Robust and tunable itinerant ferromagnetism at the silicon surface of the antiferromagnet GdRh$_2$Si$_2$

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Spin-polarized two-dimensional electron states (2DESs) at surfaces and interfaces of magnetically active materials attract immense interest because of the idea of exploiting fermion spins rather than charge in next generation electronics. Applying angle-resolved photoelectron spectroscopy, we show that the silicon surface of GdRh$_2$Si$_2$ bears two distinct 2DESs, one being a Shockley surface state, and the other a Dirac surface resonance. Both are subject to strong exchange interaction with the ordered 4f-moments lying underneath the Si-Rh-Si trilayer. The spin degeneracy of the Shockley state breaks down below ~90 K, and the splitting of the resulting subbands saturates upon cooling at values as high as ~185 meV. The spin splitting of the Dirac state becomes clearly visible around ~60 K, reaching a maximum of ~70 meV. An abrupt increase of surface magnetization at around the same temperature suggests that the Dirac state contributes significantly to the magnetic properties at the Si surface. We also show the possibility to tune the properties of 2DESs by depositing alkali metal atoms. The unique temperature-dependent ferromagnetic properties of the Si-terminated surface in GdRh$_2$Si$_2$ could be exploited when combined with functional adlayers deposited on top for which novel phenomena related to magnetism can be anticipated.

Silicon-terminated surfaces of crystalline solids are intrinsically part of conventional electronics, but their exploitation in novel materials combining two-dimensional electron states (2DESs) and magnetism, which play an important role in the development of next-generation electronics, still remains elusive. The appearance of 2DESs at surfaces or interfaces and their interplay with magnetic degrees of freedom may open an avenue for new physics in silicon-based technologies for future devices. In our world, a natural source of very strong magnetism is elemental Gd, which contains a half-filled 4f shell. Its ground state has a large pure spin moment $J = S = 7/2$ with vanishing orbital moment $L = 0$. Thus, in crystalline solids Gd will be insensitive to crystal-electric-field effects, which may strongly affect the magnetic properties of materials. In the present work, we consider the

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layered antiferromagnet GdRh$_2$Si$_2$, which crystallizes in the tetragonal body-centered ThCr$_2$Si$_2$ structure$^{4,5}$. Below the Néel temperature $T_N \sim 107$ K, the Gd 4$f$ moments become ferromagnetically ordered within the $ab$-plane, while they stack in antiferromagnetic (AFM) order along the $c$-axis$^{4,6}$. The Gd planes are well separated from each other by Si-Rh-Si trilayers. Similarly to other RERh$_2$Si$_2$ (RE = Yb, Eu) crystals$^{7,8}$, the chemical bonds within the trilayer are much stronger than those between the Gd and Si plane, therefore the surface of a cleaved GdRh$_2$Si$_2$ crystal can be terminated either by Si or Gd atoms. Silicon termination is particularly interesting, since in this case the first magnetically active layer of Gd is hidden and protected by the Si-Rh-Si buffer at the surface. This leaves us with the question, what happens to the two-dimensional electrons confined at the silicon surface, when the 4$f$ moments of the underlying Gd layer are ferromagnetically ordered, as it is schematically illustrated in Fig. 1.

As we will show below, the silicon surface of GdRh$_2$Si$_2$ reveals a remarkable property. Studying this system with angle-resolved photoelectron spectroscopy (ARPES)$^9$, we find two distinct 2DESs arising from Shockley and Dirac fermions, which reveal surface- and surface-resonant behavior, respectively. The surface electronic states theoretically predicted by I. Tamm$^{10}$ and later on by W. Shockley$^{11}$ were observed on the (111) surface of noble metals in ARPES measurements$^{12,13}$. These states, usually called Shockley states, lie exclusively in the bulk projected band gaps and in real space they are localized in a few surface atomic layers. Therefore they carry intrinsic quasi-2D electronic properties. In the recent past, Shockley states found on many metal and semiconductor surfaces attracted considerable attention due to their rich and exotic properties$^{9,14}$. Surface resonance states also have a quasi-2D nature, but in difference to the Shockley states they can penetrate into the material thus overlapping with the bulk band states. The wave function of the surface resonance in the bulk becomes essentially modified at the surface and is characterized by an enhanced probability density in the near surface region. Surface resonances could have notable $k_z$ dispersion$^8$ and can be considered as a kind of bridge connecting the properties at the surface and in the bulk of the material$^{9,15}$.

