

Atomically-Precise Texturing of Hexagonal Boron Nitride Nanostripes

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Monolayer hexagonal boron nitride (hBN) is attracting considerable attention because of its potential applications in areas such as nano- and opto-electronics, quantum optics and nanomagnetism. However, the implementation of such functional hBN demands precise lateral nanostructuring and integration with other two-dimensional materials, and hence, novel routes of synthesis beyond exfoliation. Here, a disruptive approach is demonstrated, namely, imprinting the lateral pattern of an atomically stepped one-dimensional template into a hBN monolayer. Specifically, hBN is epitaxially grown on vicinal Rhodium (Rh) surfaces using a Rh curved crystal for a systematic exploration, which produces a periodically textured, nanostriped hBN carpet that coats Rh(111)-oriented terraces and lattice-matched Rh(337) facets with tunable width. The electronic structure reveals a nanoscale periodic modulation of the hBN atomic potential that leads to an effective lateral semiconductor multi-stripe. The potential of such atomically thin hBN heterostructure for future applications is discussed.

1. Introduction

Hexagonal boron nitride (hBN) is an attractive two-dimensional (2D) material for high performance electronics and photonics applications. It competes as insulating substrate with SiO₂ or Al₂O₃, and it is also promising as protective coating for metallic films that support intense 2D plasmons,^[1,2] as a single photon emitter with deep UV lasing,^[3] or as a superconductor through suitable doping.^[4] Although hBN flakes may be obtained by mechanical exfoliation of bulk crystals, a single hBN monolayer can be readily synthesized on metal -surfaces,^[5] leading to structurally and chemically robust substrates that frequently exhibit nanoscale patterns. This makes hBN-covered metals excellent platforms to achieve functional interfaces with atoms, molecules, and aggregates, as well as

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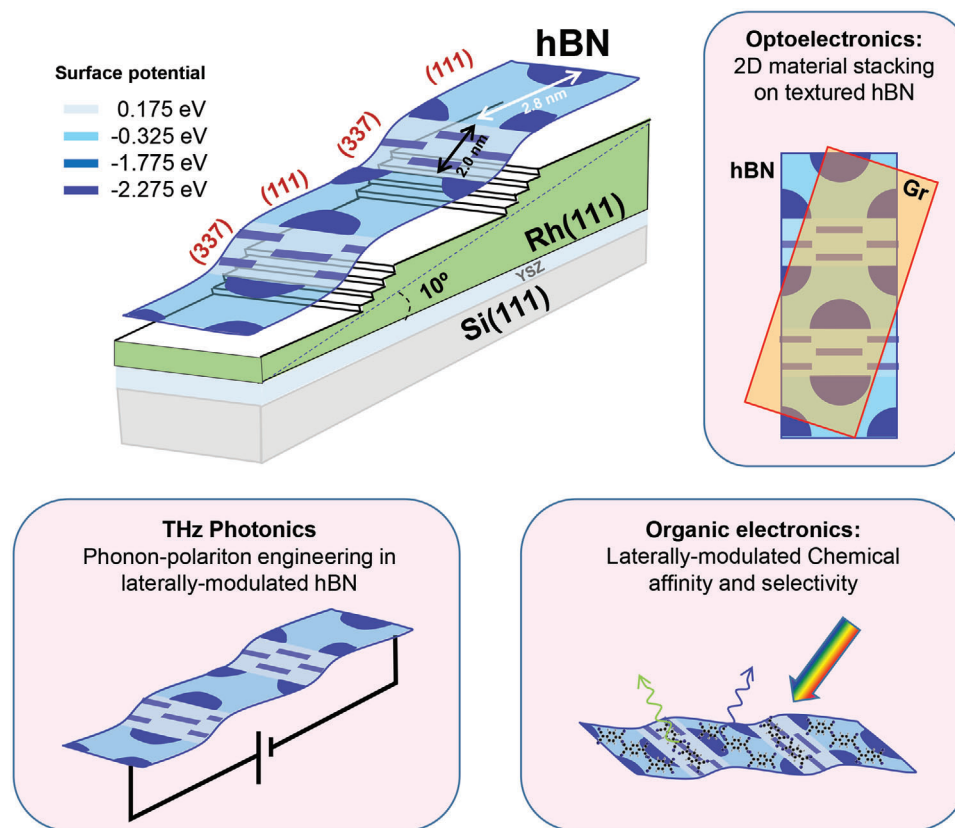


Figure 1. Lateral nanopatterning of hBN through epitaxial growth. We explore this concept by studying vicinal hBN/Rh interfaces. CVD growth of hBN induces periodic (111)/(337) faceting of the Rh substrate. The hBN monolayer uniformly coats the faceted substrate, defining an effective lateral hBN/hBN heterointerface with periodic surface potential texturing. Using commercial Rh(111) films^[17,18] one could exploit such chemical and electronic modulation of hBN to explore phonon–polariton THz excitations, selective growth of optically-active molecules, and vertical stacking with other 2D materials.

to develop hybrid 2D materials, such as twisted van der Waals stacks or 2D heterostructures.^[6–8] The latter hold a great potential for atomically thin circuitry, such as superstructures formed with isostructural graphene,^[9–13] which are optimal to engineer gaps and doping, as well as to tune and enhance spin scattering. However, exploiting fine hBN-based nanostructures requires structural quality down to the atomic scale, which lies beyond current lithographic capabilities. The bottom-up vapor growth is the alternative, which also works for 2D hybrids,^[14] although general procedures to control shape, size, and spatial order of surface phases are still lacking.

