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A ferromagnetic Eu-Pt surface compound grown below hexagonal boron nitride†

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One of the fundamental applications for monolayer-thick 2D materials is their use as protective layers of metal surfaces and *in situ* intercalated reactive materials in ambient conditions. Here we investigate the structural, electronic, and magnetic properties, as well as the chemical stability in air of a very reactive metal, Europium, after intercalation between a hexagonal boron nitride (hBN) layer and a Pt substrate. We demonstrate that Eu intercalation leads to a hBN-covered ferromagnetic EuPt₂ surface alloy with divalent Eu²⁺ atoms at the interface. We expose the system to ambient conditions and find a partial conservation of the di-valent signal and hence the Eu-Pt interface. The use of a curved Pt substrate allows us to explore the changes in the Eu valence state and the ambient pressure protection at different substrate planes. The interfacial EuPt₂ surface alloy formation remains the same, but the resistance of the protecting hBN layer to ambient conditions is reduced, likely due to a rougher surface and a more discontinuous hBN coating.

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1. Introduction

Ferromagnetic two-dimensional (2D) structures are of uttermost importance in spintronics applications. Such 2D magnetic systems can either be 2D van der Waals (vdW) ferromagnets, Or ultrathin magnetic overlayers. Both types of systems present quantum and topological phases, but achieving new exotic properties requires design and investi-

gation of novel materials and architectures. In thin magnetic overlayers, the surface contains transition and/or rare-earth metals. The interest in such 2D ferromagnetic systems is prompted by the reduced atomic-scale size and the diversity of magnetic states that arise. Some of the 2D vdW ferromagnets can be exfoliated from bulk crystals, e.g., Fe₃GeTe₂⁷ or family members.^{8,9} However, most of them are grown by ex situ chemical vapor or atomic layer deposition. 10-13 Magnetic overlayers, on the other hand side, are usually grown by physical vapor deposition under vacuum conditions, a process that does not guarantee an unique 2D layer but may lead to a multilayer system. Achieving a single 2D magnetic layer demands a detailed in situ, atomic-scale investigation, comprising structure and electronic states, as well as chemical stability in ambient conditions. 2D magnetic alloys as well as some of the vdW ferromagnets are quite reactive in air, loosing or changing their magnetic properties. Protection of such surfaces can be achieved by means of ceramic coatings, polymer protection films or deposition of non-reactive metals. Nevertheless, most of these protecting overlayers influence the magnetism of the surfaces leading to a variation of the desired properties. Recently, protection of the surfaces by graphene (Gr), 14-22 hexagonal boron nitride (hBN)15,23-28 and a mixture of both materials²⁹ is being considered. In this context, the use of a hBN protecting layer is very appealing, since it would provide close contact of the ferromagnetic material with a wide-gap semiconductor, enabling charge injection. Therefore, the ques-

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tion that arises is whether we can achieve a sufficiently protective hBN layer that preserves the magnetic properties of the 2D compound in ambient conditions.

Here, we study a hBN-protected ferromagnetic Eu-Pt surface alloy. The Eu-Pt compound is formed after Eu intercalation under the hBN film previously grown on a Pt crystal surface. Metal intercalation below Gr or hBN overlayer has been extensively studied over the last two decades. 20-22,30-32 The purpose in the majority of the works was to separate the 2D overlayer from the substrate. 33,34 Most often this is done by the intercalation of noble metal atoms like Au, Ag, or Cu.³³ Conversely, in order to force a stronger 2D material interface interaction, one can proceed with the intercalation of alkaline³⁵ or earth-alkaline metals.^{36–38} If a too-strongly interacting substrate-2D overlayer is achieved, additional intercalation of oxygen lifts again the 2D layer and re-establishes the original 2D material properties. 20 However, oxygen exposure may result in the oxidation of the protecting hBN layer.³⁹ Eu intercalation has been less investigated^{21,22,30,40,41} despite of its interesting magnetic properties, mainly due to the strong reactivity of this rare earth metal.

All Eu intercalation studies have been carried out on graphite or graphene epilayers, but no experiments exist using hBN.

