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Robust band gap and half-metallicity in graphene with triangular perforations

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Ideal graphene antidot lattices are predicted to show promising band gap behavior (i.e., $E_G \simeq 500$ meV) under carefully specified conditions. However, for the structures studied so far this behavior is critically dependent on superlattice geometry and is not robust against experimentally realistic disorders. Here we study a rectangular array of triangular antidots with zigzag edge geometries and show that their band gap behavior qualitatively differs from the standard behavior which is exhibited, e.g., by rectangular arrays of armchair-edged triangles. In the spin unpolarized case, zigzag-edged antidots give rise to large band gaps compared to armchair-edged antidots, irrespective of the rules which govern the existence of gaps in armchair-edged antidot lattices. In addition the zigzag-edged antidots appear more robust than armchair-edged antidots in the presence of geometrical disorder. The inclusion of spin polarization within a mean-field Hubbard approach gives rise to a large overall magnetic moment at each antidot due to the sublattice imbalance imposed by the triangular geometry. Half-metallic behavior arises from the formation of spin-split dispersive states near the Fermi energy, reducing the band gaps compared to the unpolarized case. This behavior is also found to be robust in the presence of disorder. Our results highlight the possibilities of using triangular perforations in graphene to open electronic band gaps in systems with experimentally realistic levels of disorder, and furthermore, of exploiting the strong spin dependence of the system for spintronic applications.

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I. INTRODUCTION

Two-dimensional materials continually gain interest and achieve huge advances towards industrial realization in a number of fields, particularly electronics and spintronics. Graphene is the most studied material within the two-dimensional family [1] due to unique properties such as high electron mobilities [2] above $10^5$ cm V$^{-1}$ s$^{-1}$, gate-tunable carrier concentration [3], and predicted long spin-relaxation lengths [4] of several μm. These studies have led to substantial efforts in fabricating and processing clean graphene systems [5] as well as pushing the limits of nanostructuring, e.g., by high-resolution lithography [6,7]. To realize graphene-based electronics and in particular transistors, opening a band gap has been one of the main drivers of both theoretical and experimental work. Many studies propose using structural modifications of graphene systems, such as nanoribbons [8], or superlattice structures imposed by periodic gating [9,10] or strain [11,12], to achieve a band gap. More recent attempts have considered chemical modification through absorption, substitution, or sublattice symmetry breaking, for example, by doping [13–16]. Periodic patterning of graphene sheets, for example, periodic perforation to form so-called graphene antidot lattices (GAL) or nanomeshes, is of particular interest since theoretical predictions suggest the possibility of obtaining sizable band gaps [17,18]. Several groups have realized these structures in the laboratory [19–22]. Band gaps induced in periodically patterned graphene are however very sensitive to disorder and defects [23]. Current fabrication methods will inevitably yield systems with a significant degree of disorder. A clear experimental signature of minibands and -gaps has yet been elusive. In the magnetic and spintronic areas, the possibility of making graphene magnetic or realizing graphene-based spintronics has also attracted a lot of attention [4]. It has been predicted that pristine graphene exhibits uniquely long spin-relaxation times $\sim 1 \mu s$, although to date experiments still find relaxation times at least two orders of magnitudes lower; reasons for this are still under debate. Inducing magnetic ordering, or at least magnetic moments, is desirable in order to achieve tunable magnetism useful for magnetic information storage or spin-manipulation devices. There have been many works, theoretical and experimental, studying magnetic moments induced by vacancy defects [27–30], adatoms [27,30,31], substrate coupling, and molecular doping [32]. Nanostructured graphene is also predicted to display significant spin polarization at certain extended edges, namely those with a zigzag (zz) geometry [33,34]. Recent experimental findings also support the prospect of magnetic zz edges even with a reasonable amount of edge roughness observed [35–37].

