calculations see the Supporting Information.

## ASSOCIATED CONTENT

**Supporting Information**

The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/acsnano.8b04945.

Additional STEM data and analyses, EDX data, DFT calculations and details on methodology (PDF)

Video S1 (AVI)

Video S2 (AVI)

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**Notes**

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