Among the different rare earth metal compounds, europium alloys are particularly interesting due to the various valence states of the Eu atoms. It may adopt a di-valent Eu²⁺, a tri-valent Eu³⁺, or even a mixed-valent state. For trivalent Eu³⁺, Eu has a $4f^6$ configuration with S = L = 3 and J = 0. The ground state J-multiplet level (in ${}^{2S+1}L_I$ configuration) is 7F_0 presenting a non-magnetic singlet. This situation differs from divalent Eu^{2+} with $\mathrm{4f}^{7}$ configuration, S=7/2, L=0 and J=7/2 leading to a ⁸S_{7/2} ground state. In the latter case, Eu²⁺ is able to form ferromagnetic compounds, e.g., europium chalcogenides. 42 The different valence states of Eu are found to depend on several factors: the surrounding material, the lattice pressure, the number and type of nearest neighbors, etc. In the particular case of Eu-Pt bulk compounds there is a smooth valence transition when changing the stoichiometry from EuPt₅ (completely tri-valent) to EuPt₂ (Eu atoms in a di-valent state).^{43–45} Additionally, valence instabilities can be induced in EuPt₃ by high pressure.46 Valence changes may also happen at the surface due to a reduced coordination. 47 Such transitions from trivalent to divalent configuration have also been observed for Eu-Ni or Eu-Pd compounds. 48,49

Here we present a Eu-Pt surface alloy formed after intercalation of Eu at the hBN/Pt interface. First, we perform a structural analysis of the interface, followed by a detailed electronic and magnetic characterization of the Eu-Pt compound. We demonstrate that the topmost layer under the hBN coat is a 2D EuPt₂ surface alloy, with di-valent Eu atoms that reveal ferromagnetic behavior at low temperature. Next, we check the efficiency of the hBN layer protection in ambient pressure, by analyzing the electronic properties prior and after air exposure. The sample is a platinum crystal curved around the (111) direction (c-Pt). This provides a smooth variation of the crystallographic orientation across the (macroscopic) surface, allowing us to extend the analysis of the EuPt₂ alloy to vicinal Pt crystal planes, characterized by a high density of atomic steps. By scanning our different experimental (electron, photon) probes on top, we can rigorously study the influence of steps and terraces in the structural, magnetic, and electronic properties of the EuPt₂ surface alloy, as well as the protecting quality of the hBN layer.

2. Experimental details

The growth and electronic properties were mainly investigated at the Nanophysics laboratory in San Sebastian, Spain using a combined system containing scanning tunneling microscopy (STM), low energy electron diffraction (LEED), X-ray photoemission (XPS) and angle-resolved photoemission spectroscopy (ARPES). Part of the electronic structure investigations have been carried out at BACH beamline of Elettra synchrotron (Trieste, Italy). The XPS setup in the laboratory is equipped with a Specs Al K_α μ-FOCUS 600 monochromator while the ultraviolet light source consists of a Specs UVS-300 discharge lamp with monochromator (Specs TMM 304) tuned to HeIIa light with $h\nu = 40.8$ eV. At Elettra synchrotron, p-polarized light was applied. All measurements were taken with the sample at room temperature. STM experiments were carried out in a Omicron VT-setup by holding the sample at room temperature and scanning with a W tip. The analysis of the STM images has been performed with WSXM software.⁵⁰ The magnetic properties were investigated at ID 32 of the European synchrotron radiation facility (ESRF) by means of X-ray magnetic circular dichroism (XMCD). For this purpose the sample was placed normal or grazing (70°) with respect to the incoming photon beam and field. The field was ramped between +6 and -6 T with the sample hold at T = 7 K. Horizontal, left and right circularly polarized light (99% polarization) was used for photon energies around the Eu M4,5 X-ray absorption edge.

As a substrate material, a cylindrical sector of a Pt (c-Pt) single crystal was used whose cylinder axis is along a [1 $\bar{1}$ 0] direction. The centre of the curved surface points towards the [111] direction, while the borders are oriented $\pm 15^{\circ}$ with respect to the (111) center (Fig. 4). This curved Pt surface was cleaned by Ar ion sputtering (room temperature) and temperature annealing (1000 K) as well as by occasional oxygen heating (2 × 10⁻⁸ mbar O₂, 950 K) followed by a flash in UHV to 1050 K. This standard procedure, as described elsewhere, 51,52 leads to sharp LEED patterns where the typical step splitting was observed.

hBN was grown by chemical vapor deposition (CVD) process from borazine precursor ($B_3H_6N_3$) (KATCHEM spol. s r.o.). For this purpose the curved Pt crystal was held at 1020 K while borazine was dosed for 20 minutes at 2×10^{-7} mbar, *i.e.*, for 240 L, in order to assure a complete hBN layer. Already a 100 L dosage would be enough for a complete layer.³² As can be observed in Fig. 1, this growth produces a sharp and well ordered moiré pattern in LEED at the Pt(111) position of the curved crystal. On the other substrate positions, corresponding

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(a)

Pt(111) +borazine, 1020K +Eu; 870K +air; +anneal. 570K

Fig. 1 LEED images along the preparation process. (a) Pt(111), (b) after borazine exposure at 1020 K producing a (9 \times 9) moiré pattern, (c) additional Eu deposition/intercalation at $T_{\text{sample}} = 870 \text{ K}$ leading to EuPt₂ layer below hBN and a different moiré pattern, (d) after 6 hours room temperature air exposure and subsequent 570 K vacuum annealing (LEED kinetic energy 70 eV).