Here, we demonstrate the bottom-up synthesis of nanostrapped hBN heterostructures with atomically sharp interfaces. We follow the standard chemical vapor deposition (CVD) growth route, using Rhodium (Rh) vicinal surfaces as one-dimensional (1D) templates. We show that the excellent registry with the Rh(337) plane triggers periodic (111)/(337) faceting of the substrate, as sketched in **Figure 1**. The hBN monolayer carpet-coats the faceted substrate, but the distinct symmetry and atomic interaction at each hBN/Rh phase give rise to a textured and periodically modulated surface potential. A 1D lateral hBN (111)/(337) heterostructure arises, featuring defect-free boundaries and significant band offsets. Since size tunability of phases can be gained by selecting the Rh vicinal plane, we envision the hBN (111)/(337) faceted sys-

tem as a model platform to mold and probe 1D phonon–polariton excitations in the THz regime,^[15] to achieve selective growth of atoms, aggregates, and molecular adsorbates for organic optoelectronics and catalysis,^[5,16] and to tailor the 3D stacking with other 2D materials, such as graphene.^[7]

2. hBN-Induced Faceting across a Rh Curved Surface

We examined striped hBN monolayers over an ample range of Rh vicinal orientations with a curved Rh surface (c-Rh). The sample is schematically depicted in **Figure 2a**. It is a cylindrical section of a Rh single crystal with the cylinder axis along $[1\bar{1}0]$ direction. The curved surface spans all vicinal surfaces (α angle) from the high symmetry (111) ($\alpha = 0$) at the right edge of the sample, passing through the (557) plane ($\alpha = 9.5^\circ$) at the sample center, and reaching the (337) surface ($\alpha = 23.5^\circ$) near the left side edge. The freshly prepared surface is heated to $T = 1000\text{K}$, and then exposed to borazine, leading to full monolayer coverage of hBN across the entire curve, as judged from scanning tunneling microscopy (STM) and low energy electron diffraction (LEED). At the Rh(111) plane, this results in a hBN nanomesh structure,^[19] which arises due to atomic lattice mismatch and variable bonding