In this paper, we propose using superlattices of triangular shaped GALs with entirely zz edges to gain large spin polarization, as confirmed by previous ab initio studies [38,39]. Graphene nanostructures which contain noncomplementary zz edges, e.g., triangles and Christmas trees (stacked triangles), display unique global ferromagnetic order [34,38,39], as we also will illustrate for the GAL case in Sec. III B below. In contrast, complementary zz-edged nanostructures, e.g., zz-edged hexagons, rhombi (two triangles back to back), or straight nanoribbons, display antiferromagnetic ordering [33,34,40]. Even before spin polarization is considered, we show through our tight-binding study how zz-edged triangular antidot lattices form exceptionally robust band gaps. When the effects of spin are included, a similarly robust half-metallicity is displayed

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findings and other important considerations. We envisage that triangular antidots could be fabricated, for example, through lithography using patterned hexagonal boron nitride as a mask. Hexagonal boron nitride naturally etches into triangular holes due to the different etch rates of the two species, i.e., boron and nitrogen [41]. Kinks or chirality within triangle edges may form during fabrication, but it is likely that they will still display a magnetic signature, albeit reduced, in accordance with theory for chiral graphene nanoribbons [42]. Our findings suggest a realistic path towards fabricating realistic spin polarized graphene nanostructures which could act as components in graphene-based spintronic devices.

The remainder of this paper is organized as follows. The system geometries and electronic and spin polarization models are described in Sec. II. Then we present our results in Sec. III, first considering several representative geometries in Sec. III A of both zz-edged and armchair(ac)-edged triangles without spin polarization. Next we focus on a single zz-edged antidot lattice and include spin interaction in Sec. III B. Finally we consider the robustness of our results by extending the tight-binding description in Sec. III C and by considering the effect of positional disorder in Sec. III D. In Sec. IV, we discuss our findings and other important considerations.

II. GEOMETRY AND MODEL

Rectangular arrays of triangular antidots are considered as shown schematically in Fig. 1. Specific geometries are denoted using \( X,Y,L_{\text{geo}} \) where \( X \) and \( Y \) represent the interantidot spacings in the two in-plane directions, \( L \) is the side length of the triangular antidot, and the index \( \text{geo} = \text{ac} \) or \( \text{zz} \) denotes the edge geometry of the triangles. \( X \) and \( Y \) take integer values and the associated antidot separations are \( Xa \) and \( Y\sqrt{3}a \), respectively, where the graphene lattice constant \( a = 2.46 \) Å. The rectangular superlattice makes for an ideal test bed for antidot lattices. The electronic properties change qualitatively with the superlattice dimensions, e.g., a semiconducting superlattice can become metallic and vice versa by changing the unit cell dimensions by just one lattice constant [17,18,43,44]. For any periodic external potential imposed onto graphene, for example, an antidot lattice, if the Fourier transformed potential is zero at the Dirac points of pristine graphene a band gap cannot form. Antidot lattices for which the Fourier transformed potentials are nonzero at the Dirac points have sizable band gaps. This criterion is from hereon referred to as the periodicity selection rules [44]. For rectangular superlattices, due to the lattice orientation chosen, the periodicity selection rules depend critically on the \( X \) spacing. All antidot lattices for which \( X = 3p \) where \( p = 1,2,3,\ldots \) are semiconducting, while for all other antidot lattices the existence of gaps or not depends on the particular antidot. Embedding the same triangular antidots into several rectangular superlattices which display different electronic behavior allows us to identify properties which arise due to the triangles themselves. The triangular antidots we consider are aligned to have either zz edges as shown in Fig. 1 or ac edges (not shown). The latter ac-edged triangles are rotated by \( 30^\circ \) with respect to those in Fig. 1 and the side length is scaled differently for the two orientations. \( L \) corresponds to a side length of \( La \) for the zigzag case and \( L\sqrt{3}a \) for the armchair case.

