Magnetism of Topological Boundary States Induced by Boron Substitution in Graphene Nanoribbons

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Graphene nanoribbons (GNRs), low-dimensional platforms for carbon-based electronics, show the promising perspective to also incorporate spin polarization in their conjugated electron system. However, magnetism in GNRs is generally associated with localized states around zigzag edges, difficult to fabricate and with high reactivity. Here we demonstrate that magnetism can also be induced away from physical GNR zigzag edges through atomically precise engineering topological defects in its interior. A pair of substitutional boron atoms inserted in the carbon backbone breaks the conjugation of their topological bands and builds two spin-polarized boundary states around them. The spin state was detected in electrical transport measurements through boron-substituted GNRs suspended between the tip and the sample of a scanning tunneling microscope. First-principle simulations find that boron pairs induce a spin 1, which is modified by tuning the spacing between pairs. Our results demonstrate a route to embed spin chains in GNRs, turning them into basic elements of spintronic devices.

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In spite of being a diamagnetic material, graphene can develop a special class of magnetism via the polarization of its \(\pi\)-electron cloud. Such \(\pi\) paramagnetism is less localized than the more conventional \(d\) or \(f\) magnetism, and can interact over longer distances. Magnetic graphene nanostructures thus offer promising perspectives for \(\text{à la carte}\) engineering of interacting spin systems with applications in quantum spintronics devices [1–4]. The vision of graphene \(\pi\) paramagnetism has been recently boosted by the development of on-surface synthesis (OSS) as a versatile bottom-up route. In OSS, nanoscale graphene flakes with customized shape and composition are fabricated over a metal substrate through the steered reactions between designed organic precursors [5,6]. Solid evidence of magnetism in flakes with zigzag edges has been revealed in scanning tunneling spectroscopy experiments [7–10].

Substituting one carbon atom of the graphene lattice by heteratoms is a potential route to induce magnetism [11,12]. A representative case is the doping of graphene with substitutional boron atoms, because it can be idealized as the removal of one electron from the conjugated bipartite lattice plus the energy upshift of a \(p_z\) state. However, boron atoms do not induce any spin imbalance in the graphene lattice, but simply behave as a point potential [11].

A prerequisite for the emergence of \(\pi\) paramagnetism is that the point defect also causes a sufficiently large rupture of the conjugated electron system, for example by completely removing lattice sites or saturating \(p_z\) orbitals [13–15], resulting in the localization of radical states.

Here we show that inserting a pair of boron atoms in the carbon lattice of graphene nanoribbons (GNRs) enables a magnetic ground state. Density Functional Theory (DFT) simulations shown in Fig. 1 reveal that a pair of

![FIG. 1. (a) Structure of the 2B-7AGNR shown over a color map representing the spin polarization density map, computed by density functional theory simulations [16] (green represents the boron moiety). (b) Spin-resolved projected density of states (PDOS) over carbon atoms around the boron dimer. A net spin polarization of one kind confirms the ferromagnetic alignment of the two magnetic moments.](image-url)
FIG. 2. (a) Organic precursors mixed in the experiments. (b) STM constant current topography image of a 2B-7AGNR ($V_b = -300$ mV, $I = 30$ pA). The green cross indicates the position from where the GNR is lifted. (right) Constant height current scan ($V_b = 2$ mV) using a CO-functionalized tip [38] of the region indicated by the dashed rectangle. (c) Tunneling current $I$ at $V_b = 25$ mV as a function of $z$ for a borylated (red) and a pristine (orange) GNR, for comparison. The grey region indicates where spectra in Fig. 3(b) were measured. The background shows results of atomistic simulations of a retraction stage shown in Fig. 4, for illustration.

substitutional B atoms in a 7-carbon-atom wide armchair GNR (7AGNR) can build up a net magnetic moment of $2\mu_B$ (two Bohr magnetons). This contrasts with the complete absence of magnetism when the pair is inserted in extended graphene [see Supplementary Material (SM) in [16]]. The spin polarization, shown in Fig. 1(a), decays towards the pristine segments with the characteristic shape of the 7AGNR end states [28] (SM, [16] for a comparison). In fact, the spin cloud emerges from the rupture of the conjugated system imposed by the borylated ring and the two neighboring Clar sextets [green in Fig. 1(a)]. This moiety behaves as a highly reflective barrier for valence band electrons [29,30], thus inducing localized end states associated with the termination of the topological 7AGNR valence band [31]. This striking result offers the vision of combining band topology of nanoribbons [31–33] and heteroatoms for shaping spin textures in graphene ribbons.