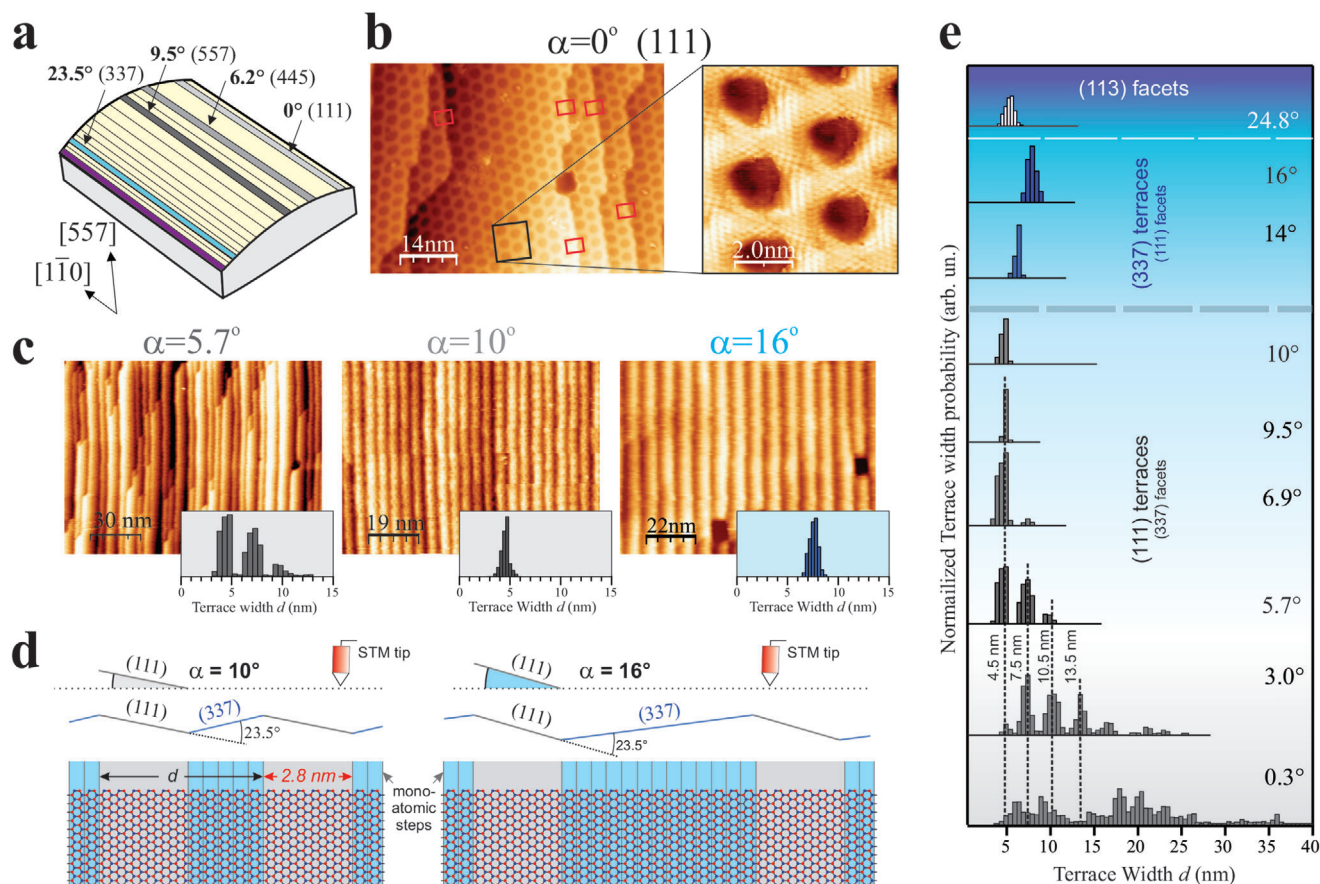


Figure 2. Evolution of the hBN nanostrapped system on a curved Rh surface. a) Sketch of the c-Rh sample with some vicinal α angles and corresponding crystal planes indicated. b) 4K-STM image and zoom-in near the hBN-covered Rh(111) plane ($\alpha = 0^\circ$). c) 300K-STM topographies and terrace width histograms at characteristic α angles: (111) terraces reveal discretized hBN nanomesh quanta ($\alpha = 5.7^\circ$), a single nanomesh quantum ($\alpha = 10^\circ$), and hBN-(337) nanostrapped ($\alpha = 16.0^\circ$). d) Description of the STM scan over the hBN/Rh interface at $\alpha = 10^\circ$ and $\alpha = 16^\circ$, where hBN-(111) and hBN-(337) nanostrapped are respectively imaged as “terraces.” e) Terrace width distribution histograms across the curved hBN/c-Rh surface. At low α (gray region), peaks correspond to multiples of the (111) nanomesh, down to a single quantum at $\alpha = 10^\circ$. Above $\alpha = 10^\circ$ (blue region), the distribution shifts to the right side, reflecting the increasing width of the (337) facets. Beyond the (337) plane ($\alpha > 23.5^\circ$, purple region), (113) faceting arises.

between the hBN layer and the Rh substrate.^[20] Images in Figure 2b, acquired with a 4K-STM, show the atomically resolved structure close to the Rh(111) plane and near substrate steps. We observe that (111) terraces are delimited by steps corresponding to Rh(111) single step height, with upper and lower terraces coated with a continuous hBN film. Within (111) terraces, the lattice constant of the hexagonal nanomesh is $l = 3.2$ nm, agreeing with former studies,^[19,20] while across the steps the pore lattice is coherently connected (red rectangles), indicating that hBN forms a continuous layer that carpets atomic steps (see also Supporting Information for a graphical demonstration).

Away from the (111) direction, the hBN monolayer interacts with the step array of the substrate elastically, leading to faceting. We examined such interaction across the entire hBN-covered c-Rh surface by STM at $T = 300$ K. Figure 2c shows STM images acquired at three characteristic points at $\alpha = 5.7^\circ$, 10° , and 16° , together with their respective terrace-width d distribution histograms. At $\alpha = 5.7^\circ$ we observe straight and parallel hBN nanostrapped, which fit in Rh(111) terraces of discrete sizes, separated by bunches of one to four Rh atomic steps (see Supporting Infor-

mation). The histogram points to an interface that is broken up in a random distribution of hBN-covered (111) terraces of width $d \approx 4.6, 7.1,$ and 9.6 nm, alternated with step bunches. Consequently, the terrace width increases approximately in multiples of the nanomesh periodicity in the perpendicular direction to the steps (nanomesh quantum $\Delta d \approx l \times \cos 30^\circ = 2.8$ nm). The lowest value $d \approx 4.6$ nm agrees with the nanomesh quantum plus the effective width of the step bunch (1.8 nm). The central panel of Figure 2c corresponds to the surface at $\alpha = 10^\circ$. At this vicinal angle, the hBN monolayer becomes a periodic structure with $d \approx 5$ nm of nanostrapped delimited by bunches of 3–5 monoatomic steps and, most important, with minimal size fluctuations around one nanomesh quantum.

The right-hand side STM micrograph of Figure 2c displays the growth of hBN on a densely stepped c-Rh area at $\alpha = 16^\circ$. In the STM image we observe well-defined hBN nanostrapped, but compared with the $\alpha = 10^\circ$ image, terraces appear wider with $d \approx 8$ nm and the step direction has reversed from uphill to downhill. This effect is schematically explained in Figure 2d. At $\alpha = 10^\circ$, (111)-oriented nanostrapped are wider than hBN-covered