Spin polarization at single-point defects, as well as at zz edges, is usually interpreted via Lieb’s theorem [45]. The theorem states that the total ground state magnetic moment of a half-filled bipartite lattice is given by the sublattice imbalance, \( M = \mu_B|N_A - N_B| = \mu_B\Delta N \), where \( N_A \) and \( N_B \) are the number of sites belonging to each sublattice. Creating a zz triangle, such as that in Fig. 1, involves removing a different number of sites from the two sublattices and results in edge atoms belonging only to a single sublattice; with the orientation shown in Fig. 1 this is sublattice B. Accordingly, zz triangles form nonzero total magnetic moments, in full compliance with Lieb’s theorem [38,39]. Rotating the antidot \( 180^\circ \) flips the triangle orientation and also swaps the edge sublattice. Thus the relative edge sublattices of two adjacent triangular antidots can be determined by a quick visual inspection. The ac triangle has both sublattices present at the edge and is not expected to exhibit spin polarization [42]. We examine both the \( \{X,Y,L\} = \{24,15,5\text{zz/3ac}\} \) and \( \{25,15,5\text{zz/3ac}\} \) geometries; i.e., two geometries differing by \( a \) along the \( x \) direction and with either zz- or ac-edged triangular perforations; we later focus on the \( \{25,15,5\text{zz}\} \) superlattice with the zz triangle displayed in Fig. 1. The side lengths of the zz- and ac-edged triangles are similar for these geometries. The two triangle orientations highlight the fundamental differences between zz-edged triangular antidots and the other antidot families represented by the ac-edged cases.

The calculations in Secs. III A and III B are performed using a nearest neighbor (NN) tight-binding Hamiltonian

\[
H_\sigma = \sum_{i} \epsilon_i \epsilon_i^\dagger \epsilon_i + \sum_{ij} t_{ij} \epsilon_i^\dagger \epsilon_j. \tag{1}
\]

The operator \( \epsilon_i^{\dagger} (\epsilon_i) \) creates (annihilates) an electron with spin \( \sigma \) on site \( i \), and the hopping parameter \( t_{ij} \) takes the value \( t = -2.7 \) eV when sites \( i \) and \( j \) are nearest neighbor sites and is zero otherwise. \( |t| \) is taken as the unit of energy throughout.

FIG. 1. Schematic of the \( \{25,15,5\text{zz}\} \) triangular antidot superlattice geometry (left) and the approximately square unit cell with \( X = 25, Y = 15, \) and \( L = 5\text{zz}\)-triangular antidot (right). The A and B sublattices of graphene are denoted by white and black circles, respectively. The antidot spacings are approximately 6 nm, and the triangular side lengths are approximately 1 nm.
the paper. In Secs. III C and III D we will consider an extension to a third nearest neighbor model (3NN) by including terms $t_2 = -0.074|t|$ and $t_3 = -0.067|t|$ connecting second and third nearest neighbor sites, respectively [46]. The inclusion of $t_2$ results in a band-center shift, which we compensate for by adding a uniform on-site shift so that the Fermi energy lies at $E = 0$. Electron-electron interactions and the resulting spin polarization are included via spin-dependent on-site energy terms found from a self-consistent solution of the Hubbard model within the mean field approximation

$$\epsilon_{i\sigma} = \pm \frac{U}{2} m_i,$$

with $+$ for $\sigma = \uparrow$ and $-$ for $\sigma = \downarrow$, the on-site magnetic moments $m_i = \langle n_{i\uparrow} \rangle - \langle n_{i\downarrow} \rangle$, and $m_{i\alpha}$ is the number operator. We use the on-site Hubbard parameter $U = 1.33|t|$ which has been shown to give results in good agreement with full ab initio calculations for nanoribbon systems [47,48]. The self-consistent Hubbard calculations are initiated with an antiferromagnetic guess, $m_i = \pm c$, with opposite signs used for the two sublattices A and B, and then iterated to convergence.

III. RESULTS

A. Unpolarized antidots with different lattice geometries

We first consider periodic structures of zz- or ac-triangular antidots in the $U = 0$ case. The band structures of zz and ac-triangular antidots, together with their total density of states and that projected onto the (edge) B sublattice, are shown in Fig. 2. The zz cases shown in Figs. 2(a) and 2(b) for the $\{24,15,5\text{zz}\}$ and $\{25,15,5\text{zz}\}$ geometries, respectively, display both sizable band gaps and dispersionless midgap states. The fivefold degenerate midgap states originate from the single-sublattice zz edges. The level of degeneracy is equal to the sublattice imbalance $\Delta N$, which also equals the number of zz chains along the triangle edges $L = 5$. Similar midgap states are also observed in other noncomplementary zz-edged nanostructures, e.g., triangular quantum dots [49–51] and wide nanoribbons [46,52], where the degeneracy is proportional to the global sublattice imbalance in the quantum dots, and to the local imbalance in the wide nanoribbons. Such zz-edge states are localized on the edge sublattice. Within the NN approximation states localized in a single sublattice remain completely dispersionless. If higher order hopping parameters are included, such states can also become dispersive, as we will discuss in Sec. III C below.