Here, we will focus on the linear-dispersive Dirac cone band, which appears at the $\Gamma$-point, and on a parabolic Shockley surface state, which can be seen within a large gap in the projected bulk bands around the $\bar{M}$-point i.e. at the corner of the surface Brillouin zone. Our ARPES study shows that when the Gd 4$f$ moments become ordered, both spectral structures are spin-split and form well-defined subbands. We have investigated the momentum-resolved temperature evolution of the spin splitting for both spectral features by ARPES and support our discussion of their origin and nature with theoretical calculations and X-ray magnetic linear dichroism (XMLD) measurements. We conclude that the two distinct 2DESs being an intrinsic signature of the Si-terminated surface of GdRh$_2$Si$_2$ exhibit itinerant magnetism at the surface. Their spin splitting arises from the strong exchange interaction with the ordered Gd 4$f$ moments lying below the Si-Rh-Si buffer. The temperature dependence of the spin splitting can be straightforwardly explained within the framework of conventional mean-field theory of the Heisenberg model$^{16}$. Our results suggest that the ferromagnetic Si-terminated surface of GdRh$_2$Si$_2$ can serve as a model substrate to induce non-trivial electronic and magnetic properties into nanostructures deposited on top which could eventually become interesting for technological applications. We present a simple example of tuning the electronic properties of the Si-terminated magnetic surface by depositing alkali metal atoms. As a result, strong energy shifts of the surface states due to the electron doping effect of the topmost layers as well as an interplay of spin-split bands of the Shockley state and surface resonances were observed, implying controllable modification of magnetism at the surface and sub-surface regions of the material. In the recent past, we have demonstrated that the Dirac fermion states can couple with ultra-heavy quasiparticles in crystalline...
4f-based systems by coupling to the electronic degree of freedom of 4f electrons. In this work, we show how the Dirac fermions can couple to the magnetic degree of freedom of 4f electrons.

Results

The coexistence of spin-polarized 2DESSs at the silicon surface from theory and ARPES. In Fig. 2a we show the surface electronic band structure obtained from our ab initio calculations for AFM ordered GdRh2Si2. To separate the surface-related electronic structure from bulk electron bands, we used a thick slab which was terminated by Gd on the one and Si on the other side. This allows us to trace simultaneously bulk-like bands, band gaps and surface-related states for both terminations. The 2D electron states that are related to the Si-terminated surface are shown in green, while the respective bands for the Gd-terminated surface are highlighted in red. In Fig. 2b we rather schematically illustrate the Fermi surface for the discussed 2DESs at the center of the Brillouin zone and at the point. Note that the lower of the two spin-split bands (1) of the Shockley state seen in (a) does not reach the Fermi energy along the direction. (c) Calculated spin-resolved electronic band structure for the Si-terminated GdRh2Si2 surface (the contribution of the topmost Si-Rh-Si block to the spin vector components is shown). Majority/minority bands are shown in red/blue. The spin-polarized 2DESs are labelled in accordance with Fig. 2a. (d) ARPES-derived Fermi surface for AFM ordered and Si-terminated GdRh2Si2 taken at a temperature of 1 K using 45 eV photons.

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When the crystal is terminated by a silicon layer, pairs of strongly dispersive bands appear in the large projected band gap around the point labelled as (1) and (2) (Fig. 2a,c). Looking closely on the behavior of these bands within the gap we can conclude that both the majority and minority states (Fig. 2c) are degenerate at the point and reveal a gap due to spin-orbit interaction. The observed spectral structure inside the gap arises from Shockley surface electron states confined within the topmost few atomic layers of the Si-terminated crystal (see next section). The essential point is that this surface state is missing at the Gd-terminated surface. In the experimentally derived Fermi surface (Fig. 2d), the respective feature can be seen as a diamond-like structure.
wrapping the Μ-point. Exploring further the spectral structure within the Μ-gap and along the Μ-Χ direction one can do a few more interesting observations. The electron-like spin-split lower subband of feature (1) does not reach the Fermi level and leaves the Μ-gap around 0.2 eV of binding energy (BE) (Fig. 2a,c). In comparison with the experimental data in Fig. 2d one may assume that this spectral feature may have a surface resonant behavior when it moves out of the Μ-gap and approaches the Χ-point. In the same region near the Χ-point another surface resonance band appears marked as (2*) which keeps its spin polarization up to the Χ-point. This band has been also detected in ARPES measurements which will be discussed below.