In our experiments, we substitutionally inserted boron pairs (2B) inside 7AGNRs (2B-7AGNRs) by adding a small fraction of 2B-doped trianthracene organic precursors [2 in Fig. 2(a)] [29,30,34–37] during the OSS of 7AGNRs using precursor 1 [5] [as schematically shown in Fig. 2(a), see Methods in SM [16]]. Scanning tunneling microscopy (STM) images of the fabricated ribbons [Fig. 2(b)] resolved the 2B unit as a topography depression at varying positions inside the GNR. Tunneling spectra showed no fingerprint of magnetism around the 2B moieties due to the strong interaction between boron and metal states [30,34,37], which quenches the eventual magnetic ground state. Therefore, to detect their intrinsic magnetic state the 2B moieties had to be removed from the metal substrate.

We used the STM tip to pick individual 2B-7AGNRs from one end [cross in Fig. 2(b)] and lift them off to lie freestanding between the tip and sample [7,39]. The (two-terminal) electrical transport through the suspended 2B-7AGNR was monitored during tip retraction $z$. At the initial stages of suspension (2B unit still on the surface), the current through the ribbon showed a weak exponential decrease with $z$ [Fig. 2(c)], as for pristine GNRs [39]. However, at a certain retraction length $z_p$, the current exhibited a pronounced peak, returning afterwards to the previous exponential decay. The peak and its position $z_p$ were reproduced for several retraction and approach cycles of the same ribbon, and appeared in all 2B-7AGNRs studied. In every case, the value of $z_p$ correlated with the distance between the 2B site and the contacted GNR end (see SM [16]), proving that the current peaks were caused by the detachment of a 2B moiety from the surface.

To explore the origin of the anomalous current peak, we measured differential conductance ($dI/dV$) spectra at positions around $z_p$ (Fig. 3). The $dI/dV$ plots show the emergence of a narrow zero-bias resonance in the spectra at $z_p = 2.7$ nm, which gradually decreases its amplitude with
tip retraction, and disappears for \( z > 2.9 \) nm. The resonance remained pinned at zero bias in all the \( z \) range observed. Its narrow line width reached a maximum value of \( \Gamma_{\text{HWHM}} \approx 6 \pm 0.4 \) mV at \( z_p \), and evolved nonmonotonously with retraction \( z \) [Fig. 3(a)] until disappearing. When the tip was approached below \( z_p \), the resonance vanished abruptly, but it was recovered by increasing \( z \) back above the \( z_p = 2.7 \) nm onset. From its narrow line shape and fixed zero-bias alignment, we conclude that the resonance is a manifestation of the Kondo effect \([40–42]\). A Kondo-derived resonance appears in \( dI/dV \) spectra when a spin polarized state weakly interacts with the conduction electrons of an underlying metal \([43]\).

To correlate these observations with the 2B-induced spin polarization predicted in Fig. 1, we performed DFT simulations of a finite 2B-7AGNR suspended between a model gold tip and a Au(111) slab \([16]\). Figure 4 shows the relaxed atomic structures of the GNR junctions before, while, and after detachment of a 2B unit, and includes the computed constant spin density isosurfaces. Before 2B-detachment from the surface \([ z < z_p \), Fig. 4(a)]\), the intrinsic magnetism around the 2B units is quenched; the projected density of states (PDOS) in the regions 1 and 2 around each boron atom is broad and spin unpolarized, contrasting with the clear spin polarization of free ribbons (shown as dashed plots). This is caused by the strong hybridization of the B atoms with the gold surface \([30,34,37]\), which appear 0.6 Å closer to the surface than the carbon backbone.