The other characteristic of zz-edged triangular antidot lattices apparent from Figs. 2(a) and 2(b) is the formation of large electronic band gaps surrounding the dispersionless midgap states. In comparison, the ac cases shown in Figs. 2(c) and 2(d) reveal that the $\{24,15,3ac\}$ is gapped and the $\{25,15,3ac\}$ is metallic. These are in full compliance with periodicity selection rules, which in rectangular lattices predicts band gaps only for cases with $X = 3p$. The zz-triangular antidots with large band gaps regardless of $X$ indicate a different band gap mechanism. This hypothesis is supported by examining the band gaps of several triangular antidot lattices.

![FIG. 2. Unpolarized band structures and densities of states (DOS) of different triangular antidot systems. (a) $\{24,15,5\text{zz}\}$, (b) $\{25,15,5\text{zz}\}$, (c) $\{24,15,3ac\}$, and (d) $\{25,15,3ac\}$ geometries, respectively. The DOS projected onto the edge sublattice B (black) is shown together with the total DOS (gray). The structures in (a) and (b) notably show very large and narrow peaks in the DOS at the Fermi level $E = 0$.](image-url)

Pedersen et al. demonstrated that a scaling behavior $E_{\text{gap}} \propto \frac{N_{\text{rem}}}{N_{\text{tot}}} \propto \frac{L}{XY}$ was followed by many gapped graphene antidot lattices [17], where $N_{\text{rem}}$ and $N_{\text{tot}}$ are, respectively, the number of atoms removed to form an antidot and the total number of atoms in the superlattice unit cell before the antidot atoms are removed. In Fig. 3, a linear behavior is clearly noted for those ac-edged systems with periodicity selection rules predicting semiconducting behavior (filled green squares) whereas those for metallic superlattices (hollow green squares) have zero band gap in almost all cases. We associate the breakdown of

![FIG. 3. Unpolarized band gaps for various geometries as the dimensionless parameters $X$, $Y$, and $L$ are varied. The zigzag (blue) and armchair (green) triangle geometries are divided into groups where the superlattice is expected to display semiconducting (sc, filled) or metallic (m, hollow) behavior, according to the periodicity selection rules. For a rectangular superlattice this distinction depends solely on the value of $X$.](image-url)
this trend for metallic systems with large $\frac{t}{\Delta}$ to large antidots in small unit cells, where additional band gap behavior is now induced by small contractions between the antidots. The $zz$ triangles are meanwhile consistently gapped (blue circles), irrespective of the behavior predicted by periodicity arguments. The band gap magnitude has an approximately linear dependence on $\frac{t}{\Delta}$, but the slope is much greater than the $ac$ case. The reason $zz$-edged triangular antidot lattices are consistently gapped is the global sublattice imbalance which induces sublattice symmetry breaking. Independent of the periodicity selection rules, sublattice symmetry breaking imposes an effective nonzero potential between sublattices in a similar manner to a mass term, i.e., a staggered on-site potential, with a different on-site potential for each sublattice. In other systems where sublattice symmetry is broken, for example, by doping such a term also opens a band gap [15,16].