(C)

to the vicinal surfaces in the mentioned $\pm 15^{\circ}$ range around (111), the LEED reveals less ordered structures with line like features pointing to a multi-facet structure.

(b)

 (1×1)

Eu was deposited in a third step on top of this hBN/c-Pt substrate while the sample was held at an elevated temperature to allow Eu intercalation below hBN. As pointed out earlier,³⁰ the high temperature is quite important to immediately protect the Eu from oxidation. We used substrate temperatures between 570 and 870 K. The deposition process was carried out in UHV systems with a base pressure prior to Eu deposition below 1×10^{-9} mbar not surpassing 3×10^{-9} mbar during deposition. For lower substrate temperatures incomplete intercalation takes place and part of the Eu stays on top of the hBN, see ESI† for details. The oxidation protection experiments consisted in exposing the sample to ambient pressure conditions (6 hours, room temperature, 80% humidity). The Eu thickness given throughout the manuscript corresponds to the values obtained for the calibration of the evaporator with a quartz microbalance at the measurement position. Since the sticking coefficient for both the quartz microbalance and the sample are unknown, but probably different, the indicated numbers can be compared with each other but are not suitable for absolute values.

3. Results and discussion

3.1. Formation of Eu-Pt surface alloy below hBN

3.1.1. Eu intercalation in the hBN/Pt(111) interface. The structural evolution of the hBN/Eu/Pt intercalated system can readily be monitored with low energy electron diffraction (LEED) experiments. LEED patterns in Fig. 1 correspond to the (111) plane on the Pt curved crystal. The clean Pt(111) pattern is shown in Fig. 1(a), which transforms, after borazine dosing at T = 1020 K, into the characteristic hBN (9 × 9) moiré of Fig. 1(b).⁵³ For successful Eu intercalation, there is a threshold temperature of the substrate of T = 770 K. Below this temperature part of the rare-earth material stays on top of the hBN coat, being subject to rapid contamination/oxidation (see

ESI†). At the lower range of the complete intercalation temperatures, right above 770 K, the LEED pattern only shows the progressive extinction of the hBN moiré. When rising the temperature to T=870 K a new $\approx(\sqrt{3}\times\sqrt{3})\text{R}30^\circ$ pattern emerges [Fig. 1(c)], with some weak satellite spots. A detailed inspection of the $\approx(\sqrt{3}\times\sqrt{3})$ structure reveals a $10/11\cdot(\sqrt{3}\times\sqrt{3})\text{R}30^\circ$ geometry with respect to Pt(111). This pattern reflects the presence of a EuPt₂/Pt(111) moiré-like coincidence lattice, similar to those found in rare earth RE–Au and RE–Ag surface alloys with RE–Au₂ and RE–Ag₂ composition. The ($\sqrt{3}\times\sqrt{3}$) ordering arises from the 1:2 Pt:Eu stoichiometry of the alloy at the local atomic-scale. The pre-factor emerges from the lattice mismatch of the 2D RE-noble metal alloy layer and the substrate.

(d)

The hBN/Eu/Pt(111) system is different from the Gr/Eu/Ir (111) one,³⁰ where the superstructure LEED spots belong to the graphene diffracted beams. In that case it was proposed that Eu forms a floating layer between the Ir(111) substrate and the graphene layer. In the here considered hBN/Eu/Pt system, however, the strongest LEED spots correspond to the EuPt₂ layer at the Pt interface. As indicated in Fig. 1(c), we can still detect extra satellite spots around the $10/11 \cdot (\sqrt{3} \times \sqrt{3})$ R30° structure, which arise from the coincidence lattice defined by the mismatched hBN/EuPt2 interface. Being all LEED structures properly identified, one can calculate real space lattice parameters out of the pattern. Taking into account the Pt lattice constant of a_{Pt} = 3.92 Å, we obtain the EuPt₂ coincidence lattice constant of $11 \cdot a_{Pt} / \sqrt{2} = 30.5$ Å, from which we deduce the EuPt₂ lattice parameter $a_{\text{EuPt}_2} = a_{\text{Pt}}$ $\sqrt{2.10/11.\sqrt{3}} = 5.29$ Å, with a nearest neighbor distance of 3.05 Å. The lattice mismatch of the EuPt₂ layer with the hBN lattice on top (2.504 Å) is quite large, but explains the 4.6×4.6 weak superstructure spots, marked by blue arrows in Fig. 1(c). After 6 hours of air exposure and a soft annealing to 570 K to remove air adsorbates the LEED pattern still reveals the 10/ $11 \cdot (\sqrt{3} \times \sqrt{3}) R30^{\circ}$ structure.