The detachment of the 2B moiety from the metal surface causes the emergence of a net spin polarization, clearly reflected in the PDOS and spin density isosurfaces [Figs. 4(b) and 4(c)]. At the intermediate snapshot of Fig. 4(b), only one of the two B heteroatoms is detached from the surface, and the ribbon hosts a net spin \( S = 1/2 \) extending towards the freestanding segment (region 4). For the fully detached 2B case [Fig. 4(c)], both regions around the two B atoms (regions 5 and 6) are spin polarized, recovering the \( S = 1 \) state of the isolated 2B-7AGNR [Fig. 1(b)]. Based on these simulations, we interpret that the most probable origin of the experimental Kondo resonance is the intermediate configuration pictured in Fig. 4(b). There, the Kondo effect is caused by the spin \( 1/2 \) of region 4 interacting weakly with the surface when the first boron atom is detached. Although the \( S = 1 \) state of Fig. 4(c) could also produce a Kondo state \([9]\), one would expect that it shows a larger extension and is accompanied by inelastic triplet-singlet side bands. Instead, the zero-bias resonance in the experiments disappears abruptly after a second kink \( \sim 2 \) Å higher [Figs. 2(c) and 3(b)] that we associate with the cleave of the second B atom, in consistency with the B-B distance. The experimental results are thus consistent with the spin polarization around freestanding 2B moieties.

Although the Kondo signal vanishes quickly with retraction, DFT finds that the \( S = 1 \) state of the free ribbon remains, and is clearly favored over an antiparallel alignment by \( \sim 14 \) meV per isolated 2B pair. The presence of a triplet state is striking; the two spin clouds at each side of the 2B center extend symmetrically over opposite sublattices of the 7AGNR, what usually favors an antiparallel kinetic exchange \([15]\). A detailed analysis reveals that the
hopping matrix elements between the two localized states at the sides of one 2B unit are very small ($t_{\text{intra}} \sim 18$ meV, see SM [16]). Consequently, the moiety formed by a 2B-doped ring surrounded by two Clar sextets is a very stable element that blocks conjugated electrons from hopping across. This explains the presence of a magnetic state because the borylated element acts as a barrier for valence band electrons of the 7AGNR segments [29], and induces spin polarized boundary states due to the nontrivial topology of this band [31]. Additionally, the 2B barrier also disconnects the boundary states at each side, and hinders the (antiparallel) kinetic exchange between them. The stabilization of the triplet configuration is then the result of the weak direct overlap between both spin-polarized boundary states through the 2B barrier, which, due to the tiny hopping between them, dominates the exchange interaction and induces the ferromagnetic alignment of the spins according to Hund’s rule.

We also studied GNRs with two consecutive 2B moieties like in Figs. 5(a) and 5(b), spaced by 1.2 nm. Transport experiments through these GNRs (i.e. (2B)$_2$-7AGNRs) as a function of tip-sample distance [Fig. 5(c)] also reveal deviations from an exponential decay with $z$, but now showing two peak features at retraction distances $z_{p1} \approx 1.60$ nm and $z_{p2} \approx 3.05$ nm. These values are related to the positions of the 2B units (nominally $\sim 2.0$ and $\sim 3.2$ nm from the contact point, respectively). A map of (normalized) differential conductance as a function of bias and $z$ [Fig. 5(d)] shows that both current features are also caused by narrow zero-bias $dI/dV$ resonances appearing at ranges $z < 1.9$ nm and $2.9$ nm < $z < 3.5$ nm, respectively. Their line shape [Fig. 5(e)] is similar to the resonances observed for the single 2B case, and can also be attributed to Kondo states, which reflect the emergence of spin polarization in the ribbon as each 2B unit is detached from the surface.

