Few-layer antimonene: electrical properties

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1. Introduction

Recent works have shown how to produce antimonene and few-layer (FL) antimonene with a variety of different procedures [1–6], allowing the exploration of its promising properties [7–11], including its application in optoelectronics [12–15]. One of the properties making antimonene unique among other 2D materials is its strong spin–orbit coupling (SOC). Elemental bulk antimony is a topological semimetal on account of an inverted bulk band order [16]. On the other hand, a band gap in the range of 0.76–2.28 eV has been predicted for single layer antimonene [17–23], which makes it very attractive for electronics and ultrafast optoelectronic applications. The electronic properties of antimonene change quite drastically from the single layer to the FL regime. Although antimony does not present a bulk gap, its nonzero topological invariant guarantees the presence of topologically protected surface states (TPSS), coexisting with bulk bands at the Fermi energy [24–26]. For a small number of atomic layers, antimonene could already behave as a 3D topological insulator because quantum confinement opens a gap in its bulk bands [27], but, when this occurs, the TPSS on opposite surfaces couple to each other and a gap opens at the Dirac point, partly degrading their topological properties. A minimum of approximately 7–8 layers (∼ 3 nm) is needed for a full decoupling of the TPSS on opposite surfaces [25,28], but bulk bands already cross the Fermi energy at this small thickness [26,27]. In this sense, FL antimonene is not so different from actual 3D topological insulators where the contribution of bulk bands at the Fermi energy is difficult to eliminate. Angle Resolved Photoemission Spectroscopy (ARPES) [9,25,26] measurements and, to a much lesser extent, transport experiments [27,28]...
have confirmed this issue in binary compounds such as Bi$_{1-x}$Sb$_x$, Bi$_2$Se$_3$ or Bi$_2$Te$_3$ [26,29,30]. Ternary [31] and quaternary [32] compounds have also shown surface contribution in their conduction properties. Since the surface/bulk conductivity ratio is typically a small fraction, it is difficult to isolate and benefit from the properties of the TPSS [33]. In fact, high-throughput numerical searches have shown that, in theory, stoichiometric topological semimetals outnumber topological insulators [34], thus is of much interest to show that topological semimetals still exhibit properties related with their non-trivial topology [35]. Therefore, finding topological semimetals with a large surface-to-bulk conductivity ratio where the exotic properties of TPSS can manifest in a more direct manner and, additionally, with simple structures and high stability, would increase their possibilities of practical use.

In this work, we present a local morphological and electronic study of mechanically exfoliated FL antimonene flakes of thicknesses between ~ 2 and 40 nm (see Methods). Within this thickness range, the band structure fully reveals the decoupling of the top and bottom TPSS and the bulk bands present a finite contribution at the Fermi energy.

2. Results

Fig. 1 shows DFT calculations (see Methods) of the electronic band structure of FL antimonene in vacuum for different number of layers. The same calculation including adsorbed water molecules on the surface, to partially mimic ambient conditions, does not present significant changes (see Supplementary Figs 1 and 2). The insets represent the Fermi surface of each thickness, in the frame of the Brillouin zone (BZ, in blue). The band structure evolution with the number of layers shows two important features. First, the formation of Dirac cones around $\Gamma$ with the Dirac point about 0.3 eV below the Fermi energy. The Dirac hole band bends upwards to also cross the Fermi energy, giving rise to a peculiar Fermi surface with six hole pockets surrounding $\Gamma$. These pockets have a clear surface and helical character near $\Gamma$, but this character is partially lost as you move away from $\Gamma$ and they progressively become bulk states [27]. Second, the appearance of three fully bulk electron pockets crossing the Fermi level close to the $M$ points for over 5 layers. Moving away from $\Gamma$ towards the $M$-point ($K$-point), ignoring the crossing of the bulk electron pockets, one crosses the Fermi level three times (once) as expected for a topological material. The localization properties of these surface states have been studied in Ref. [27].