Bulk EuPt₂ exists and crystalizes in a MgCu₂ Laves phase structure, as shown in Fig. 2(a). A bulk lattice constant a_{Laves} between 7.64 and 7.73 Å has been reported. ^{43,44,61} However, in

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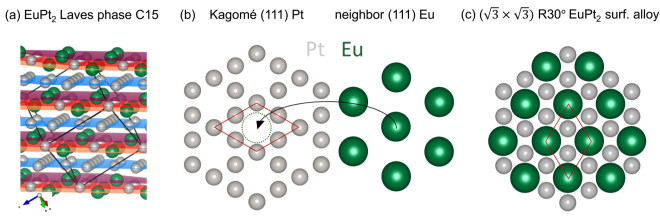


Fig. 2 Structural Model of the EuPt₂ surface. (a) Laves phase (C15) of bulk EuPt₂ in the MgCu₂ structure, (b) central Pt Kagomé (blue layer in (a)) and neighboring Eu(111) layer stacking inside bulk EuPt₂ that would result in a PtEu₃ composition with (2 × 2) superstructure after Eu incorporation, (c) $(\sqrt{3} \times \sqrt{3})$ R30° monolayer surface alloy structure below a 2D hBN layer.

such bulk EuPt₂ structure, and along the [111] direction, one cannot find any stoichiometric EuPt2 plane. Contiguous (111) planes contain either Pt and Eu solely, as shown in Fig. 2(b). The plane containing the Eu atoms is shifted by 3/8 of the densely packed Pt planes (approx. 0.8 Å). The fundamental Pt containing (111) plane is formed by a Kagomé lattice [blue layer in Fig. 2(a)], with the Eu atoms sitting above and below each Pt hexagon of the Kagomé lattice. Considering the Eu-Pt bilayer, this defines a (2×2) superstructure with a EuPt₃ composition. The here found EuPt₂ structure formed below hBN is therefore not related to bulk EuPt2 and can be understood by the simple incorporation of Eu atoms in the uppermost Pt (111) surface plane. Due to the larger size of the Eu atom, the interatomic distance at the surface increases, and a mismatch with the Pt(111) substrate underneath arises, leading to the $10/11 \cdot (\sqrt{3} \times \sqrt{3}) R30^{\circ}$ coincidence lattice.

The chemical characterization of the Eu intercalation process is carried out by means of X-ray photoemission spectroscopy. Results are shown in Fig. 3. In the bulk EuPt2 compound Eu atoms are in a di-valent Eu2+ configuration, while Eu atoms in compounds with higher Pt content become mixed-valent or completely tri-valent.44 The latter situation would be the case for Eu interstitial atoms, e.g., those that diffuse into the bulk and are surrounded by Pt completely. Fig. 3 reveals the Eu 3d core level for sub- and monolayer preparations at different temperatures. Submonolayer Eu deposition and intercalation at low substrate temperature (T =570 K) leads to a (nearly) complete divalent configuration while preparations at higher T result in the appearance of an additional Eu³⁺ signal. We interpret these observations as follows: at lower temperature only a partial Eu intercalation takes place, leading to the formation of EuPt2 patches below hBN (intercalated) and di-valent metallic Eu above hBN (not intercalated). For higher temperature, the Eu intercalation is complete, but together with the Eu²⁺ species of the EuPt₂ interface below hBN, Eu3+ component arises, which is ascribed to Eu interstitials in the Pt bulk, or simply to the buildup, under the topmost EuPt₂ patches, of EuPt_n (n > 2) alloys, giving rise

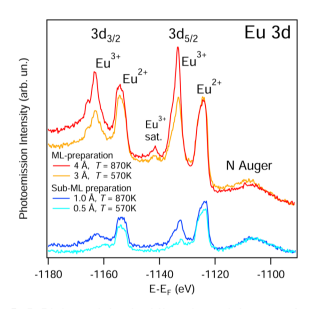


Fig. 3 Eu 3d photoemission signal. X-ray photoemission spectra for the Eu 3d edge taken at $h\nu=1486.6$ eV (Al $K_{\rm cl}$) for sub- and monolayer preparations at sample temperatures T=570 K and 870 K, respectively.

to a tri-valent or a mixed-valent situation. The Eu_2O_3 oxide has also Eu atoms in a tri-valent state. However, the presence of significant amounts of Eu_2O_3 are ruled out by the absence of an O 1s emission in XPS. The spectral region of the latter can be found in the $ESI.\dagger$ At higher coverage (>3 Å), preparations at both low and high temperatures already force extra Eu atoms to diffuse below the completed $EuPt_2$ layer, leading to similar di- and tri-valent contributions in the Eu 3d XPS spectra, as shown in the top part of Fig. 3. This tri-valent contribution for low-temperature preparation prove that a considerable part of Eu intercalates already at this temperature. Increasing the temperature at high coverage enhances the bulk diffusion, and leads to even stronger Eu^{3+} emission compared to Eu^{2+} .