At the silicon surface, another remarkable feature labelled as (3) can be seen at the Γ-point, which apparently reveals a split pair of linear bands. The linear-dispersive conical shape corresponds to Dirac fermions and is imposed by the two-dimensional square symmetry of the layered crystal structure. A similar Dirac cone band has been seen in homologous systems like EuRh₂Si₂, YbRh₂Si₂ and YbCo₂Si₂ where its properties and interplay with the 4f states have been characterized.

Figure 3. Spin splitting of the Shockley state and Dirac cone. ARPES data taken from a Si-terminated GdRh₂Si₂ sample using 55 eV photons. The band maps were obtained near the Μ-point at 117 K (a) and 19 K (b) and near the Γ-point at 72 K (c) and 19 K (d). The measurements were performed along the Χ-Μ and Χ-Γ directions, respectively. The white vertical lines indicate the energy-distribution curves, which were further used for the analysis of the spin splitting in the T-dependent studies. The surface- and bulk-related spectral features are labelled in accordance to the theoretically derived bands seen in Fig. 2.
Thus, the results of our slab band structure calculations clearly point out the coexistence of two 2DESs at the silicon surface of GdRh$_2$Si$_2$. Moreover, our calculations suggest that in AFM ordered GdRh$_2$Si$_2$ spin degeneracy is lifted for both Shockley and Dirac cone states (Fig. 2c). In order to verify this prediction and to study how this phenomenon is related to the magnetic ordering in the system, we explored experimentally these electron states, their band splitting and their temperature dependence.

Surface-sensitive ARPES is ideally suited to explore the momentum-dependent electronic structure of a 2DES trapped at a crystal surface. In Fig. 3 we present the ARPES-derived spectral structure of a Si-terminated GdRh$_2$Si$_2$ surface as a function of temperature around the $\Gamma$- and the $\mathbf{M}$-points of the surface Brillouin zone. The bulk Néel temperature $T_N \sim 107$ K marks the onset of in-plane ferromagnetic alignment of the Gd 4f moments, which stack antiferromagnetically along the $c$-axis. We thus cleaved the crystal at 120 K well above $T_N$. Si termination can be easily identified by the presence of the intense Shockley surface state at the $\mathbf{M}$-point, which is absent for Gd termination. The high-temperature band map in the paramagnetic state is shown in Fig. 3a. Comparing the ARPES data to the results of our calculation, we can see that the Shockley state is unsplit. This changes dramatically when gradually cooling down the sample. To monitor the emergence of the spin splitting, we choose an energy distribution curve (EDC) at $k_{\parallel} \sim 1/3$ of the $\mathbf{X}$-$\mathbf{M}$ distance near the $\mathbf{M}$-point as indicated by the vertical line in Fig. 3b. With bare eyes, the appearance of the spin splitting becomes visible near $\sim 90$ K notably below the Néel temperature. Upon further cooling, the splitting becomes well resolved and its value rapidly increases, reaching $\sim 160$ meV. We have similarly followed the spectral structure of the Dirac cone at the $\Gamma$-point as a function of temperature, using the EDC at $k_\mathbf{0}$ marked by the vertical line in Fig. 3d. In this case, the band splitting becomes apparent only at temperatures below $\sim 70$ K. Figure 3c shows the spectral pattern at 72 K without evident spin splitting, whereas in Fig. 3d the splitting is nicely seen at 19 K with a maximum value of $\sim 70$ meV. A detailed analysis of the measured temperature dependence of the band splitting will be given below.