The sublattice-projected densities of states (DOS) for $zz$-triangle lattices in Figs. 2(a) and 2(b) show that each sublattice contributes equally to the DOS at all energies except at the $E = 0$ edge states which reside only on the B sublattice. However the local density of states (LDOS), shown in Fig. 4, reveals a more complex picture. The edge state localization is clear in Fig. 4(a) where the LDOS is mapped at $E = 0$ by circles whose radius is proportional to the LDOS at that site. White and black circles are used for sites on the A and B sublattices, respectively, and we note that only large black circles near the triangle edges are found at this energy. Despite the equal contributions from sublattice projected DOS at other energies, the LDOS distributes inhomogeneously around the triangles. This is shown in Fig. 4(b) for the conduction band energy $E = 0.1|t|$, where we note that the B sublattice contribution to the DOS is now spread throughout most of the unit cell, but is significantly larger near the triangle edges. The A sublattice has a vanishing LDOS in this region and its DOS contribution is mostly distributed at sites midway between neighboring antidots. The dispersion of the states at this energy is due to the large regions where both sublattices have a significant occupation. The different electron distributions for the A and B sublattices suggest different effective scattering potentials for the different sublattices. The inhomogeneous LDOS distribution, together with the band gap formation regardless of periodicity selection rules, suggests that sublattice symmetry breaking is the driving mechanism behind band gap formation and not the periodic selection rules usually forming band gaps in graphene antidot lattices. Importantly, this suggests that band gap behavior in $zz$ triangles should be stable against geometrical variations as long as the sublattice imbalance is maintained. Since the dimensions $X$ and $Y$ of the antidot lattice play a minor role, one may expect that lattices made of triangular $zz$-edged antidots are robust against disorder, as we discuss in Sec. III D below.

B. Effect of spin polarization

A nonzero Hubbard interaction $(U = 1.33|t|)$ leads to spin dependence in $zz$-edged triangle systems through the formation of magnetic moments $m_i$.

The self-consistent solution to the Hubbard model using the $[25, 15, 5zz]$ geometry is shown in Fig. 5. Different superlattice and triangle dimensions always yield a similar pattern, namely a distribution with antiferromagnetic alignment between moments on the different sublattices. The magnitude of the moments is maximum at the $zz$ edges, decreases slightly towards the corners of the triangles, and quickly decays perpendicular to the $zz$ edges. Similar moment distributions have been reported in $ab\text{ initio}$ studies of triangular perforations [38]. Triangles with large side lengths have long segments with approximately constant magnetic moments with a maximum $m_i \sim 0.31 \mu_B$. Only below $L < 5$ do these constant-moment segments vanish and the maximum moment decreases. All the geometries considered are consistent with Lieb’s theorem such that $M = \sum_{i} m_i \mu_B = \mu_B \Delta N \equiv \mu_B L$. The triangle corners are geometrically similar to the kinks arising in chiral graphene nanoribbons, which display a similar drop in moment values [42]. The magnetic moment profile is found to be almost completely independent of the superlattice geometry, suggesting that nearby triangles do not influence each other unless they are very near.

The spin-split band structure of the $[25, 15, 5zz]$ system is shown in Fig. 6(a), together with the spin up (red) and
spin down (blue) DOS in Fig. 6(b). As before, the lighter regions show the total DOS, and the darker regions show its projection onto the edge B sublattice. There are a number of key differences from the unpolarized band structure of the same geometry system in Fig. 2(b) compared to the spin polarized band structure in Fig. 6(a). The fivefold degenerate dispersionless bands are no longer present at zero energy, and the band gap is considerably reduced by the presence of dispersive bands at the energies $E = \pm 0.02|t|$. These bands have opposite spin orientations on either side of $E = 0$, as do the five low-dispersive nondegenerate bands in the energy range $(\pm 0.1|t| \to 0.15|t|)$. To examine the formation of this band structure the Hubbard interaction $U$ is varied from a low $U = 0.2|t|$ to $U = 1.33|t|$ in Fig. 6(c), left to right. The band structures in these panels correspond to the region shown by the dashed box in Fig. 6(a). We denote three low energy spin down bands $V$, $VI$, and $VII$ at low Hubbard-interaction strength $U = 0.2|t|$, the fifth through seventh lowest energy spin down bands in this region. In the unpolarized band structure, band $V$ corresponds to one of the fivefold degenerate dispersionless bands whereas $VI$ and $VII$ form the conduction bands. The $V$, $VI$, and $VII$ bands are labeled both at the left- and rightmost panels for clarity. These panels reveal how the formerly degenerate and dispersionless bands undergo different degrees of spin splitting. The highest of these ($V$) initially at low $U$ (left) appears below both bands $VI$ and $VII$ and finally at high $U$ (right) appears above said bands. The degree of spin splitting is determined by the LDOS distribution and the magnitude of the magnetic moments. High degrees of spin splitting can be attributed to a LDOS localized around areas with large magnetic moments, which is confirmed by examining the spin polarized LDOS.