3.1.2. Eu intercalation in vicinal hBN/Pt(111) interfaces. After characterizing the Eu intercalation below the hBN mono-

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layer on Pt(111), we focus on surfaces vicinal to the Pt(111) plane, investigated with the curved sample sketched in Fig. 4. The negative sign of the vicinal angle α corresponds to surfaces with A-type steps ({100} microfacets), and the positive to B-type step arrays ({111} microfacets).⁵¹ STM images in Fig. 4 correspond to three representative points of the curved substrate, namely the Pt(223) position ($\alpha = -11.5^{\circ}$), a low vicinal angle (α = -2.2°) close to Pt(111), and the Pt(554) surface ($\alpha = 5.8^{\circ}$). Prior to hBN growth, all vicinal surfaces exhibit well-ordered 1D step arrays, either at low and high vicinal angles.⁵¹ However, the hBN monolayer induces drastic structural changes, leading to a more complex nanoscale landscape. Close to the (111) position, large hBN/Pt(111) areas develop, which alternate with densely bunched steps. At larger vicinal angles the step bunching process remains, and the surface becomes a faceted structure. At the (554) position one observes a rather well ordered structure consisting of (111) terraces and side facets tilted at approx. 12°. At the (223) position one does not get a clear long range order. The latter situation is similar to stepped Ni crystals covered by hBN,62 while the ordered structures are rather close to the Rh case, where hBN growth on stepped surfaces leads to periodically arranged nanofacets. 63 One interesting question is whether the hBN forms a continuous, defect-free coat over the hill-and-valley structure underneath, since this requires "bending" of the hBN layer over Pt substrate facets. Due to step-bunching/faceting, the hBN monolayer must bend at the facet borders, i.e., step edges of the underlying Pt substrate must be overgrown by hBN to create a defect-free film. Such a defect-free connection of adjacent terraces has been shown for hBN growth on Cu(110)⁶⁴ and Cu(111).65 In the former case, the closure of the film was attributed to the fact that the Cu(110) step height s of s =

1.27 Å is smaller than the hBN bonding length (1.44 Å).66 However, for Cu(111) with s = 2.08 Å this explanation alone would not allow for "carpeting". Nevertheless, the edgedocking probability of the B6N7 seed to the different A- or B-type steps was observed to be different⁶⁵ resulting in a crystalline film. For Pt(111) with an even higher step height s =2.27 Å this connection ability is unknown. For a similar material, Ru(0001), a continuous film is difficult to achieve due to the intrinsic mismatch between individually nucleated h-BN domains on the same terrace as well as between adjacent terraces. 67 In general terms, however, one expects an increasing number of defects for an increasing presence of facet/step boundaries at large vicinal angles.

Eu deposition and intercalation on the vicinal surfaces changes the facet periodicity, size and inclination, as observed in the STM images. At the A-step type Pt(223) position, the rather disordered hBN/Pt(223) structure transforms into a well ordered array after Eu intercalation. On B-type steps, however, the Eu intercalation is leading to smaller facets. A statistical analysis of the STM images reveals an increasing average facet distance [(111) + side facet] of 20 facets per um close to (111), 25 facets per μm at (223), and 65 facets per μm at (554). This means that at A-type steps the surface rugosity decreases, contrary to B-type steps. On the other hand, XPS spectra (see below) reveal a similar Eu²⁺/Eu³⁺ relation on the different crystal positions, with a slightly higher di-valent amount at (111) compared to stepped surface planes.

3.2. Magnetism of the hBN/EuPt₂ system in the (111) plane

Divalent Eu $(4f^7, J = 7/2)$ has a low-temperature ferromagnetic state in bulk EuPt2 and similar compounds.68 We measured the magnetic properties of our hBN-protected EuPt2 surface

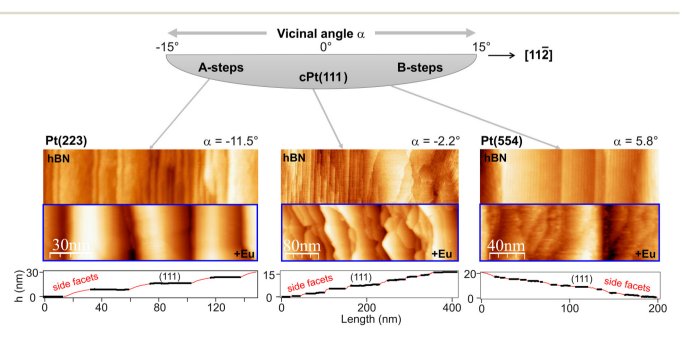


Fig. 4 STM images of the (Eu)/hBN/c-Pt(111) system. Large scale STM images of a hBN monolayer prior and after Eu intercalation collected at three different positions on the curved c-Pt(111) substrate (scanning parameters: I = 0.13 nA; U = 0.5 V). The line scans were taken close to the top of the images corresponding to the Eu intercalated systems. (111) and side facets are indicated by different color.