**Dirac cone and Shockley state: Temperature dependence of their spin splitting.** A deeper understanding of the properties of the discussed 2DEs can be obtained from the analysis of the orbital composition.
and visualization of the spatial extension of the calculated eigenstates. Representative electron densities of the Dirac cone near the $\Gamma$-point and the Shockley state at the $\Sigma$-point are shown in Fig. 4a,b, respectively. An analysis of the orbital character of the 2DESs reveals that the Shockley state is built up mainly by Si 3s and 3p (53%), Rh 4d (30%) and Gd 5d (14%) states. Our calculations suggest that the Shockley state is located exclusively within the first four atomic layers Si-Rh-Si-Gd and only 3% of this state penetrates in the vacuum. The Dirac cone is essentially built by Rh 4d $t_{2g}$ (74%) orbitals with an admixture of the Si 3px,3py states (23%) and a tiny Gd 5d (3%) contribution. Its conical structure is a consequence of the in-plane square symmetry of the layered material.8,17,18 The fourfold warping of the Dirac cone can be well recognized in its three-dimensional representation derived from our calculations (Fig. 4c). The theoretically derived Shockley state that is shown in Fig. 4d, is in excellent agreement with our ARPES results: the spin splitting of the electron- and hole-like bands as well as the appearance of the spin-orbit gap at the $\Sigma$-point are nicely reproduced.

Despite the 2D character of the Dirac cone, the electron density distributions in Fig. 4a,b show that its wave function penetrates deeper into the bulk than the Shockley state which is more strongly confined to the topmost four layers. In contrast to the Shockley state, the Dirac cone has the character of a surface resonance.9 Its wave function is resonantly enhanced at the surface but keeps a finite contribution in the bulk having therefore notable $k_z$ dispersion.8 We have recently explored this property in details for EuRh$_2$Si$_2$.8 The Shockley state decays exponentially into the crystal and is well localized at the surface region. We have verified that with increasing slab thickness, the orbitals from atoms below the first Gd layer and further in the bulk, mainly from Rh 4d, contribute to the Dirac cone, confirming its resonant nature. Moreover, the band structure in Fig. 2a,c shows that the Dirac cone essentially overlaps with the bulk bands, which appear as a bunch of gray lines in the slab calculation (Fig. 2a), where small hybridization gaps open up at the band crossings. However, both 2DESs show a spin splitting as a consequence of the exchange coupling to the Gd 4f moments. One may thus anticipate that the fundamental differences between the Dirac cone and the Shockley state substantially affect the strength of the exchange interaction with the ordered Gd 4f moments near the silicon surface and the temperature evolution of the splitting.

In Fig. 5, we present the evolution of the spin splitting for both 2DESs as a function of temperature. The splitting values have been deduced from fits of the energy distribution curves, which have been chosen slightly away from the Fermi level crossing and in the case of the $\Sigma$-point well away from the spin-orbit gap (vertical lines in Fig. 3b,d). A few exemplary EDCs and details about the fitting procedure can be found in Supplementary Fig. S1.
For the Shockley state, the splitting sets in at a temperature of \(\sim90\) K and rapidly approaches to a value of \(\sim160\) meV. At higher temperatures, the splitting seems to be much smaller than life-time and instrumental broadening and therefore cannot be resolved. Note that the monotonous increase of the splitting with decreasing temperature is interrupted by a “kink” around 60 K highlighted by a dotted circle, which will be discussed further. We found that the splitting of the Shockley state is actually anisotropic with values ranging from \(\sim160\) meV up to the largest observed value of 185 meV (inset in Fig. 5) in a direction parallel to \(\chi\) slightly away from the \(\text{M}\)-point.

The experimentally established spin splitting of the Shockley state is in agreement with the results of our theoretical studies. Interestingly, the averaged spin splitting of this state is larger than that observed for EuRh\(_2\)Si\(_2\) \((\sim150\) meV\(^2\)) but not as much as one might anticipate.

To model the temperature evolution of the Shockley state splitting, we fit the data with a magnetization curve obtained in the framework of the Weiss molecular-field approximation to the Heisenberg model\(^{16}\) (See Supplementary Note). For the localized Gd 4\(f\) moments, we assume a pure spin moment of \(J = S = \frac{7}{2}\). Neglecting the kink at 60 K, which we will discuss later on, the curve perfectly fits the ARPES data. This supports our finding from the band structure calculation, that the band splitting is mediated via exchange coupling to the localized Gd 4\(f\) moments. The fit confirms, that the splitting indeed vanishes at a temperature of \(\sim90\) K and saturates at \(\sim160\) meV at the considered point of the Brillouin zone. As we have already mentioned, the spin splitting of the Shockley state is highly anisotropic.