Although these results apparently suggest that each 2B behaves as an independent spin center, DFT simulations of free (2B)$_2$-7AGNRs [Fig. 5(b)] find that the two singly occupied boundary states between neighboring 2B elements interact strongly and open a large hybridization gap [30], forcing them into a closed shell configuration. As a consequence, the spin polarization vanishes between two 2B sites, but persists outside this region as two uncompensated spin 1/2 clouds [Fig. 5(b)] with barely no preferred relative spin alignment. From our electronic structure calculations [16,30] we can characterize this hybridization by a relatively large effective hopping term $t_{\text{inter}}$ between boundary states of neighboring 2B units, which contrasts with the weak hopping $t_{\text{intra}}$ across each 2B unit. In fact, the electronic structure close to the Fermi level of a sequence of borylated units can be mapped onto the Su-Schrieffer-Heeger (SSH) model [45], characterized by two alternating hoppings along a 1D wire. Since $t_{\text{inter}} > t_{\text{intra}}$, an alternative way to understand the spin-polarized states in Fig. 5(b) is as zero-energy topological modes of a very short SSH chain. These simulations allow us to predict that a $S = 1$ spin chain will emerge for larger inter-2B spacing, when both hopping terms become smaller than the Coulomb charging energy $U$ of the boundary states [16].

To explore if inter-2B interactions survive in the experimental geometry, we simulated a (2B)$_2$-7AGNR suspended between a tip and sample. Figure 5(f) shows the spin polarization of a snapshot with one 2B moiety completely detached and the second partially bound to the surface. In contrast with the large spin cloud around the lifted single 2B in Fig. 4(c), here there is no spin density between the two 2B moieties, but a net $S = 1/2$ cloud. 

FIG. 5. (a) (Left) Constant current STM image of a borylated GNR ($V_b = -300$ mV, $I = 30$ pA). (Right) Constant height current image of the marked rectangular region ($V_b = 2$ mV) using a CO-functionalized tip. (b) DFT simulation of the magnetization of a (2B)$_2$-7AGNR. (c) Cotunneling current $I$ vs $z$ through the ribbon in (a) suspended between tip and sample. (d) Normalized differential conductance of the suspended (2B)$_2$-7AGNR as a function of $V_b$ and $z$ (see SM [16]). Two zero-bias resonances are observed. (e) Representative $dI/dV$ spectra measured at the indicated $z$ positions, with fits (dashed) using Frota functions [44]. (f) DFT relaxed structure of a suspended (2B)$_2$-7AGNR. Constant spin-density isosurfaces are shown over the atomic structure; red arrows indicate the position of the B atoms. The inset shows the indicated region of the suspended GNR from a different angle.
above, confirming the presence of interactions between 2B units. The second $S = 1/2$ boundary cloud is expected to appear below the lower 2B only after the last boron atom is detached, this being responsible for the more extended Kondo effect observed in the experiment above $z_{p2}$. These results confirm the spin polarization predicted at the interface between (2B)$_2$ units and pristine 7AGNR segments [31], which in essence are zero-energy modes of the 7AGNR valence band of similar nature than those created by a single 2B unit at every side.

The peculiar spin polarization of single 2B units and dimers is a remarkable consequence of the large and long-range exchange interactions present in GNRs [7,8,31]. Our results support that spins survive in freestanding GNRs and can form $S = 1$ spin chains for low concentrations. Tuning the separation between 2B units is a promising strategy to control spin polarization through a change in the correspondence between inter-2B and intra-2B interactions. Furthermore, the stronger sensitivity of substitutional boron heteroatoms to chemical bonding endows these systems with ideal properties to manipulate complex spin states in chains.

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Magnetism of Topological Boundary States Induced by Boron Substitution in Graphene Nanoribbons

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I. MATERIALS AND METHODS

A. Sample preparation

To prepare the Au(111) substrate we perform a cleaning cycle by sputtering the crystal with Neon ions for 10 minutes and afterwards annealing the crystal in UHV conditions \([ p < 10^{-9} \text{ mbar}]\) at \(T = 740 \text{ K}\) for 15 minutes. We prepare our samples by simultaneously evaporating DBBA and B$_2$-DBTA molecular precursors on the clean Au(111) surface [1–3]. Afterwards an Ullmann reaction and cyclodhydrogenation are induced by successive annealing the sample to 200°C for few minutes and flashing to 400°C for 30 seconds.