alloy with X-ray circular magnetic dichroism (XMCD) at the (111) position in the curved crystal. Results are shown in

Fig. 5. The X-ray absorption (XAS) spectrum in part (a) reveals a mixture of Eu2+ and Eu3+ contribution after Eu intercalation below the hBN/Pt(111) surface. This observation confirms the coexistence of the two Eu valences and it is consistent with the XPS results, namely di-valent Eu atoms at the EuPt2 surface and tri-valent Eu atoms diffused into the Pt bulk. The XMCD signal results from the difference of the absorption spectra of left and right circularly polarized light and is shown in the bottom of Fig. 5(a). At the applied field of 6 T, it shows the

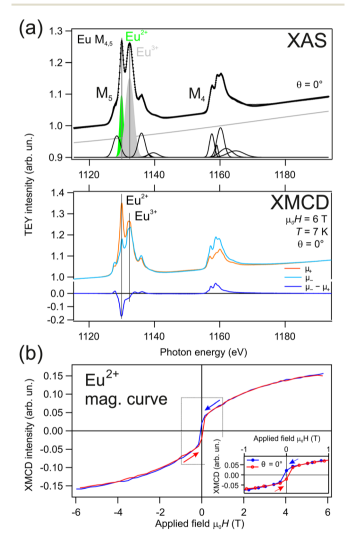


Fig. 5 Magnetic properties of intercalated Eu below hBN/Pt(111). (a) X-ray absorption spectrum (total electron yield - TEY) of horizontally (top, XAS) and circularly polarized light with opposite sign and its resulting difference spectrum shown below (XMCD) of 3 Å Eu intercalated below hBN/Pt(111) at a substrate temperature of 770 K. The XAS spectrum was fitted with several gaussian profiles (black, grey, green) and a linear background (grey line) for each of the two main Eu²⁺ and Eu³⁺ contributions. (b) Magnetization curve taken at the maximum of the Eu M_5 XMCD signal for a variable applied field from +6 T to -6 T (blue) and in the opposite direction (red), respectively. The sample was oriented perpendicular to the applied field at temperature T = 7 K. The inset presents a zoom-in to the small field region.

typical lineshape of pure di-valent Eu. 69 This is expected since, as stated before, Eu³⁺ has a 4f⁶ configuration with S = L = 3and J = 0, hence the Eu³⁺ signal should not contribute significantly to the anisotropy.

Ferromagnetism can be probed by measuring the Eu XMCD signal while varying the applied magnetic field. The XMCD signal is proportional to the magnetization M in the system. The resulting magnetization curve is shown in Fig. 5(b). It reveals a "S" shape-like behaviour with some jump close to zero-field. Ferromagnetism is confirmed by the small hysteresis opening, which is better observed in the inset of Fig. 5(b). The corresponding XMCD spectrum after removing the applied field (remanent state) is shown in the ESI.† Another method for determining the ferromagnetic state is the Arrott plot analysis, 70 which is also presented in the ESI.† For the inplane geometry, with the sample at 70° with respect to the magnetic field and light incidence, the magnetization curve is even more "S"-shaped at small fields, pointing to an out-ofplane easy axis (see ESI† for more details).

Exposure of the hBN/Eu/Pt system to air

Fig. 6(a) displays the complete evolution of the XPS spectra during the intercalation of 4 Å-Eu in the hBN/Pt(111) interface and after exposure to air. Prior to Eu intercalation, we obtain the characteristic shape and positions in the B 1s and N 1s core levels for the weakly-coupled hBN/Pt(111) system.⁵³ Individual contributions of the emissions can be extracted from line fittings, that are shown with further details in the ESI.† After Eu intercalation B 1s and N 1s core levels notably change their shape and energy, reflecting the fact that the hBN contact interface is now different. There are emissions from areas where the hBN layer is still contacting to Pt without Eu intercalation, giving rise to the right hand side emission, and areas for hBN on EuPt2, to which we assign the additional peaks that appear shifted to high binding energy. Such a shift is also observed for the hBN/Rh(111) interface, pointing to a similarly strongly interacting hBN/PtEu₂ interface.⁵³ With respect to the Eu core level, as mentioned above, it proves that Eu atoms are present in two configurations, di-valent Eu²⁺ for the 2D EuPt₂ alloy and tri-valent Eu³⁺ for Eu atoms incorporated into the Pt bulk below the EuPt₂ layer. After sample exposition to ambient conditions (6 hours, room temperature, 80% humidity) and a vacuum annealing to 770 K to remove at least part of impurities from the air exposure, the ratio of Eu²⁺/Eu³⁺ drops to one third. Note, that also the other core level intensities shrink. This relates to the appearance of adventitious C 1s (see ESI†) and O 1s emissions. The O 1s core level is detected at the higher binding energy side of the Pt 4p_{3/2} spectrum. A detailed view (inset) indicates a double-peak with energy positions of 530.5 eV and 532 eV, respectively. The former is in good agreement with the binding energy in the tri-valent Eu₂O₃ oxide. ^{71,72} The 532 eV emission is interpreted as due to hydroxide -OH, typical of Eu samples exposed to air. On the other hand, B 1s and N 1s core levels partially recover their shape and energy prior to Eu exposure. Especially the very strong interacting component of the hBN/EuPt2 areas disNanoscale Paper