In Fig. 5 we also show the temperature-dependent spin splitting for the Dirac cone. Here, the splitting can be well resolved below \(\sim60\) K. The overall splitting is less than half as large as for the Shockley state. Therefore, similar to the Shockley state, an evaluation of the splitting remains inaccessible as soon as it drops below \(\sim50\) meV already slightly above 60 K, i.e. at even lower temperatures. Furthermore, as the Dirac cone has mainly Rh 4\(d\) character, spin-orbit coupling (SOC) has to be taken into account, which mixes with the exchange splitting. From our calculations, we estimate the spin-orbit splitting to be of the order of 40–45 meV for the Dirac cone close to the \(\Gamma\)-point, which is of the same order as the observed splitting around 60 K. As SOC does not depend on temperature, it might dominate the splitting at temperatures above \(\sim65\) K and even remain above \(T_N\). Ignoring all SOC effects, we nevertheless tried to fit the Dirac cone splitting in the same Weiss theory framework as the Shockley state and interestingly find the same critical temperature of \(\sim90\) K as in the previous case. Our findings imply that the spin splitting observed by ARPES is driven by the fundamental magnetic exchange interaction, which in the
reflect the magnetism of GdRh$_2$Si$_2$ in the bulk and, at least to a large part, at the Si-terminated surface. It is possible where the splitting of the resonant Dirac cone band has been detected. We therefore believe that the XMLD data is a surface resonance state penetrating deeper in the bulk. Thus, the Dirac cone state might open an additional channel for alignment between 4f moments at the surface and those deeper in the bulk has appeared along the c-axis. This is perfectly fine with our observation of the spin splitting onset of the Dirac cone, which is a surface resonance state penetrating deeper in the bulk. Thus, the Dirac cone state might open an additional exchange channel leading to an accelerated growth of macroscopic magnetic domains - ordered Gd 4f moments near the surface. In that regard we may propose that below ~60 K the magnetic systems at the surface and in the bulk become linked together and further growth of magnetic domains takes place simultaneously upon cooling at the surface and in the bulk. It should be noted that the XMLD signal is an average over aligned moments within the beam spot size $S_{\text{beam}}$ and thus also depends on the domain size $S_{\text{domain}}$ if $S_{\text{domain}} < S_{\text{beam}} = 100 \times 20 \, \mu\text{m}^2$. We believe that the dynamics of the domain growth and the role of the Dirac cone might be an interesting challenge for further studies on GdRh$_2$Si$_2$, a material which offers a rich playground for investigating the magnetic interplay between localized and itinerant electrons at the surface and in the bulk. One essential remark needs to be made about the XMLD experiment. In contrast to ARPES measurements, there is no direct evidence that during our XMLD measurements we have exclusively investigated the Si-terminated surface of GdRh$_2$Si$_2$. Nevertheless, the obtained results are meaningful and important when combined with the ARPES data. First, the T-dependence of the measured XMLD signals in TEY (surface) and TFY (bulk) modes are different, implying rather different magnetic properties near the surface and in the bulk. Second, the onset of magnetic order in the surface region and in the bulk of GdRh$_2$Si$_2$ by simultaneously looking at the total electron yield (TEY) and the total fluorescence yield (TFY) signal, respectively. The TEY signal probes less than ~20–30 Å near the surface with the main contribution coming from the first Gd layer. In TFY the probing depth is of the order of 100 nm and thus an almost pure bulk signal is seen. In Fig. 6 we plot the results of our XMLD measurements. The magnetic part of the linear dichroism was separated from the natural linear dichroism by subtracting the XLD signal measured at 140 K from all spectra. From the TFY data (open symbols), the onset of AFM order in the bulk can be seen at ~107 K in perfect agreement with the bulk Néel temperature determined from macroscopic measurements. However, from the TEY signal (solid symbols) it becomes evident that the magnetic order at the surface sets in substantially below the bulk Néel temperature - at around 90 K, which agrees well with the onset of the Shockley state splitting seen in ARPES (see Fig. 5). Furthermore, the TEY curve reveals an interesting kink-like feature near 60 K, like in the spin splitting curve derived from our ARPES measurements. From the difference of the TFY and TEY curves above ~60 K we may assume, that in this temperature range the magnetism at the surface is mainly mediated by the Shockley state, and is different from that in the bulk. Below ~60 K, the slope and shape of the TEY curve quickly approach those of the TFY curve. This effect might indicate that in this temperature range a new channel for alignment between 4f moments at the surface and those deeper in the bulk has appeared.