With this picture in mind, we proceed to the experimental characterization of our exfoliated FL antimonene flakes (see Supplementary Fig. 3 for characterization of FL antimonene flakes with different techniques). We start by analyzing flakes with different number of layers and lateral sizes in the range of several microns, deposited on highly doped Si substrates, with low energy and photoemission electron microscopy (LEEM/PEEM), in combination with synchrotron based X-ray photoelectron spectroscopy (XPEEM) (Fig. 2). The samples were first cleaned in ultra-high vacuum (UHV) (see Methods), ensuring flakes were free of oxide and contaminants, as demonstrated by the 5d 4d core level spectrum (inset in Fig. 2a), which shows the doublet with the 4d 5/2 at a binding energy of around 32 eV. Fig. 2a shows a LEEM image of several FL antimonene flakes. XPEEM images and spectra of the same flakes acquired in the maximum of the 5d 4d core level (Supplementary Fig. 4) reveal the antimony nature of all the flakes. Fig. 2b shows the AFM topography corresponding to the same flakes in Fig. 2a, presenting thicknesses from tens of nm down to terraces of just 1.5–2.5 nm (insets at the bottom left and top left corners respectively in Fig. 2b). We also carried out the electronic structure characterization by measuring the PEEM signal at the Fermi energy level ($E_F$) (Fig. 2c). The direct comparison of the AFM topography and the PEEM signal (see Fig. 2d) shows that the photoemitted intensity is mainly constant across the flakes and independent of the thickness in the range ~ 20–40 nm, while it is strongly quenched in certain areas marked by white arrows. These areas correspond to the thinnest zones, only 2–5 atomic layers (as shown in the top left inset in Fig. 2b), since the minimum height of a monolayer, when measured with AFM, is ~ 1 nm due to molecules trapped between the substrate and the flake [1]. A detailed analysis can be found in Supplementary Fig. 5.

These results are consistent with the DFT calculations shown in Fig. 1. For 1-2 layers the material is fully insulating, while for 3-4 layers, corresponding to the majority of the thinnest flakes in Fig. 2, the Fermi surface (see Fig. 1) is barely developed. For thicker flakes, the integrated contribution to the density of states at the Fermi energy comes mostly from the electron pocket at $\Gamma$ and the hole pockets around it. Note also that, due to the low electron mean free path at the recorded kinetic energy, the PEEM signal comes from the few topmost 2 or 3 surface atomic layers, around 100 meV below the $E_F$. Hence, as the TPSS have their weight centered there, it is not preposterous to think that this PEEM signal could be associated with the TPSS of the top layer and not from the bulk pockets that extend throughout the flake width. Next, we carry out an electrical characterization of FL antimonene flakes of different thicknesses. Kelvin probe force microscopy (KPFM) measurements on flakes with thicknesses between ~ 2 and 9 nm (~ 2-3–21 layers) [1] (see Supplementary}
Material Section S4) provides information on the Contact Potential Difference (CPD) variation, which is related to the work function change. As in other 2D materials [36], the measured CPD varies with increasing thickness and above a given number of layers, it saturates (see Supplementary Figs. 6 and 7). We also performed conductive AFM (C-AFM) measurements contacting the flakes with gold nanowire electrodes using the scanning-probe-assisted nanowire circuitry (SPANC) technique [37] and a metal-coated AFM tip as a second mobile electrode on flakes deposited on SiO₂. Fig. 3a shows FL antimonene flakes of different thicknesses contacted through SPANC. Gold nanowire electrodes are positioned on one end of the flakes and the conductive AFM tip is used to acquire current vs. voltage (IV) curves at different distances of the gold electrodes (see Supplementary Fig. 8) with nm resolution in ambient conditions (see Methods for details on the IV curves acquisition procedure). Supplementary Fig. 9 shows one of the flakes from Fig. 3 imaged more than one year later, showing no noticeable degradation, which accounts for the high stability of antimonene under ambient conditions [1−5].

This local measurement of the conductive properties allows us to obtain resistance vs. length (RL) plots from different thicknesses (Fig. 3b). From the RL plots and the geometry of the flakes, we can calculate the sheet resistance for the different thicknesses of the FL antimonene flakes (Fig. 3c), obtaining robust values regardless of the contact resistance (Supplementary Fig. 10). Within the sensitivity of our measurements, we can obtain an average value of ρ₂D = 1200 ± 300 Ω sq⁻¹, independent of the flake thickness. This result is remarkable considering that several bulk bands cross the Fermi energy in our thin flakes. For instance, the evolution of the bottom of the second bulk conduction band (located at M) with the number of layers (Supplementary Fig. 6) shows that, for 16 layers (approximately 6 nm, which closely corresponds to our thinnest flake), this second band already crosses the Fermi level. Thus, a higher number of bulk bands is expected to be present in the thicker flakes. Even so, the increasing number of bulk bands does not seem to appreciably increase the conductivity, suggesting a dominant contribution of the TPSS.