The samples are prepared in situ and transferred directly into a homebuild STM kept with liquid Helium at \(\approx 6 \text{ K}\). We use a PtIr tip with an atomically sharp gold termination. The tip is prepared by controlled indentation into the clean Au(111) surface.

B. Lifting procedure

The lifting is performed as following. After stabilization of the tip above the clean Au(111) surface with a fixed set of parameters \([I = 30 \text{ pA}, V_b = -1000 \text{ mV}]\), we open the feedback loop of the control electronics. Afterwards, we position the tip above the termination of the GNR and approach towards the surface with a bias \(V_b = 25 \text{ mV}\). At some point during the approach a sudden increase in the current is observed. This corresponds to the formation of a tip-molecule bond and defines \(z = 0\) during each experiment. After the tip-molecule bond is formed we are able to partially lift the GNR from the surface.

Custom cleaving trajectories are employed for the different molecules, depending on the strength of the tip-molecule bond. In the most robust cases we cleave the molecule by simple retraction along the \(z\)-axis. In less strongly bound cases, we need to move additionally in the \(xy\)-plane, following a trajectory along the direction of the GNR-backbone on the surface. However, we find that variations in the lifting trajectory do not influence at which \(z_p\) we observe the peak in \(I\) and the Kondo-resonance in the differential conductance spectra.

C. Normalization of differential conductance spectra

Each individual spectrum taken at one \(z_0\) was normalized following \(G_{\text{norm}}(V, z_0) = \frac{G(V, z_0) - G_{\text{min}, z_0}}{G_{\text{max}, z_0} - G_{\text{min}, z_0}}\), where \(G(V, z_0)\) is the measured \(dI/dV\)-signal at \(z_0\) and \(G_{\text{min}, z_0} [G_{\text{max}, z_0}]\) is the minimal [maximal] value of \(G(V, z_0)\) in the analysed bias interval. The normalization is only applied to the data presented in Fig. 5(d) of the main text.

D. Presentation of experimental data

The figures presenting experimental data were prepared using WSxM [4] and the python matplotlib library [5], perceptual continuous color scales for color maps are based on [6].
E. Details on ab-initio calculations

First-principles electronic structure calculations are performed using density functional theory as implemented in the SIESTA package [7, 8], where the valence-electron wave functions are expanded using a linear combination of numerical atomic-orbitals as a basis set and the core electrons are replaced by norm-conserving Troullier-Martins pseudopotentials [9]. A double-ζ plus polarization (DZP) basis set is adopted for the surface Au atoms, where an extended basis is considered for the top atomic layer optimized for the description of the Au(111) surface [10]. The tip model is defined by a gold rod with its axis along the Au(100) crystalline direction and its valence-electrons described using a DZP basis set for s-orbitals and single-ζ non-polarized (SZ) basis for d-orbitals (i.e. DZP-SZd). Such geometry and basis set has been shown to provide a reasonable description of the contact, with a smooth local density of states at the tip apex around the Fermi level [11]. For the atoms defining the borylated ribbons a double-ζ non-polarized (DZ) basis set is employed, where the orbital radii are defined using a 30 meV energy shift [8]. Dispersive interactions between the borylated ribbon and the metallic surface are considered using the van der Waals density functional by Dion et al. [12] with the modified exchange by Klimeš, Bowler and Michaelides [13]. The real-space grid for integrations is defined using a 300 Ry energy cutoff [8]. The smearing of the electronic occupations is defined by an electronic temperature of 300 K with a Fermi-Dirac distribution. The self-consistency cycle is stopped when variations on the elements of the density matrix are lower than $10^{-5}$ as well as $10^{-5}$ eV for the Hamiltonian matrix elements.