At $U = 1.33|t|$ and $E = 0.155|t|$, corresponding to the $zz$-edge states band $V$ and shown in Fig. 7(a), the LDOS is localized almost entirely on magnetic edge sites, consistent with a large degree of spin splitting. Meanwhile, the LDOS of the spin polarized conduction bands at $E = 0.02|t|$ shown in Fig. 7(b) is mostly localized near the triangle corners which have smaller magnetic moments, consistent with a small degree of spin splitting. Further, the dispersion of the conduction bands is shown to emerge due to a nonzero occupation of the A sublattice as shown in Fig. 7(b). In the unpolarized case bands $VI$ and $VII$ define the conduction band edge, but as $U$ increases [see Fig. 6(c)], the spin-down versions flatten and increase in energy, whereas the spin-up versions broaden and decrease slightly in energy. We noted earlier that the unpolarized cases displayed LDOS contributions from both sublattices, which overlapped to form dispersive conduction bands. When spin polarized, this distribution is quite different for each spin. The LDOS of the spin-down band shown in Fig. 7(c) is localized almost entirely on B sublattice sites near the center of the $zz$-edge sections, which leads to a flattening of the dispersion and an upwards energy shift. Conversely, the LDOS of the spin-up bands shown in Fig. 7(d) is localized both on the B sublattice near the antidot corners and on sites from both sublattices further away from the triangle. The more homogeneous distribution of the spin-up bands leads to further broadening and a weaker downwards shift from spin splitting.
The band gaps for spin polarized zz triangles are shown for a range of geometries in Fig. 8, where we note a decrease of approximately one order of magnitude compared to the unpolarized cases. In fact, the gaps of semiconducting ac triangles are larger than those for polarized zz triangles. However, spin polarized zz-edged antidots display another interesting feature. The dispersive states surrounding the band gap are completely spin polarized, so that a spin-selective half-metallicity can be induced by small $E_F$ shifts applied using a back gate. This suggests that such geometries may be employed in a range of spintronic components to filter spins of different orientations.

Many of the features we have described in both unpolarized and polarized zz triangles depend on the inhomogeneous electron distributions and in particular the localization on the edge sublattice and near zz edges. It is important to determine if such features are artifacts of the NN model we employ for our calculations and whether they are robust in the face of disorder. The latter point is of interest as many effects induced by superlattices tend to vanish at any realistic disorder [23]. We now briefly address both issues.

C. Effect of higher order hopping terms

Within the NN model, states which occupy only a single sublattice appear completely dispersionless. In comparison, a 3NN model enables intrasublattice coupling by the inclusion of the 2NN terms, and the parametrization we use has been shown to accurately describe zz nanoribbons [46]. For the unpolarized case, we note that the introduction of these additional hopping terms leads to an energy splitting of the previously degenerate midgap states, see Figs. 9(a) and 9(b). This leads to a shift of the Fermi energy relative to the bulk valence and conduction bands in order to satisfy half-filling, increasing the electron-hole asymmetry already introduced by the 2NN hoppings. The NN-model band gap can be identified in the 3NN band structure between the energies $E = -0.025|t|$ and $E = 0.1|t|$ but is now slightly smaller and more importantly contains multiple midgap states. In particular a dispersive channel opens at $E \sim 0.05|t|$, similar to that seen near zz-ribbon edges when a 3NN model is employed [53]. Disregarding these midgap states, the 3NN band gap between $E = -0.025|t|$ and $E = 0.1|t|$ scales similarly to the NN model when varying the system dimensions. The emergence of dispersive states in the band gap could of course limit the applicability of these systems. However we note that in many cases they have either very little dispersion, or are spaced far enough apart in energy, so as to still offer reasonable band gap or transport gap behaviors.