3. Discussion

It is instructive to compare the mentioned results with the case of graphene, where a linear increase of the sheet conductance (the inverse of the sheet resistance) has been observed with the number of layers [38,39] in contrast with our case, where it remains constant. We compare the sheet resistance (ρ₂D = 1200 ± 300 Ω sq⁻¹) with that obtained following the same procedure for FL graphene [37], ρ₂D,G = 670 ± 60 Ω sq⁻¹. Graphene has 4 Dirac cones, with intervalley scattering typically suppressed by long-range scattering potentials. The Dirac cone of FL antimonene is surrounded by hole pockets which, to a first approximation, altogether give a similar carrier concentration to that of graphene in ambient conditions (from the band structure we estimate n₂D ~ 3.5 × 10¹³ cm⁻²). The helicity of the states in the hole pockets near the Dirac cone are opposite to that of the nearby Dirac states which also prevents inter-pocket back-scattering [40]. Thus, the resistivity of FL antimonene is expected to be comparable but larger than that of
graphene, as observed experimentally. Combining the estimation of the carrier concentration from the band structure with the sheet resistance obtained from the transport measurements we can also tentatively estimate the mobility, $\mu$, by $\mu = \frac{\sigma_{2D}}{n_{2D} e}$, where $\sigma_{2D}$ is the bidimensional conductance ($\sigma_{2D} = 1/\rho_{2D}$), $e$ the electron charge, and $n_{2D}$ the bidimensional carrier concentration. We obtain a high mobility of $\mu \sim 150 \text{ cm}^2\text{V}^{-1}\text{s}^{-1}$ in ambient conditions. To put this figure in perspective, one can compare it with the mobility of black phosphorus, which is reported to be $\mu_{BP} \sim 50$–1000 cm$^2$V$^{-1}$s$^{-1}$ depending on the thickness [40].

In support of our observations suggesting a relevant role of the TPSS in the electronic properties, we have carried out two complementary calculations: a ballistic conductance calculation of defect-free FL antimonene contacted on the surface (Fig. 4a) and conductivity calculations in the presence of surface disorder (Fig. 4b). The results of the conductance (per unit length) are shown in Fig. 4a (see Methods for details and Supplementary Fig. 11 for several examples of transmission curves). The conductance increases with the number of layers, but at such slow pace that, for 9 layers, it is not even twice that of 3 layers (where only the Dirac cone contributes to the conductance) and visibly saturates. From these calculations, however, we cannot fully conclude whether the slow conductance increase is due to the full development of the TPSS in the center of the BZ or to the increasing presence of bulk states (for instance those at M). In order to assess this issue, we show in Fig. 4b the (diffusive) Kubo conductivity of our thicker FL system in the presence of disorder on both top and bottom surfaces (see Methods for details). We can calculate the conductivity from the TPSS alone or from all the states in our multilayers, surface and bulk. Here we plot the ratio between the TPSS conductivity of the top surface (actually considering only surface states, namely, excluding the halves of the hole pockets which correspond to bulk states), and the total conductivity (top TPSS + bulk states) for different disorder realizations as a function of the disorder strength affecting the bottom surface. In general, the TPSS contribution represents a significant fraction over the total conductivity (around 50%), despite introducing “only surface disorder”. In fact, increasing the strength of disorder on the bottom layer with respect to the top surface, the relative contribution of the surface conductivity with respect to the total one increases up to 70% (in our parameter range). This behavior is expected, as bulk states feel the disorder on the bottom layer more effectively, and it is practically (experimentally) relevant since, while the top surface can be treated or cleaned, the bottom one always remains affected by the substrate. This behavior is shared by all disorder cases up to a point where the bottom disorder gets strong enough as to also affect the top surface states in the hole pockets, which cannot be dissociated from the bulk states, reducing the TPSS conductivity. For thicker flakes, this turning point is likely to shift to larger bottom disorder strengths. The overall picture emerging from these two different calculations implies that the electrical transport is amply dominated by the TPSS, as suggested by our experimental results.