The freestanding 2B-GNRs, as the one shown in Fig. 2A in the main text, are computed within a simulation cell that includes 20 and 40 Å of empty space in the directions along and perpendicular to the ribbon, respectively, to avoid interactions with the periodic replicas. Geometry optimizations are performed using the conjugate gradient (CG) method until all forces are lower than 20 meV/Å. For the calculations involving the suspended borylated GNRs, the Au(111) surface is represented by a 3-layer thick slab within a simulation cell of dimensions $46.16 \times 24.98 \text{ Å}^2$ along the periodic directions. A H passivation is employed at the reverse side of the slab in order to prevent spurious effects due to interaction between surface states belonging to the top and bottom surfaces of the slab [14]. A $1 \times 2 k$-point mesh is used to sample the bidimensional Brillouin zone. The slab defined in this way comprises 640 atoms [7680 orbitals], where the atoms from top Au layer are allowed to relax with the CG method until forces are lower than 20 meV/Å (the H passivation and the two bottom Au layers are kept fixed).
II. SUPPLEMENTARY TEXT

A. Boron doping in extended graphene

In Fig. 1 we present the spin polarized DOS obtained from DFT simulations on extended graphene monolayers with boron doping. Inserting a single substitutional boron atom into the graphene lattice effectively removes one electron from the bipartite lattice. The defect acts as a weak point potential, but does not induce any spin imbalance in the surrounding graphene sub-lattice (Fig. 1a). This is also the case when a 2B moiety is inserted in the extended graphene monolayer (Fig. 1b).

These results contrast with the substitutional insertion of the same 2B moiety into a 7AGNR. As shown in Fig. 1 of the main text, this induces a net spin polarization due to the rupture of the conjugation of the valence band.

FIG. 1: Boron doping in a graphene monolayer: Spin-polarized, projected density of states (PDOS) on C atoms in the region around (a) one substitutional B atom in graphene, and (b) a pair of B atoms in opposite sublattices of the graphene (shown in the drawings on top). In every case, there is not net spin polarization in the system.
B. Lift of GNRs with 2B sites at different positions

In Fig. 2 we present two further examples of $I$-$z$ retraction curves of two 2B-GNRs with the 2B site at 2.0 nm and 3.6 nm from the contact point, respectively. The peaks in the current are observed at $z_p = 1.2$ nm and $z_p = 2.4$ nm, in agreement with the different 2B to contact point distances. In both cases the peak in the current is caused by a zero-bias Kondo resonance, like in the case presented in the main text.

FIG. 2: **Fingerprint of magnetism in borylated GNRs**: (a), (b) STM constant current topography scan of two borylated 7AGNRs [$V_b = -300$ mV, $I = 30$ pA] with the doping position at different distances from the lifting position. The green cross indicates the position from where the GNR is lifted. Scale bar is 3 nm. (c) Cotunnelling current $I$ as a function of $z$ for the GNRs depicted in (a) and (b). The value of $z$ at which the current peak is observed correlates with the position of the doping site in the ribbon. The lifting experiment is performed at $V_b = 25$ mV. (d), (e) Spectrum over a larger bias window, taken at $z = 1.4$ nm ($z = 2.4$ nm) with GNR 1 [GNR 2] bridging tip and sample. The dotted grey lines are Frota functions, obtained by fitting to the experimental data, with HWHM = 5.0 ± 0.3 mV [(d), GNR 1] and HWHM = 5.7 ± 0.3 mV [(e), GNR 2].
C. Evolution of the Kondo resonance line width with retraction distances $z$

Figure 3 shows full width at half maximum (FWHM) and amplitude ($A$) of the Kondo resonances shown in Fig. 3 of the main manuscript. These are obtained by fitting a Frota function [15] plus an offset to the spectra. For $z \geq 2.92 \text{ nm}$ the Kondo resonance vanished and $A$ dropped approximately by one order of magnitude. The Frota function was fitted to the data in the interval of $|V_b| < 20 \text{ mV}$. We tested the stability of this procedure by increasing the size of the interval up to $|V_b| < 50 \text{ mV}$. No qualitative changes and only minor quantitative changes of the FWHM were observed. $A$ was unaffected by the interval’s size.