Considering the polarized case, the band structures and DOS in Figs. 9(c) and 9(d) are remarkably similar despite the large changes we have discussed in their associated unpolarized versions. The most significant change now between NN and 3NN models is the expected (minor) electron-hole asymmetry. Notably the system remains half metallic with spin-dependent dispersive states close to the Fermi level. The excellent agreement between NN and 3NN models in this case can be understood by the fact that the features introduced by the additional 3NN terms in the unpolarized case, namely dispersion and splitting of the midgap states, also result independently from the inclusion of the spin-dependent potentials.

We note that the 3NN model, both with and without spin polarization, also agrees qualitatively with previous $ab$ initio calculations, which display similar band structures [38]. Although the 3NN model serves to correct the missing intersublattice interaction, it appears that the most important behavior in polarized systems is captured by the lower order NN model.
D. Robustness against disorder

One of the major obstacles in inducing band gaps using graphene antidots is that the large gaps predicted in atomically precise systems are extremely fragile in the presence of even mild geometrical disorders \[23\]. The gap mechanism for usual antidot arrays, namely the periodicity selection rules, relies on pristine conditions and regular antidot spacing. We have shown that zz-edged triangular antidots behave very differently from other antidots and that their behavior arises from the breaking of sublattice symmetries around individual antidots. We further demonstrated that these effects were independent of the superlattice geometry, which suggests also that they should be more stable than, for example, ac-edged antidots, in the face of disorder. While a full-fledged disorder analysis is beyond the scope of the present paper, we highlight the essential effects by examining a \(4 \times 4\) geometrically disordered array of antidots in a repeated superlattice. We present one particular random configuration but also note that an additional 10 different configurations have been examined all showing qualitatively the same behavior. The triangle centers are randomly shifted by \(\Delta \mathbf{r} = [\delta x, \delta y(\sqrt{3} a)]\) with \(\delta x = \delta y \leq 3\), as shown for zz triangles in Fig. 10(a) and ac triangles in Fig. 11(a). The same size triangles as before are considered, but for computational efficiency we use smaller \(\{X = 15, Y = 9\}\) “blocks” to compose the supercell, essentially cutting down on the amount of pristine graphene between perforations. Note that according to the periodicity selection rules these superlattice geometries are predicted to form band gaps \[44\].

![FIG. 10. (a) Schematic of a disordered 4 \times 4 array of \{15,9,3ac\} triangular antidots. (b) Pristine band structure. (c) Disordered system for \(U = 0\). (d) Pristine system for \(U = 1.33|t|\). (e) Disordered system for \(U = 1.33|t|\).](image-url)
IV. CONCLUSION

We have discussed the electronic properties of triangular antidots systems in graphene sheets, with a particular focus on zigzag edged geometries whose geometry breaks the symmetry between graphene’s two sublattices. In order to shed light on the possibility of magnetic states at such edges, we have analyzed systems in both the spin polarized and the unpolarized cases. We further have illustrated the robustness against disorder by individually displacing the antidots of a 4 × 4 array unit cell. Spin unpolarized superlattices of triangular zz-edged antidots form band gaps significantly larger than similarly sized ac-edged counterparts. Gap opening occurs irrespective of conventional rules governing the formation of band gaps in, for example, ac-edged triangular antidots and scales with the triangular antidot side length. Furthermore zz- as opposed to ac-edged triangles are far more robust against geometric disorders. We conclude that these unique features are caused by a gap-opening mechanism related to sublattice-symmetry breaking. In contrast to conventional graphene antidot lattices, this mechanism is less sensitive to experimentally unavoidable imperfections in lattice spacings. The zz-edged triangular antidots become half-metallic over a wide range of energies when spin polarization is included, with a high degree of spin selectivity achievable by gating. Spin splitting of the unpolarized cases leads to the emergence of dispersive spin-dependent states and subsequent reduction of the band gaps compared to the unpolarized cases. The half-metallic behavior of zz-edged triangles also appears more robust against geometric disorder compared to ac-edged counterparts. These findings suggest a robust path to realize devices based on nanostructured graphene with robust band gaps. Further, devices with half-metallic and spin-selective properties appear feasible.

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