To conclude, we show the electrical characterization of mechanically exfoliated FL antimonene flakes within the ~2-3 to 100 layers regime, where topologically protected surface states might be playing a key role. We detect a strongly quenched photoemission signal at the $E_2$ for the thinnest flakes and a mainly constant value as the thickness increases. Our DFT calculations show that a transition from low to high surface density of states close to the Fermi level takes place at about 6 layers. We determine a sheet resistance of $\sim 1200 \Omega \text{sq}^{-1}$ above this threshold independent on the sheet thickness. We also estimate the mobility of FL antimonene in ambient conditions, resulting in $150 \text{ cm}^2\text{V}^{-1}\text{s}^{-1}$. Such a high value, in combination with its stability and simple structure, turns an-
antimonene into a promising candidate for nanoelectronics and optoelectronics applications. Hence, we envision that the richness in properties and phenomena of this material (including optoelectronic and energy-related) will pave the way for further studies on its many opportunities both from fundamental and practical points of view, with a particular focus in topological surface state physics, as for example for fault-tolerant quantum computation or as conducting channels with reduced dissipation in spintronic devices.

4. Methods

4.1. Sample preparation

We obtained FL antimonene flakes by mechanical exfoliation [1]. We placed a macroscopic freshly cleaved crystal of antimony (Smart Elements) in adhesive tape and after repetitive peeling, we directly transferred FL antimonene flakes to SiO$_2$/Si and highly doped Si substrates. A primary optical microscopy inspection allowed us to locate the thinnest flakes that we later imaged by AFM in contact mode to measure their thickness [41]. We used OMCL-RC800PSA cantilevers from Olympus with a nominal spring constant of 0.39 N m$^{-1}$ and low forces of the order of ~1 nN to ensure that the flakes were not deformed by the tip. We used WSXM software (www.wsxm.eu) both for the acquisition and processing of the AFM data [42,43].

4.2. PEEM/LEEM measurements

The experiments have been carried out at the PEEM experimental station of the CIRCE beamline at the ALBA Synchrotron [44]. All measurements were done in a low energy and photoemission electron microscope from micrometer-sized FL antimonene crystals. Prior to the measurements, we carried out a careful cleaning process in ultra-high vacuum (UHV), consisting of several annealing cycles at 400°C for 5 min in a hydrogen atmosphere (10$^{-6}$ mbar). This cleaning protocol ensured the flakes were free of oxide and contamination.

4.3. KPFM measurements

We carried out simultaneous dynamic mode AFM for the topography and frequency modulation mode for the KPFM [45] in a single-pass scheme, using metallized AFM tips (Budget Sensors ElectriMulti75-G). We applied an AC bias voltage of amplitude 5 V and frequency 7 kHz to the tip. We performed the KPFM measurements in an inert Ar atmosphere to avoid CPD shielding by the presence of adsorbed water on the surface of the samples [36].

4.4. C-AFM measurements

We contacted the FL antimonene flakes deposited on SiO$_2$/Si substrates using gold nanowire electrodes through the SPANC technique [37]. Briefly, we deposited gold nanowires on the substrates with FL antimonene flakes and assembled them into nanoelectrodes by AFM manipulation. In this case, the so-fabricated gold nanoelectrodes were connected to a microscopic gold electrode, we then used a metallized AFM tip (Budget Sensors ElectriMulti75-G) as a second mobile electrode to acquire IV curves at different locations. We employed dynamic mode AFM to image the samples, with an amplitude set point of 15 nm (cantilever free amplitude 20 nm). Then, we stopped the tip over the points of interest, and we brought it down into contact. There we acquired several IV curves and after this we brought the tip back to dynamic mode AFM. Before and after each set of IV curves we checked that the tip had not changed by acquiring IV curves on the gold nanoelectrode, ensuring tip stability along the whole set of measurements. To avoid artifacts, we also always carefully check the dependence of the conductance with the applied load, selecting the optimal conditions [46]. This procedure ensures that any possible layer of contamination or oxide on the surface is pierced and the current measured would respond to the intrinsic properties of the material [47]. In addition, we obtain the sheet resistance of the few-layer antimonene flakes from the flakes geometry and the slope of the Resistance vs. Lenght (RL) plots, which is independent of the contact resistance, thus obtaining reliable sheet resistance values within the uncertainty in the measured currents. RL plots acquired on the same flake but with different AFM tips can present different contact resistances, but they present the same slope (and there-
fore they lead to the same sheet resistance) (see Supplementary Fig. 10).