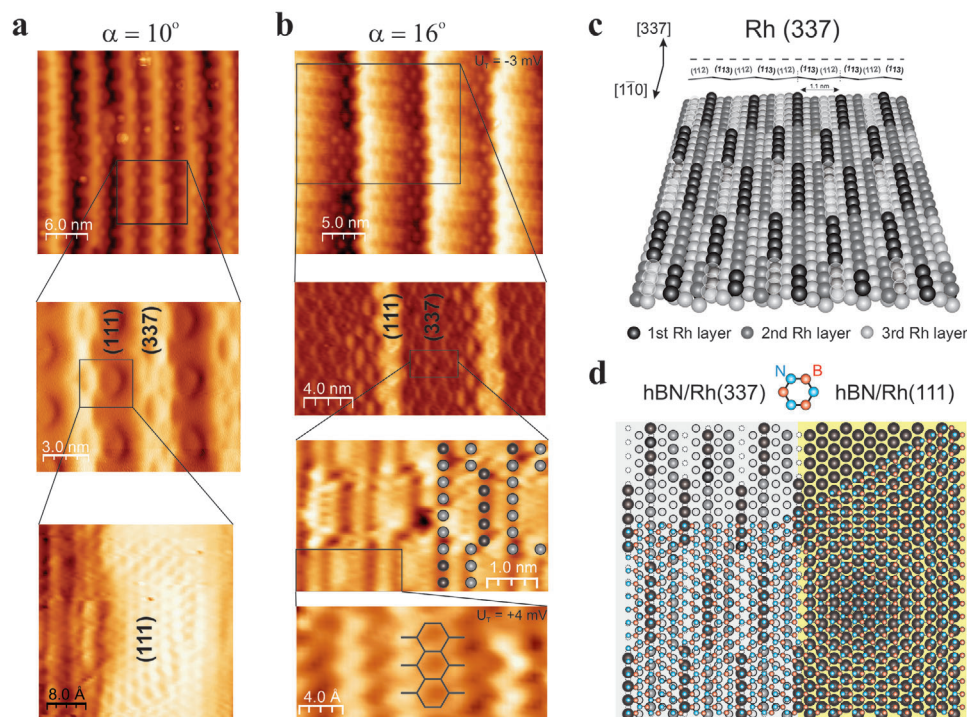


Figure 3. Atomic structure of hBN-covered Rh(111)-(337) facets. a) and b) 4K-STM topographies (top) and progressive filtered zoom-ins (second to bottom) at $\alpha = 10^\circ$ and $\alpha = 16^\circ$, respectively. b) The atomic positions of the step in the reconstructed (337) surface appear overlaid (third image), as well as the hBN lattice (fourth image). c) Model for the Rh(337) surface with selective step atom removal (white dashed balls). Black balls mark the outermost surface atoms, while gray balls become lighter in successive surface levels. d) Model for a continuous hBN monolayer (blue-red balls and sticks) in a coherent (111)/(337) lateral interface. In hBN/Rh(111), the coincidence lattice leads to nanomesh pores around N/Rh on-top sites. In hBN/Rh(337), black balls mark the out-protruding step atoms, which correspond to N/Rh on-top sites of the hBN overlayer.

step bunches, and hence (111) nanostructures appear as terraces in the STM image. At $\alpha = 16^\circ$ (111) nanostructures are narrower than stepped facets, which are thus imaged as terraces, while hBN-(111) nanostructures appear as downhill steps. 4K-STM, ARPES, and LEED experiments (see below) prove that the orientation of the stepped facets is the Rh(337) direction. This transition in the orientation of the apparent terrace beyond $\alpha = 10^\circ$ is reflected in the terrace-width distribution analysis performed across the entire surface, shown in Figure 2e. This provides a unified picture of the hBN/c-Rh interface. In the $\alpha < 10^\circ$ range, (111)-oriented terraces are discrete multiples of the perpendicular nanomesh. With increasing α , the center of gravity of the terrace width distribution shifts toward the first nanomesh quantum. For $\alpha > 10^\circ$, hBN-(337) nanostructures turn into apparent terraces, and hence, the histogram shifts to the right side reflecting the increasing (337) terrace width. Importantly, a sharp size distribution remains, demonstrating that hBN-(337) nanostructures have a single characteristic width at each vicinal angle over the $10^\circ < \alpha < 23.5^\circ$ range. Beyond 23.5° , hBN-(111) nanostructures disappear and hBN-covered (113) facets emerge.

3. The hBN-Covered Rh(111)/Rh(337) Interface

Next we focus on the hBN-induced Rh(111)-Rh(337) periodic faceting, which exhibits straight, and sharply defined hBN nanostructures with coherent coupling between (111) and (337) phases. The atomic arrangement at the hBN/Rh interface of each phase is

investigated in detail using 4K-STM. Images in Figure 3a,b have been acquired again at $\alpha = 10^\circ$ and $\alpha = 16^\circ$ on the c-Rh surface. The larger view in top panels corresponds to the direct topography, while lower panels are image derivatives showing progressive zoom-ins acquired at varied tip bias. Such image processing removes the hill-and-valley modulation, allowing us to define the lateral (111)/(337) interface, which shows no sign of discontinuity. From 10° to 16° , (337) terraces widen from $d_{337} = 2$ nm to ≈ 8 nm, while the (111) nanostructure width barely changes. The internal structure of each phase is visualized in the bottom panels of Figure 3a,b. This includes the hBN atomic lattice in both hBN-(111) and hBN-(337) nanostructures, as indicated with the overlaid structure. The hBN-(111) nanostructure shows a wavy wire running parallel to the step [bright protrusion in top panel of (b)], with half-pores at nanostructure boundaries. The hBN-(337) nanostructure exhibits a characteristic texture, with the $l = 3.2$ nm nanomesh periodicity in the parallel direction, and a reduced one of 1.1 nm perpendicular to the steps, both defining a 2D structure with uniaxial symmetry. Such nanoscale texture in the hBN layer is explained by the periodic re-arrangement of the Rh(337) substrate steps described in Figure 3c.

Adsorption-induced faceting of vicinal surfaces is driven by the optimal registry of the adlayer atomic lattice with the step spacing, such as to reduce interface strain.^[21] For hBN/metal systems, characteristic facets are expected for step periodicities that match an entire number of hBN cells.^[22] In the hBN/Rh system, we may compare the step spacing in the Rh(337) plane with the distance

between contiguous N rows in hBN. As shown in Figure 3c, the (337) plane can be viewed as an alternating sequence of (112) and (113) terraces, resulting in a $l_{337} = 1.1$ nm lattice constant perpendicular to the steps. This closely corresponds to five rows of N atoms ($5 \times a_{\text{hBN}} \times \cos 30^\circ = 1.08$ nm) in the hBN monolayer with $a_{\text{hBN}} = 2.504 \text{ \AA}$ being the hBN lattice constant. Thus, by slightly (2%) stretching the hBN monolayer, only N atom rows are in coincidence with the out-protruding step-edge. Yet, the mismatch along the steps leads to periodic off-registry (strained) N-bridge positions. The substrate reconstruction proposed in Figure 3c solves this problem.