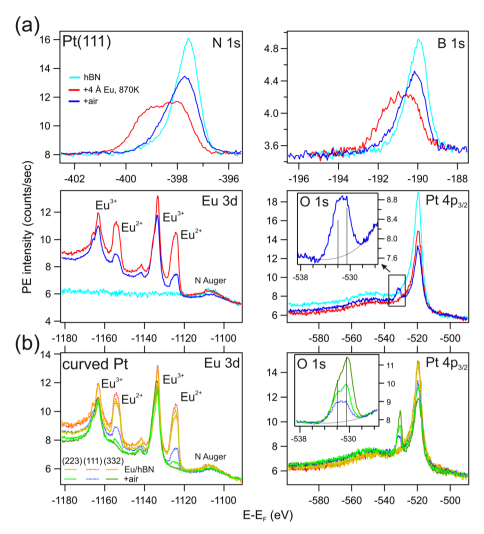


Fig. 6 XPS analysis of Eu intercalation below hBN on the curved Pt crystal. (a) N 1s, B 1s, Eu 3d, O 1s, and Pt $4p_{3/2}$ X-ray photoemission spectra taken at $h\nu = 1486.6$ eV (Al K_{c}) for the Eu 4 Å preparation before and after exposing to ambient conditions at Pt(111). (b) Comparative Eu 3d, O 1s, and Pt $4p_{3/2}$ core levels at three positions of the curved Pt substrate, at Pt(332), Pt(111) and Pt(335) positions, respectively.

appeared while the emissions of weak interacting hBN/Eu and hBN/Pt are preserved or even increased. The latter suggests that hBN areas on oxidized Eu have a similar core level emission as the hBN/Pt system, being both very low interacting interfaces. Another result from the XPS analysis is that oxidation of the hBN layer is not observed.

The hBN protection for the intercalated EuPt₂ alloy in vicinal Pt substrates is examined in Fig. 6(b), in direct comparison with the Pt(111) plane. Here we show the Eu 3d, Pt 4p_{3/2} and O 1s core levels at the Pt(332) and Pt(223) surfaces. At the Pt(332) plane, no di-valent signal remains in the Eu 3d spectrum after exposure to air. At the Pt(223) position the Eu²⁺/Eu³⁺ is strongly reduced, but the Eu²⁺ peak is still visible. Again, the O 1s spectrum suggests that the strong reduction of the number of di-valent Eu atoms is due to the formation of tri-valent Eu oxides and hydroxides. The overall O 1s intensity increases as the Eu²⁺ peak decreases at the stepped surfaces, although a more detailed peak analysis indicates that the intensity of the Eu-OH peak at 532 eV is similar in all three

cases, and it is the 530.5 eV peak from $\rm Eu_2O_3$ the one that scales reciprocally with the $\rm Eu^{2+}/Eu^{3+}$ ratio. The complete oxidation of the B-type (332) surface correlates with the high facet density observed in the STM analysis of Fig. 4, since this allows a higher number of hBN bending or breaks at facet borders. The rather small divalent signal that remains in Pt (223) may reflect the presence of larger (111) and side facets where the intercalated $\rm EuPt_2$ alloy remains better protected at ambient conditions.