**Magnetic properties in the bulk and at the surface from XMLD.** To shed further light on the temperature-dependent spin splitting of the 2DEGs, which is directly linked to the ordering of the Gd 4f moments, we performed X-ray magnetic dichroism (XMLD) experiments at the Gd M$_4$ (3d $\rightarrow$ 4f) absorption edge. XMLD is sensitive to both FM and AFM order because the measured signal is proportional to the square of the ordered magnetic moment (M$^2$)\textsuperscript{24,25}. Using XMLD, we detected the onset of magnetic order in the surface region and in the bulk of GdRh$_2$Si$_2$ by simultaneously looking at the total electron yield (TEY) and the total fluorescence yield (TFY) signal, respectively. The TEY signal probes less than 20–30 Å near the surface with the main contribution coming from the first Gd layer. In TFY the probing depth is of the order of 100 nm and thus an almost pure bulk signal is seen. In Fig. 6 we plot the results of our XMLD measurements. The magnetic part of the linear dichroism was separated from the natural linear dichroism by subtracting the XLD signal measured at 140 K from all spectra. From the TFY data (open symbols), the onset of AFM order in the bulk can be seen at ~107 K in perfect agreement with the bulk Néel temperature determined from macroscopic measurements. However, from the TEY signal (solid symbols) it becomes evident that the magnetic order at the surface sets in substantially below the bulk Néel temperature - at around 90 K, which agrees well with the onset of the Shockley state splitting seen in ARPES (see Fig. 5). Furthermore, the TEY curve reveals an interesting kink-like feature near 60 K, like in the spin splitting curve derived from our ARPES measurements. From the difference of the TFY and TEY curves above ~60 K we may assume, that in this temperature range the magnetism at the surface is mainly mediated by the Shockley state, and is different from that in the bulk. Below ~60 K, the slope and shape of the TEY curve quickly approach those of the TFY curve. This effect might indicate that in this temperature range a new channel for alignment between 4f moments at the surface and those deeper in the bulk has appeared along the c-axis. This is perfectly fine with our observation of the spin splitting onset of the Dirac cone, which is a surface resonance state penetrating deeper in the bulk. Thus, the Dirac cone state might open an additional exchange channel leading to an accelerated growth of macroscopic magnetic domains - ordered Gd 4f moments near the surface. In that regard we may propose that below ~60 K the magnetic systems at the surface and in the bulk become linked together and further growth of magnetic domains takes place simultaneously upon cooling at the surface and in the bulk. It should be noted that the XMLD signal is an average over aligned moments within the beam spot size $S_{\text{beam}}$ and thus also depends on the domain size $S_{\text{domain}}$ if $S_{\text{domain}} < S_{\text{beam}} = 100 \times 20 \, \mu\text{m}^2$. We believe that the dynamics of the domain growth and the role of the Dirac cone might be an interesting challenge for further studies on GdRh$_2$Si$_2$, a material which offers a rich playground for investigating the magnetic interplay between localized and itinerant electrons at the surface and in the bulk. One essential remark needs to be made about the XMLD experiment. In contrast to ARPES measurements, there is no direct evidence that during our XMLD measurements we have exclusively investigated the Si-terminated surface of GdRh$_2$Si$_2$. Nevertheless, the obtained results are meaningful and important when combined with the ARPES data. First, the T-dependence of the measured XMLD signals in TEY (surface) and TFY (bulk) modes are different, implying rather different magnetic properties near the surface and in the bulk. Second, the onset of the TEY signal is seen at ~90 K which is in fine agreement with the onset of the splitting of the Shockley state of the Si-terminated surface seen in ARPES. Third, the “kink” seen in the T-dependence of the TEY signal appears at the same temperature of about 60 K, where the splitting of the resonant Dirac cone band has been detected. We therefore believe that the XMLD data reflect the magnetism of GdRh$_2$Si$_2$ in the bulk and, at least to a large part, at the Si-terminated surface. It is possible that within the 100 by 20 microns spot size there are also Gd-terminated parts of the surface. One may assume that those could contribute to the constant background seen in our experiment.