4.5. Band structure calculations

Band structure and Fermi surface contour calculations in Fig. 1 were performed using DFT as implemented in the Vienna Ab initio Simulation Package (VASP) code [48,49]. We employed the local density approximation (LDA) [50] together with the projector-augmented-wave (PAW) [49] method which has been shown to correctly describe Sb films and predicts the bulk lattice constant [21] in excellent agreement with the experimental value of 4.30 Å [51]. Calculations are performed considering SOC and a plane wave basis set was used with a cutoff energy of 300 eV on a $21 \times 21$ Monkhorst-Pack [52] k-point mesh. A vacuum region of 16 Å along the z direction was used for each system in order to minimize the interaction between the periodic repetitions of the cells. Full structural relaxations were performed for each of the 2L-7L systems until atomic forces were smaller than 0.001 eV Å$^{-1}$ [53]. The relaxed structure of the 7L system was used for the 9L system employed for the conductivity calculations (see below). We have taken defects into consideration as a perturbation (to all orders), expanding the eigenstates of the system + defects in a basis of Bloch states near the Fermi energy of the defect-free system (see Ref. [27] for details).

4.6. Ballistic transport calculations

We have obtained the room temperature conductance of the pristine multilayers thermal averaging the Landauer expression, $G = e^2/h \cdot T(\mu)$, where the transmission, $T$, is computed with the non-equilibrium Green’s function formalism [54]

$$T(\mu) = \frac{1}{N^2} \sum_k \Gamma(k, \mu) G(k, \mu) \Gamma(k, \mu) G^{-1}(k, \mu)$$

We have found that the most convenient way to contact the surface layer (mimicking the experiment) is, as shown in the insets of Fig. 4a, to continue to infinity this layer along the x direction, but gated with a potential of 2.0 eV that sets the Fermi level into the valence band where its conductance is maximum. The multilayer is finite in the x direction (with the maximum length permitted by our computational limitations) and periodic in the y direction with a minimal unit cell of width $a$ (as shown in the insets). Thus, the Green’s functions, G, and coupling matrices, $\Gamma$, depend on $k$ in this direction. The summation over $N$ values of $k$ in the first Brillouin zone divided by the actual width in the perpendicular direction, $Na$, gives the conductance per unit length. The Green’s functions and coupling matrices are evaluated from the non-collinear DFT Hamiltonian obtained with the OpenMX code [55] (http://www.openmx-square.org/), computing the whole system shown in the insets in Fig. 4a periodically repeated. OpenMX is based on norm-conserving pseudo-potential method with a partial core correction and a linear combinations of pseudo atomic orbitals (LCPAO) as a basis function here specified by Sb7.0-s3p2d2 (meaning that the cutoff radius is 7.0 Bohr and that three primitive orbitals for s components, and two primitive orbitals for each of p and d components are used). The exchange-correlation functional was the spin-polarized GGA-PBE. All calculations were performed until the change in total energy between two successive iteration steps converged to less than $10^{-6}$ Hartree. A cutoff energy of 220 Ry and an $11 \times 11 \times 1$ k-grid have been used in all presented results. The unit cell structure was also geometrically optimized. The quasi-Newton eigenvector following method was executed for structural relaxation of all structures until the change in forces between two successive iteration steps was less than $10^{-4}$ Hartree/Bohr. SOC was also employed and van der Waals interaction (vdW) correlation is considered by using the semiempirical dispersion-corrected density functional theory (DFT-D2) force-field approach. We have verified that the obtained band structure is essentially similar to that shown in Fig. 1 (see Supplementary Fig. 12).

4.7. Kubo conductivity calculations

We have chosen a system of 9 antimonene layers for the calculation of the Kubo conductivity. This number guarantees that the bulk pockets are well developed and that the TPS5 of opposite surfaces are completely decoupled. We first compute the DFT Kohn-Sham Hamiltonian with the CRYSTAL code [56,57] to which we have added SOC after self-consistency [58]. The lattice constant was considered (based on our previous work, Ref. [58]) to be $a = 4.27$ Å and the intra- and interlayer distances $h = 1.52$ Å and $d = 3.68$ Å, respectively. For these calculations we utilized a small-core pseudopotential basis set [59] with 23 valence electrons. For better agreement with plane-wave calculations we considered 0.94 scaling factor for the last filled $p$-orbital. The SOC enhancement factor 65, was also defined based on our previous work where we verified the SOC implementation vs. fully relativistic non-collinear VASP calculation (Ref. [58]). LDA exchange [60] and VBH correlation [61] functionals were used on a 32 $\times$ 32 Monkhorst-pack k-grid. For convergence, we applied a Fock (Kohn-Sham) matrix mixing of 97% between subsequent SCF cycles and the convergence on total energy was set to $10^{-9}$ Hartree.