The Rh(337) unit cell contains two rows of step atoms at different levels, with the outermost one corresponding to the downhill step of the (112) terrace. As shown in Figure 3c, removal of 7 out of 12 step atoms in this row (dashed-white transparent balls) reverses the (112)–(113) series and moves the contiguous step segments at the topmost surface position. Now, the outermost surface layer (black balls) is made of 5–6 step-atom segments, defining a 2D lattice identical to that imaged in Figure 3b. In Figure 3d we overlay the hBN lattice on top of a (111)/(337) interface. The Rh(337) step re-arrangement has the double effect of eliminating 7 off-registry N-bridge step sites, while creating six new registry N on-top step sites. Thus, the Rh(337) reconstruction relieves strain by improving registry, making the hBN/Rh interface more robust through exclusive N/Rh on-top bonding. Note that the $l = 3.2$ nm nanomesh periodicity remains along the steps, enabling the coherent coupling with neighboring hBN-(111) nanostripes, which in turn explains the smooth coating of the faceted substrate (see also Figure S8, Supporting Information for a graphical explanation). Similar hBN/Rh step-lattice registry and density of N on-top step anchoring points can be obtained when considering separate hBN/Rh(112) and hBN/Rh(113) interfaces, as shown in Figure S3, Supporting Information. However, in (112) and (113) planes, off-registry N bridge-bonding to step atoms remains, in contrast to the sole N on-top bonding of Figure 3c, which only the Rh(337) plane allows.

4. Uniaxial π Band Symmetry in the hBN-(337) Surface

The structurally different hBN/Rh(111) and hBN/Rh(337) phases exhibit distinct electronic bands, which we display in Figures 4a,b. The (111)-nanomesh modulation triggers the splitting of all hBN electron levels,^[19,23] from vacuum to valence and core bands, in contrast to the simple sequence of electronic states in non-interacting or registry hBN monolayers.^[24] Figure 4a shows the electron bands measured with ARPES along the $\bar{\Gamma}\bar{M}_{\text{hBN}}$ and $\bar{\Gamma}\bar{K}_{\text{hBN}}$ directions in the hBN/Rh(111) system ($\alpha = 0$). The Rh substrate emission dominates the spectrum from the Fermi energy down to -3 eV, while σ and π bands of hBN appear below this energy. The characteristic π_α – π_β splitting is observed,^[23] due to the different interaction with the Rh substrate at pores and wires.^[25] As explained from density-functional-theory (DFT),^[23,25] the highly bound π_β band originates in the highly interacting on-top N atoms in the pores, while the π_α branch comes from off-registry N atoms in the wires.

The pore/wire nature of the π_α/π_β split-bands in the hBN/Rh(111) interface is well captured in our electron-plane-wave-expansion (EPWE) and photoemission model.^[26] Using

Table 1. π band properties in hBN/metal systems. π band bottom ($\bar{\Gamma}_{\text{min}}$), top (\bar{K}_{max}), and bandwidth ($\Delta_{\Gamma M}$) measured with ARPES in hBN/metal systems. Data for hBN/Ni(111) and hBN/Au/Ni(111) are from Ref. [24]. $\langle V_{\text{N}} \rangle_w, p$ and m^* respectively are the average surface potential at N atoms for wires and pores, and the effective mass used in the EPWE model, which renders the semiconducting gap (E_{F} gap). All magnitudes are expressed in eV.

Interface	$\bar{\Gamma}_{\text{min}}$	\bar{K}_{max}	$\Delta_{\Gamma M}$	E_{F} gap	$\langle V_{\text{N}} \rangle_w$	$\langle V_{\text{N}} \rangle_p$	$\langle m^* \rangle$
hBN/Au/Ni(111)	−8.2	−2.22	4.87	5.08	0.0	0.0	1.00
hBN/Ni(111)	−10.0	−4.45	4.53	5.08	−1.690	−1.690	1.05
hBN/Rh(111)	$−9.8 \pm 0.2$	$−3.0 \pm 0.2$	4.4 ± 0.1	4.20	−0.325	−2.275	1.03
hBN/Rh(337)	$−8.8 \pm 0.2$	$−3.3 \pm 0.2$	2.86 ± 0.1	4.87	0.175	−1.775	1.10

only four parameters, namely, the different scattering potentials at N and B atoms, a rigid potential shift ΔV from wires to pores (first Fourier component of the nanoscale potential), and an effective mass m^* , we fit the experimentally observed energies at $\bar{\Gamma}$, \bar{M} , and \bar{K} points, and obtain the theoretical bands of Figure 4c (see also Figure S4, Supporting Information for more details). We indeed find that a π band splitting arises for $\Delta V < 0$. To fit ARPES bands we require $\Delta V = -1.95$ eV and $m^* = 1.03 \cdot m_e$, with m_e being the free electron mass (see Table 1). Actually, due to the (13×13) periodicity, the simple EPWE model delivers multiple π band replicas within the hBN Brillouin zone, which are averaged out by introducing broadening in the photoemission probability. Note that experimental bands in Figure 4a only mirror the hexagonal symmetry of the atomic lattice, but do not show nanomesh umklapps, sometimes seen for graphene/metal moiré systems.^[27,28]