We now compare the essential differences in properties of EuRh$_2$Si$_2$ and GdRh$_2$Si$_2$. It is worth noting that in EuRh$_2$Si$_2$ the onset temperature of the Shockley state spin splitting ($T_s$)\textsuperscript{23} also differs from the bulk Néel temperature ($T_N = 24.5$ K), but in this case $T_s$ is more than 30% larger than $T_N$, while in GdRh$_2$Si$_2$ $T_s$ is ~16% below $T_N$. We propose that this difference between the two systems might arise from the competition between an enhanced...
in-plane exchange within the topmost Eu/Gd layer due to the strongly polarized Shockley states on the one hand, and a reduction of the overall magnetic coupling at the surface due to a reduced coordination number on the other hand. The former effect is likely of similar strength in both systems, since the spin splitting of the Shockley states is similar in both Eu- and Gd-based materials. On the other hand the “reduced coordination” effect is expected to be much stronger in the Gd- than in the Eu-based compound, because the exchange coupling between adjacent rare earth layers deep in the bulk is very strong in GdRh$_2$Si$_2$, whereas it is rather weak in EuRh$_2$Si$_2$. This is evidenced by the magnitude of the field required to get saturation magnetization, i.e. to rotate the ferromagnetic rare earth layers from AFM to FM stacking. This field is huge in GdRh$_2$Si$_2$, about 50 T$^4$, but tiny in EuRh$_2$Si$_2$, merely about 0.1 T applied along the basal plane$^{28}$. Thus EuRh$_2$Si$_2$ is already a quasi-two-dimensional magnetic system in the bulk, while GdRh$_2$Si$_2$ is truly three-dimensional. Altogether, at the surface of EuRh$_2$Si$_2$ the ordering temperature is only slightly pushed down by the removal of adjacent Eu layers, but strongly pushed up by the additional exchange, mediated through the Shockley states, thus $T_s > T_N$. In contrast, in GdRh$_2$Si$_2$, the removal of adjacent Gd layers pushes the ordering temperature significantly down, while this effect cannot be compensated by the additional in-plane exchange, thus $T_s < T_N$. The much stronger coupling along the $c$-direction in the Gd-based compound is likely due to the hybridization provided by the Gd 5$d$ electrons.

Our combined experimental and theoretical studies of GdRh$_2$Si$_2$ show that its silicon-terminated surface reveals rich and unique magnetic properties below ~90 K. It bears two distinct 2DESs arising from Shockley (surface state) and Dirac fermions (surface resonance state). Both are subject to strong exchange interaction with the hidden and magnetically active 4$f$ moments of Gd. This interaction lifts up the spin degeneracy of both electron states leading to the appearance of spin-split subbands with largest splitting values of 185 meV and 70 meV for the Shockley and Dirac state, respectively. Exploring the temperature evolution of the ordering of the 4$f$ moments at the surface we find that between ~90–60 K surface and bulk magnetism behave independently. Below ~60 K, the resonant Dirac cone state seems to link the surface and bulk magnetic subsystems. Evidently, this material still leaves plenty of unanswered questions and offers a rich playground for studying magnetic phenomena at silicon-terminated surfaces coupled to magnetically active overlayers. The surface magnetism of this compound might be further linked to a functional surface layer of organic molecules or ordered magnetic materials with metallic or semiconducting properties. To demonstrate this, we deposited potassium onto a freshly cleaved Si-terminated surface of GdRh$_2$Si$_2$. The respective data are shown in Fig. 7. As a result a systematic shift of the spin-split Shockley surface state to higher binding energies is observed that is caused by charge transfer from K into this state. Its spin splitting not only survives upon K deposition, but one can clearly see that the spin-split bands labelled as (1) strongly interfere and hybridize with the bands marked as (5) lying at the periphery of the $\overline{\text{M}}$-gap. Additionally, a new parabolic band appears at the $\overline{\text{M}}$-point and touches the Shockley state that is possibly derived from a respective unoccupied surface state of the clean Si-terminated surface (see Fig. 2). The observed surface electron doping effects together with the spin-derived hybridization lead to essential modifications of the 2DESs implying also changes of the magnetic properties at the topmost layers of the material. In the end we would like to add that previously, we have demonstrated that the Dirac fermion states can couple with ultra-heavy quasiparticles in crystalline 4$f$-based systems$^8$, i.e. couple with the electronic degree of freedom of 4$f$ electrons, while here, we show that the Dirac fermions may also couple via exchange to the magnetic degree of freedom of 4$f$ electrons.