Finally, we analyze the impact of the air exposure on the hBN protecting layer. For this purpose angle-resolved photoemission spectroscopy (ARPES) is particularly appropriate, since it is highly sensitive to the hBN valence band, as shown in Fig. 7. The photoemission intensity map of Fig. 7(a) corresponds to the hBN/Pt(111) interface (centre of the curved sample), and has been measured using the He II $_{\alpha}$ excitation energy ($h\nu$ = 40.8 eV). The strongly dispersing, intense features are the hBN π bands, with minimum at the $\bar{\Gamma}$ point of the Brillouin zone and maximum at \bar{K} . The emissions between

(a) (b) (c) (d)

hBN +1 Å Eu +5min. air +annealing 0

-2 - 1 - 2

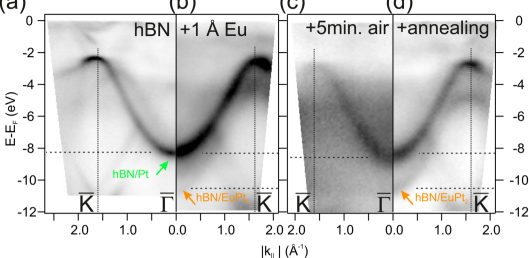


Fig. 7 Angle resolved Photoemission analysis of the hBN layer. He II $_{\alpha}$ ($h\nu$ = 40.8 eV) photoemission intensity maps along $\bar{\Gamma}\bar{K}$ direction of the hBN band structure for (a) hBN/Pt(111), (b) after intercalation of 1 Å of Eu at T = 770 K, (c) exposure of that sample to air, and (d) after another annealing to 770 K to desorb air contamination, respectively. The most intense features correspond to the π -band of hBN at the indicated interfaces.

approx. 2 and 8 eV binding energy entirely belong to hBN, while the Pt valence band features appear closer to the Fermi level. After intercalation of a very small amount of Eu (1 Å), the main π band appears unaltered, although a replica of this band emerges below, at 2 eV higher binding energy (Fig. 7(b)). The unaltered band corresponds to hBN π -band emissions of pure Pt areas without Eu while the shifted band arises in areas where the Eu intercalates. The band shift is explained by the enhanced interaction of the hBN with the EuPt2 substrate, with a net electron transfer from Eu atoms to the hBN layer similar to other systems. 40,73-75 After air exposure all bands get quite blurry, due to adsorbates. The adsorbate layer is disordered, which has the consequence that substrate and interface emissions are not well diffracted, decreasing the band intensity and increasing the background in second electron emissions. The dominating hBN π band is still visible, slightly shifted to higher binding energies. Interestingly, both the pure hBN/Pt and the Eu-intercalated bands can be recovered by annealing the sample again to 770 K, which removes adsorbates from the surface. First, this indicates that the oxygen exposure does not affect (oxidize) the hBN layer, which remains intact. Second, the photoemission intensity mapping still includes the higher binding energy replica, indicating that at least part of the strongly interacting hBN/EuPt2 patches remain, despite the oxidation of another part of the EuPt₂ layer as observed in the XPS. Third, above the oxidized part, the pi-band of the hBN layer is expected to present an energy position similar to the original hBN/Pt(111) π band position since the charge transfer from the metallic Eu has disappeared and the interaction at hBN/Pt or hBN/oxide interfaces is weak in both cases. This leads to two very similar π -bands that cannot be resolved. In the supporting material we present data from the $\bar{\Gamma}$ -point emission of a preparation of 4 Å Eu where this effect is better observed (see ESI†).

Conclusions

We have investigated the structural, magnetic and electronic properties of Eu after intercalation between a Pt substrate and a hBN monolayer. We used a Pt sample curved around the (111) direction in order to additionally assess the role of substrate steps. Our LEED analysis of the (111) interface shows a $\sim (\sqrt{3} \times \sqrt{3}) \text{R} 30^{\circ}$ pattern, revealing the presence of the EuPt₂ surface alloy under the hBN layer. We find that Eu atoms in this EuPt2 layer are divalent, while Eu atoms that have diffused further inside the Pt bulk during the intercalation process are trivalent. Interestingly, the Eu2+/Eu3+ ratio is not affected by the presence of steps at the Pt substrate. XMCD magnetization curves on the di-valent Eu atom reveal a ferromagnetic behavior. Air exposure of the sample leads to a partial protection of the divalent Eu atoms at the (111) plane, while at vicinal surfaces the protecting role of the hBN layer is less efficient, as reflected in the larger attenuation of the divalent Eu state. Such incomplete protection of vicinal planes may be related to a larger number of defects and domain boundaries in a more discontinuous hBN layer, since this covers a much rougher hill-and-valley faceted structure. This facilitates oxygen diffusion, intercalation and the EuPt2 alloy oxidation. In contrast, the hBN layer itself remains intact upon both Eu intercalation and air exposure.

Author contributions

A. M. I. B., K. A., and F. S. conceived and designed the work. A. M. I. B., K. A., A. A. M., I. P., F. B., L. F., and F. S. carried out structure and photoemission data collection while the data were analyzed and interpreted by A. M. I. B., K. A., and F. S. The magnetic measurements were performed

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by A. M. I. B., R. S., R. C., Y. H., L. F. and F. S. The article was drafted by A. M. I. B., J. E. O., and F. S. All authors contributed in the critical revision of the article and its final approval.

Conflicts of interest

There are no conflicts to declare.

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