In the hBN/Rh(337) bands, the uniaxial symmetry brought in by the nanoscale texture is reflected in the band structure, shown in Figure 4b. Data correspond to $\alpha = 24^\circ$, that is, very close to the Rh(337) surface. Due to experimental restrictions we were not able to measure in the exact $\bar{\Gamma}\bar{K}_{\text{hBN}}$ direction but only close to it as indicated in the middle panel of Figure 4e. Nevertheless, we now observe a striking asymmetry from the perpendicular ($\bar{\Gamma}\bar{M}_{\text{hBN}}$) to the parallel ($\bar{\Gamma}\bar{K}'_{\text{hBN}}$) direction along substrate steps. Perpendicular to the steps, the main dispersing feature is broken up in segments, separated by ≈ 0.3 meV mini-gaps (see arrows; the second derivative of the image enhances gap visualization, see Supporting Information), whereas in the parallel direction the π band splits in a set of continuous sub-bands (inclined arrows). Such distinct band topology is correctly reproduced in EPWE calculations of Figure 4d, which are based on the atomic model depicted in Figure 4e. Note that despite the apparent atomic complexity, the hBN/Rh(337) interface is simple in terms of bonding, since all the outermost Rh step atoms bind to N in atop positions. For the EPWE model, it is thus reasonable to assume the same surface potential reduction at N atoms on steps that was used for pores, $\Delta V = -1.95$ eV. In excellent correspondence with the experimental observations in Figure 4b, the model reveals the presence of a fivefold π band splitting at $\bar{\Gamma}$, which leads to a set of dispersing bands along $\bar{\Gamma}\bar{K}_{\text{hBN}}$, and to five mini-bands separated by mini-gaps at Rh(337) Brillouin zone edges ($n \times \pi/d_{337}$) along $\bar{\Gamma}\bar{M}_{\text{hBN}}$.

The hBN/Rh(337) system represents the first successful case of uniaxial symmetry imprinted onto the electronic structure of the hBN monolayer. In pursue of this objective, (110)-oriented

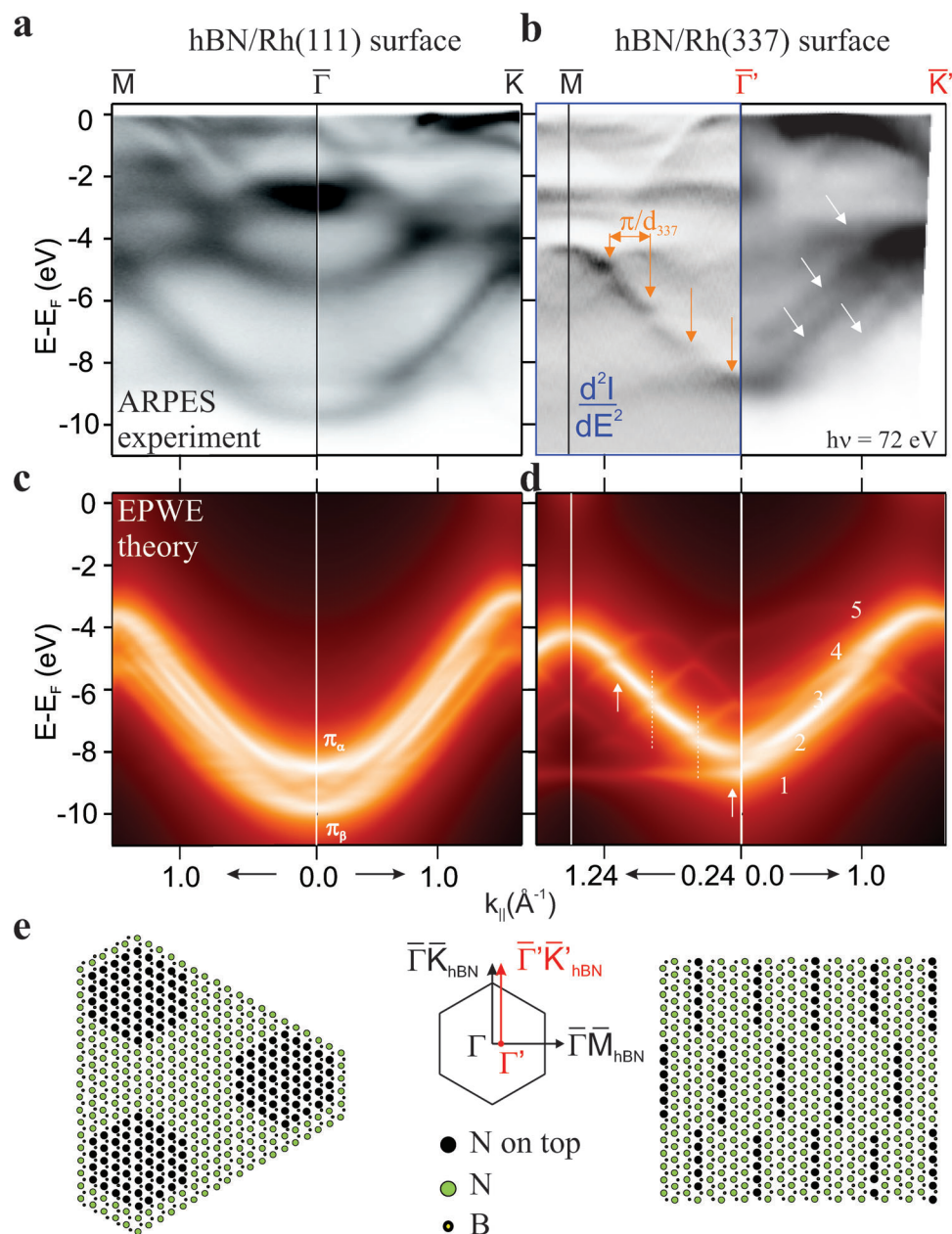


Figure 4. hBN electron bands on Rh(111) and Rh(337). a,b) Experimental hBN bands along the $\bar{\Gamma}\bar{M}$, $\bar{\Gamma}\bar{K}$, and $\bar{\Gamma}'\bar{K}'$ directions measured with ARPES in the Brillouin zones of hBN/Rh(111) and hBN/Rh(337) as indicated in (e). For hBN/Rh(337) in (b), arrows mark mini-gaps at $n \times \pi/d_{337}$ Brillouin zone edges along $\bar{\Gamma}\bar{M}$ and the π band manifold along $\bar{\Gamma}'\bar{K}'$. c,d) Theoretical hBN bands calculated with the EPWE model, assuming the different atomic potentials at pores/steps (N black) and wires (N green) shown in (e). The fivefold π band splitting along $\bar{\Gamma}'\bar{K}'$ is numbered from bottom to top in (d).