D. DFT simulations of the edge states and boron-induced inner states in 7AGNR

The boundary states induced by the substitutional boron pairs exhibit the same shape, symmetry, and decay towards the pristine segments as the 7AGNRs end states, localized in the zigzag terminations. Fig. 4 below presents a comparison between two 7AGNRs end states facing each other [shown in the top] and the 2B-induced bonding and anti-bonding states [bottom]. Notice that, interestingly, for 2B centers the “bonding” state is higher in energy than the “anti-bonding” linear combination.

![Wavefunction of edge states and 2B-induced states in 2B-7AGNRs](image-url)

FIG. 4: Wavefunction of edge states and 2B-induced states in 2B-7AGNRs: Bonding (a) and anti-bonding (b) boundary states localized around the boron dimer [bottom] as compared to two 7AGNR edge states [top]. Red arrows indicate the position of the boron atoms.
E. Electronic structure of 2B-7AGNR on lift-off configuration versus free standing

FIG. 5: Density of states projected on the C atoms within the boxed regions shown in Fig. 4 of the main text, here displayed in a larger energy window: solid lines are related to the configurations where a 2B-7AGNR is bridging a gold tip and a Au(111) surface [Fig. 4, main text], whereas the dashed lines are taken from a corresponding free standing 2B-7AGNR. Panels (a)-(c) corresponds to the three different configurations in Fig. 4 of the main text.

The density of states projected on the C atoms around each B atom (PDOS), inside the marked regions in Fig. 4 of the main text, are shown here in Fig. 5 for a wider energy window. Before detachment, when the B atoms are close to the Au substrate [regions 1, 2, and 3], besides having the magnetism quenched, the hybridization with the substrate is reflected in the larger broadening of the PDOS at lower energies. This is in accordance with previously reported results for fully borylated 7AGNR on Au(111) surface.[3] For the detached regions [4, 5, and 6], the intrinsic magnetism near the B atom is restored and the PDOS resemble the free standing ones [dashed lines]. The small misalignment between the solid and dashed lines in these cases is due to the Fermi energy on the free standing 2B-7AGNR being arbitrarily defined somewhere inside the gap.
F. Minimal model of an isolated 2B center in 7AGNR

**FIG. 6:** Minimal model of an isolated 2B center in a 7AGNR: Two boundary states L and R, one at each side of the 2B pair and characterized by a charging energy $U$ (Hubbard parameter), are connected by a hopping matrix element $t_{\text{intra}}$. We also add a Coulomb exchange interaction $J_{\text{intra}}$ between electrons in different levels. The corresponding effective Hamiltonian, expressed in terms of spin $\hat{S}$ and occupation $\hat{n}$ operators at each site, is shown above in the relevant limit here, $U \gg t_{\text{intra}}$.

A single 2B moiety in a 7AGNR is found to build up a total spin $S=1$ due to the predominant ferromagnetic interaction between the two singly occupied boundary states [B-states] localized at each side. The purpose of this section is to formulate the minimal model [depicted in Fig. 6] capable of explaining the low-energy electronic structure and the magnetism of isolated 2B pairs.

The effective Hamiltonian describing an isolated 2B pair also can be seen in Fig. 6 in the limit $U \gg t_{\text{intra}}$. Exact and mean field solutions of this model are readily obtained and can be compared with the relative energies and the band structures of different DFT solutions. In particular, we use information from DFT simulations [both non-spin-polarized and spin-polarized with either ferromagnetic or anti-ferromagnetic spin alignment] to estimate the following values of the effective parameters describing the system according to the model of Fig. 6: the charging energy of each boundary state $U \sim 260$ meV, the effective hopping matrix element between both boundary states $t_{\text{intra}} \sim 18$ meV, and the Coulomb exchange interaction $J_{\text{intra}} \sim 17$ meV.