**Methods Experiment.** ARPES studies were carried out at the Swiss Light Source (SIS X09LA instrument), the Diamond Light Source (I05 beamline) and BESSY-II (One-cubed ARPES instrument) and are described in details elsewhere$^{20,23}$. The ARPES spectra were acquired using a Scienta R4000 electron energy analyzer. The overall energy and angular resolutions were 10 meV and 0.1 degree, respectively. High quality single-crystalline samples of GdRh$_2$Si$_2$ were cleaved in situ in ultra-high vacuum at a base pressure better than $8 \times 10^{-11}$ mbar. Surface regions terminated by a Si layer were selected as the beam was scanning across the sample by looking at the ARPES map in the vicinity of the $\overline{\text{M}}$-point of the surface Brillouin zone, where the Shockley state settled in a gap of bulk-projected bands as a characteristic of Si termination was observed. The surface origin and two-dimensional character of the electron state at the $\overline{\text{M}}$-point were additionally confirmed by $h\nu$-dependent measurements, which indicate the absence of any dispersion along the $k_z$-direction. The beam spot size was set to 20 × 80 μm$^2$. The temperature-dependent measurements were always performed going from high to low temperatures in order to avoid fast sample aging.

XMLD experiments at the Gd $\text{M}_\text{I}$ edge were carried out at the European Synchrotron Radiation Facility (soft X-ray beamline ID32) using the high-field magnet end station$^{27}$. The samples were cleaved in UHV at 140 K in the high-field magnet immediately prior to the measurements. A small field of 100 mT was applied along the (110) axis of the crystal which was aligned to the E vector of the incident light for linear horizontal polarization. The temperature dependence of the XMLD was measured by slowly ramping the temperature down to 10 K and back up to 140 K and continuously measuring the absorption with linear horizontal (LH) and linear vertical (LV) polarization, simultaneously detecting the total electron yield and total fluorescence yield from the sample. The beam spot size at the sample was set to 100 × 20 μm$^2$.

**Theory.** As in the previous study$^{20,23}$, we used the generalized gradient approximation (GGA) to the exchange-correlation potential$^{33}$. The Hamiltonian contained scalar-relativistic corrections and spin-orbit coupling was taken into account by a second variation procedure$^{34}$. We set the energy cutoff for the plane-wave expansion of wave functions to 256.5 eV and sampled the two-dimensional Brillouin zone with a $12 \times 12 \times 1$ $k$-point grid. In order to correctly describe the Gd 4$f$ and Rh 4$d$ states, we used the GGA + $U$ approach$^{35,36}$. For the Gd 4$f$ electrons values of $U = 6.7$ eV and $J = 0.7$ eV were chosen, while the Rh 4$d$ electrons were treated with $U = 3.5$ eV and $J = 0.6$ eV. The GdRh$_2$Si$_2$ (001) surface was simulated by a 32-layer-thick asymmetric slab with the topmost (lowermost) surface terminated by Gd (Si).
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Author Contributions
C.K. and D.V.V. designed the research. K.K.I., N.C.-C., C.G. and C.K. prepared and characterized the samples for spectroscopic experiments. ARPES measurements were done by M.G., A.G., A.F., A.C., S.D., S.S. and D.V.V. XMCD experiments were performed by K.Ku. Theoretical studies were performed by M.M.O., E.V.C. and Y.M.K. Operation of the ARPES facilities was carried out by M.S. and M.R. at the Swiss Light Source and by P.D., T.K.K. and M.H. at Diamond, respectively. The obtained results were discussed together with C.L. The manuscript was written by C.K. and D.V.V. All authors have read and approved the decisive version of the manuscript.

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