metal surfaces were tested in the past,^[29–32] but the hBN film was observed to break up in rotational phases. On Rh(110),^[33] hBN nanostriped domains appear, reflecting the scarcity of favorable interface (N on-top) bonding sites compared to reconstructed Rh(337). Nevertheless, away from high symmetry directions, other hBN/metal surface combinations should exist with optimal lattice matching. Uniaxial vicinal planes with different terrace orientations, such as (110), (110), or (201), as well as other metals, such as Pt, Pd, Ru, or Cu should be explored.^[5] hBN-induced faceting appears to be the rule rather than the

exception,^[22] and this can serve to pinpoint new “magic,” well-matched surfaces, such as Rh(337), and to find nanostriped patterns with varied textures.

In Table 1 we summarize the π band properties for fundamental hBN/Rh systems, namely, free-standing (Au/Ni), 2D atomic-registry [Ni(111)], 2D nanomesh [Rh(111)], and 1D step-registry [Rh(337)], from which we derive some trends. The π band minimum at $\bar{\Gamma}$ ($\bar{\Gamma}_{\min}$) is connected to the ratio of substrate-interacting N atoms in top positions to the total number of N atoms. Such ratio increases from 20% in Rh(337) steps, to 40% in Rh(111)

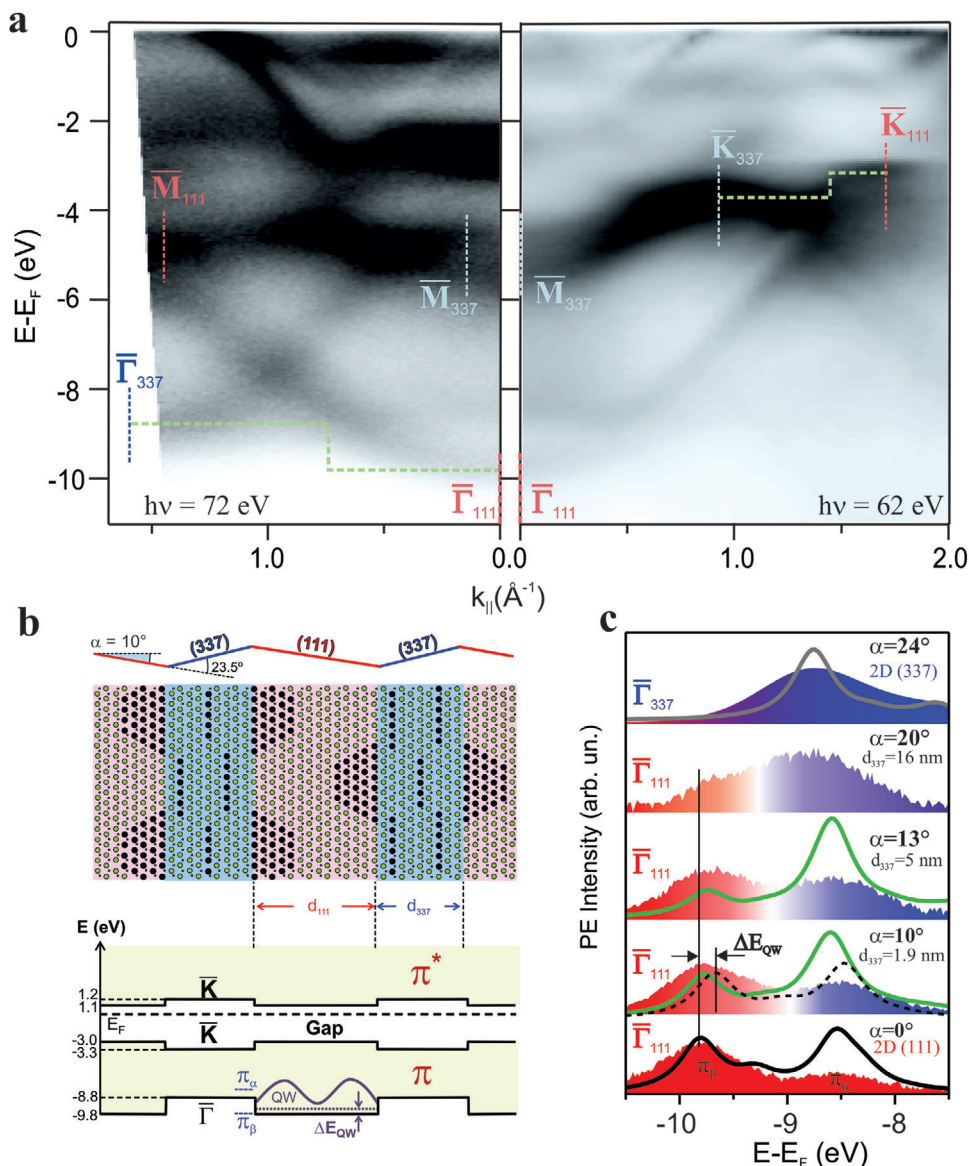


Figure 5. Electronic states in lateral hBN heterostructures. a) Electron bands for the hBN (111)/(337) superlattice measured with ARPES ($h\nu = 72$ eV) at $\alpha = 10^\circ$ in the hBN/c-Rh sample. The faceted geometry leads to separate emission from (111) and (337) facets. The respective $\bar{\Gamma}$, \bar{M} , \bar{K} symmetry points and the resulting heterostructure offsets are indicated. b) Atomic model and band diagram for the hBN (111)/(337) lateral heterostructure, as determined from data and calculations of Figure 4. c) Evolution of the ARPES intensity at $\bar{\Gamma}_{111}$ in the (111)/337 nanostructured system at increasing α . The spectrum on top corresponds to the $\bar{\Gamma}_{337}$ point of the 2D hBN-(337) system. Lines are EPWE spectra at $\bar{\Gamma}$ for heterostructures of fixed $d_{111} = 2.8$ nm and variable d_{337} . ΔE_{QW} marks the expected EPWE shift of π_β in fully decoupled 2.8 nm d_{111} nanostructures.