While the Coulomb exchange $J_{\text{intra}}$ favours parallel alignment, the kinetic exchange (KE) term, given by $4(\frac{t_{\text{intra}}^2}{U})$ [Fig. 6] tends to align spins antiferromagnetically. The small value of $t_{\text{intra}}$ relative to $U$ translates into a small KE interaction, of the order of $\sim 1.2$ meV, i.e., around 14 times smaller than the Coulomb exchange within each 2B pair. Therefore, the ground state of an isolated 2B pair corresponds to a total spin $S=1$ as expected from Hund’s rules.
G. Electronic and magnetic interactions between neighbouring 2B centers

We now consider the case of periodic doping of 7AGNRs with 2B centers. The distance between two 2B sites is given by the number \( n \) of 7AGNR unit cells defining the periodicity of the system. Fig. 7 compares the computed hopping matrix elements between B-states at each side of a given 2B pair \( (t_{\text{intra}}) \) and those between two neighbour 2B pairs \( (t_{\text{inter}}) \). These values are obtained by fitting to the Su-Schrieffer-Heeger (SSH) model [16] the B-derived in-gap states appearing in the non-spin-polarized DFT band structures in Fig. 5 of Ref.[3] for 2B-(7AGNR)\(_n\) with \( n = 3, \ldots, 13 \). The fitting is rather accurate for \( n > 3 \), while for \( n=3 \) there is a somewhat larger uncertainty due to the interaction between the lower B-derived state and the valence band of the 7AGNR near the \( \Gamma \)-point. As mentioned above we obtain a very small effective hopping parameter between the boundary states across the diboron

![Diagram of electronic interactions between B-derived states from DFT calculations](image)

**FIG. 7:** Electronic interactions between B-derived states from DFT calculations: (a) Schematic model of a 2B-AGNR with intra-2B and inter-2B hopping terms indicated. \( J_{\text{intra}} \) represents the direct Coulomb exchange resulting in ferromagnetic alignment of the spins within each pair. (b) Computed values of intra-2B and inter-2B hopping terms from the analysis of non-spin-polarized DFT band structures of the periodic systems [3]. The inter-2B spacing given by \( n \), the number of 7AGNR unit cells between 2B sites. The correspondence with the Su-Schrieffer-Heeger (SSH) model is evident. The inset shows the exponential decay of \( t_{\text{inter}} \) and magnifies the crossing point where the topology of the non-spin-polarized band structure of the system changes. However, as discussed in the text, the non-polarized solution is not relevant in the limit where \( t_{\text{inter}} < U \) that clearly holds for \( n \geq 5 \).
 moiety $t_{\text{intra}}$, which amounts to 18 meV for $n \geq 5$.

The effective electron hopping between proximal B states $t_{\text{inter}}$ is much larger than $t_{\text{intra}}$ for $n < 7$, and decreases exponentially with the separation, resulting residual above $n \geq 7$. The spacing between diboron sites in the results in Fig. 5 of the main text corresponds to $n = 3$. The inter-diboron hopping is very large in this case [$\sim 350 \text{meV} > U$], resulting in a closed-shell state with no spin polarization in the region between them. The small electron hopping across a 2B site protects the outer B states, which each maintain a net spin polarization of $S=1/2$, as detected in Fig. 5.

If the larger hopping parameter is the hopping from one site to its neighbour, i.e. $t_{\text{inter}} > t_{\text{intra}}$, the system resembles a non-trivial bulk-boundary correspondence, as described by the SSH model. According to this model, for the smallest spacings between diborons [$n=3, 5$] boundary states are expected at the terminations since $t_{\text{intra}} < t_{\text{inter}}$. However, if we consider Coulomb correlations, only the case $n=3$ shows end states because it fulfils $U < t_{\text{inter}}$. The $n=5$ case, for which already $U > t_{\text{inter}}$, results in a chain with a spin texture formed by $S=1$ spins localized at the 2B sites and aligned antiferromagnetically along the chain, due to the large kinetic exchange associated with $t_{\text{inter}}$.

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