We calculate the dc conductivity using finite-size Kubo formalism at room temperature. We have essentially followed the methodology developed in our previous work [27], but here we have fully implemented the exact evaluation of momentum matrix elements, as explained in Ref. [62]. The red bands shown in Fig. 1 are the relevant ones in our calculations. The exfoliation procedure guarantees that the internal layers of our flakes are defect-free, but not the surfaces. We also expect the bottom layer to be more influenced by disorder than the top surface due to the substrate effects (substrate charges and intercalated molecules). For simplicity, we assume the disorder not to mix states between bulk pockets or between bulk and surface pockets. This way the total conductivity is simply the sum of the individual pocket conductivities. For these calculations we have used a square sample of 45 nm in size and studied different disorder configurations defined by the strength of the top and bottom surface disorder, $\delta_t$ and $\delta_b$, and the top and bottom surface disorder concentrations, $n_t$ and $n_b$, respectively. In order to achieve statistical precision, we have considered at least 100 configurations of disorder on each surface up to the point that the behaviour of the curves in Fig. 4b was not changing anymore. Furthermore, we have smoothed the curves by averaging over the Fermi energy interval containing typically 11 levels.

Declaration of Competing Interest

The authors declare that they have no competing interests.

CRediT authorship contribution statement

**Pablo Ares:** Conceptualization, Methodology, Validation, Formal analysis, Investigation, Writing – original draft, Writing – review & editing, Visualization, Supervision. **Sahar Pakdel:** Software, Investigation, Writing – review & editing. **Irene Palacio:** Conceptualization, Methodology, Formal analysis, Investigation, Writing – review & editing, Visualization. **Wendel S. Paz:** Software, Validation, Investigation. **Maedeh Rassekh:** Software, Investigation. **David Rodriguez-San Miguel:** Validation, Investigation. **Lucia
Aballe: Investigation, Writing – review & editing. Michael Foerster: Investigation, Writing – review & editing. Nerea Ruiz del Árbol: Investigation. José Ángel Martín-Gago: Conceptualization. Methodology. Investigation, Writing – review & editing. Supervision, Funding acquisition. Félix Zamora: Conceptualization, Resources, Writing – review & editing. Funding acquisition. Julio Gómez-Herrero: Conceptualization, Methodology, Resources, Writing – review & editing, Supervision, Funding acquisition. Juan José Palacios: Conceptualization, Methodology, Formal analysis, Investigation, Resources, Writing – original draft, Writing – review & editing, Visualization, Supervision, Funding acquisition.

Availability of data and materials

The data generated during this study are available from the corresponding authors on reasonable request.

Acknowledgments

We acknowledge financial support through the “Marlade Maetzu” Programme for Units of Excellence in R&D (CEX2018-000805-M), the Spanish MINECO through projects PCI2018-093081/FISB-2018-0434-P, PID2019-109539GB-C43, PID2019-106268CB-C31 and -C32, MAT2016-77068-C3-1-P and -3-P, MAT2013-46753-C2-2-P and MAT2017-85089-C2-1-R, the EU Graphene Flagship funding (Graphene Flagship Core 3 881603 and JTC2017/2D-St&Ge), the EU via the ERC-Synergy Program (Grant ERC-2013-SYG-610256 NANOCSOMOS), the Comunidad Autónoma de Madrid through MAD2D-CM, S2018/NMT-4321 (NanomagCSTM-C) and the European StructuralFunds via FotoArt-CM project (S2018/NMT-4367), and the Fundación Ramón Areces. J.J.P. acknowledges the computer resources and assistance provided by the Centro de Computación Científico of the Universidad Autónoma de Madrid and the RES. S.P. acknowledges financial support by the VILLUM FONDEN via the Centre of Excellence for Dirac Materials (Grant No. 1174).

Supplementary materials

Supplementary material associated with this article can be found, in the online version, at doi:10.1061/japm.2021.101132.

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