pores, and to 100% in lattice-matched Ni(111), and directly correlates with the average $\langle V_N \rangle$ potential in the EPWE model. Despite the strong difference in interface bonding, the hBN gap remains unchanged in Ni and Au/Ni, whereas nanoscale texturing in Rh(111) and Rh(337) effectively reduces the gap.

5. Electronic States in Lateral hBN (111)/(337) Heterostructures

The straight boundaries with coherent atomic bonding, together with the distinct band properties of (111)- and (337)-oriented hBN nanostructures define an ideal atom-thick, lateral semiconductor

heterostructure. In Figure 5a we show electron bands as obtained from ARPES measurements at $\alpha = 10^\circ$ along $\bar{\Gamma}\bar{M}_{\text{hBN}}$ and $\bar{\Gamma}\bar{K}_{\text{hBN}}$, where the superlattice is formed by $d_{111} = 2.8$ nm and $d_{337} \approx 2$ nm hBN nanostructures. This nanostructured system is depicted in Figure 5b, together with its band diagram, where we have assumed the energies of Table 1 to define valence and conduction band offsets. Interestingly, the hill-and-valley geometry of the surface plane leads to separate intensity from (111)- and (337)-oriented facets in ARPES scans (see Figure S6, Supporting Information for details). This allows us to track the evolution of the π band emission from each phase, and hence to prove (111)/(337) electronic coupling. This is done for the 2.8-nm-wide d_{111} nanostructures in

Figure 5c. Red-blue shaded spectra correspond to $k_{\parallel} = 0$ in the (111) plane ($\bar{\Gamma}_{111}$ point), whereas green curves are EPWE calculations for the heterostructure of Figure 5b, assuming d_{337} nanostripes of increasing width. The spectral density clearly evolves from the hBN-(111) system (bottom) to hBN-(337) (top) as a function of α , that is, from the 2D (111) nanomesh to increasingly sparse $d_{111} = 2.8$ nm nanostripes. Because of the 1 eV valence band offset at the interface, the π_{β} peak, which lies below the band minimum in the (337) plane (Γ_{337}), is weakly coupled to the (337) bands. Consequently, π_{β} behaves as a (111)-terrace-like state, which shifts up in energy and has its intensity reduced when the (111) surface evolves into a narrow stripe. In contrast, the π_{α} feature that lies above the Γ_{337} minimum resonates with the (337) band and hence remains intense. Note that we are exclusively probing emission from d_{111} -oriented areas of the surface. Thus, Figure 5c demonstrates that d_{111} nanostripes embedded within large d_{337} areas at $\alpha = 20^{\circ}$ produces emission that is coherent with the hBN-(337) system. In fact, the asymmetric intensity variation of π_{α} and π_{β} , as well as the small (50 meV) shift of π_{β} , are reproduced in our calculations for electronically-coupled 111/337 interfaces. In contrast, our calculations permit us to discard a fully decoupled, quantum-well-like d_{111} nanostripe (black dotted spectrum in Figure 5c), which would produce spectra exhibiting a symmetric π_{α} - π_{β} intensity quenching, and a substantially larger peak shift ($\Delta E_{QW} = 150$ meV).

6. Conclusions

We have explored a novel approach to nanopatterning a material with practical technological impact, namely growing hBN monolayers on vicinal surfaces. Growth of hBN on vicinal Rh(111) leads to periodic (111)/(337) faceting, triggered by excellent atomic registry of hBN with a step-reconstructed (337) surface plane. The hBN film forms a continuous layer over the faceted system with alternated (111) and (337) phases, each exhibiting a characteristic internal texturing, and strain-free atomic bonding at the mutual interface, thereby defining an electronically-coherent hBN lateral heterostructure.

In the context of the rich phenomenology exhibited in hBN/metal interfaces,^[5] we devise a vast number of possibilities for vicinal surfaces. These are known to drive the nucleation and growth of azimuthally-oriented hBN flakes,^[34,35] and here we show that they also trigger 1D texturing, as depicted in Figure 1. Beyond the (111)/(337) system, other hBN/metal orientations may also exhibit faceting with distinct electronic properties and textures, and this could be systematically explored with curved surfaces. The nanoscale corrugation of the surface potential can help to imprint, for example, a periodic modulation of energy and filling of molecular levels in organic adsorbates.^[16] It can also have a strong influence in epitaxial growth, for example, in the vertical stacking of transition metal dichalcogenides (TMDs) or graphene, where it may help to tune, in general, electronic properties^[36] and, in particular, the twist angle.^[7] The potential for optoelectronics of such 2D material stacking should also be explored for applications, such as band structure modulation of graphene and TMDs. The hBN (111)/(337) heterostructure is also an ideal platform to manipulate phonon-polariton excitations and their dynamics.^[15] The band structure modulation is expected to affect the optical response of the material, thus

adding an appealing alternative to the suite of existing possibilities in the THz region.

Supporting Information

Supporting Information is available from the Wiley Online Library or from the author.

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Conflict of Interest

The authors declare no conflict of interest.

Data Availability Statement

Research data are not shared.

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