Tuning electronic structure of graphene via tailoring structure: theoretical study

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Abstract

Electronic structures of graphene sheet with different defective patterns are investigated, based on the first principles calculations. We find that defective patterns can tune the electronic structures of the graphene significantly. Triangle patterns give rise to strongly localized states near the Fermi level, and hexagonal patterns open up band gaps in the systems. In addition, rectangular patterns, which feature networks of graphene nanoribbons with either zigzag or armchair edges, exhibit semiconducting behaviors, where the band gap has an evident dependence on the width of the nanoribbons. For the networks of the graphene nanoribbons, some special channels for electronic transport are predicted.

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I. INTRODUCTION

Graphene has been a hot issue in both fundamental and application research fields, since stable single graphene layer at room temperature was achieved experimentally [1–4]. Such two-dimensional hexagonal honeycomb has novel electronic [5], mechanical [6] and thermal [7] properties. For example, a perfect graphene sheet is semimetallic, being an extremely good conductor. Meanwhile, graphene exhibits astonishing transport properties: electron mobility in graphene was predicted to be as high as 15000 cm$^2$/Vs at room temperature. Moreover, atomic thickness, chemical inertness and almost perfect structure makes it promising material for device application. Therefore, graphene holds a sound prospects for a new generation of graphene-based electronics [8–12].

However, most electronic and optoelectronic devices require a semiconductor with a finite gap. To broaden its applications, many efforts have been devoted to turn the semimetallic graphene into a gapped semiconductor, through loading strain [13], introducing defects [14], and performing chemical functionalization [15, 16]. Of these efforts, designing superlattice scale defects in graphene sheet, so called nanopatterning, has drawn much attention.

Experimentally, patterning of a graphene sheet by e-beam lithography with features as small as ten nm was available [17–19], indicating that patterning graphene with the desired features is possible. In fact, a quasi-one-dimensional graphene nanoribbon (GNR) has been successfully achieved, and its electronic structures have a strong dependence on the width and chirality [20–22]. Quite recently, nanoscale square array of holes have been fabricated on the graphene in experiment, and the existence of a transport gap was found [23, 24].

On the theoretical side, graphene antidot lattices, graphene sheets with regularly spaced holes, have been proposed as a platform for quantum confinement of electrons in graphene [25–27]. Based on the tight-binding (TB) calculations, the electronic structures of the antidot lattice with triangular arrays or roughly holes in graphene sheet were illustrated by two groups [25, 27]. Such arrays of nanoscale perforations in graphene manipulate the electronic structures of the systems significantly, by giving rise to the localized states, or inducing the band gap near the Fermi level. Very recently, such antidot lattices proposed by Pedersen et al were more accurately studied by using the density functional theory calculations [28]. Band gaps ranging from 0.2 to 1.5 eV were reported. However, all these concerned antidots were arranged in a hexagonal unit cell. If the triangular (hexagonal) holes are arranged in
a rectangular supercell, the antidots display different geometry features. Whether do the array features have significant influence on the electronic behaviors?

On the other hand, although square arrays have been achieved experimentally, theoretical efforts on such defective patterns have not been reported yet, to our best knowledge. Structurally, such square arrays behave as the networks of armchair edged GNRs (a-GNRs) and zigzag edged GNRs (z-GNRs). As we know, a-GNRs and z-GNRs display different electronic properties: the former exhibits an interesting three-family behavior, while the latter has a direct band gap which decreases with increasing width of the z-GNR \[22\]. For the networks consisting of a-GNRs and z-GNRs, electronic properties of the systems are still puzzling.

In this work, we firstly investigate the electronic structures of graphene sheet with triangular and hexagonal holes in a rectangular supercell, based on the first principles calculations. Then rectangular holes are introduced into a graphene sheet, which feature the networks of a-GNRs and z-GNRs. We systematically explore the electronic structures of such patterned graphene with varying the width of either a-GNR or z-GNR. It is found that the graphene sheets with rectangular holes are all semiconductors. Meanwhile, tuning band gap can be practiced through different patterning. Moreover, the size-effect of different structural feature on electronic structures is also discussed.

II. COMPUTATIONAL METHOD

Our calculations are performed by using the SIESTA program \[29\] at the level of local density approximation, in which the norm-conserving pseudopotential are taken into account. Spin-polarized calculations are performed for all our concerned cases. For both C and H atoms, double-ζ basis sets \[30\] are chosen for calculations. For each concerned system, the periodic boundary conditions along x and y axes are considered. Meanwhile, a spacing of more than 10 Å is applied along z axis to neglect the interaction between the graphene monolayers. All atomic positions and the lattice constants of the concerned systems are allowed to be fully optimized, with the residual force convergence value of less than 0.02 eV/Å. The Brillouin zone is sampled with 12 × 12 × 1 according to the Monkhorst-Pack scheme \[31\], which is tested to be enough for our calculations. In the investigations of the electronic structures, more k-points with sampling of 18 × 18 × 1 are employed.
III. ELECTRONIC STRUCTURES

Three kinds of structural patterns (triangular, hexagonal and rectangular holes) in graphene sheets are concerned, which are shown in Fig. 1. For the systems with triangular or hexagonal holes, the lattice constants of the supercell are chosen to be 25.56 and 24.60 Å along $x$ axis and $y$ axis respectively. While for the systems with rectangular holes, they are set to be 25.56 and 19.68 Å. Fig. 1-(a) is a generated graphene with a triangular void by removing a triangular portion with zigzag edges. In addition to the graphene with absence of a triangular portion shown in Fig. 1-(a), three more cases with different size of the voids are considered, which are formed by removing the carbon atoms at the edges marked with $E_1$, $E_2$ and $E_3$ in Fig. 1-(a) in turn. For convenience, we notate the systems with triangular holes from small size to large size as tri-1, tri-2, tri-3 and tri-4 respectively. In our calculations, the carbon atoms with two-folded coordinates near the edges are terminated with hydrogen atoms for all our concerned cases.

The calculated density of states (DOS) for the cases of tri-1, tri-2, tri-3 and tri-4 are illustrated in Fig. 2-(a-d), in which the DOS of the corresponding perfect sheet is plotted as a reference (in Fig. 2-(d)). Clearly, the majority spin states and the minority spin states split, and the peaks strikingly appear near the Fermi level, which is consistent with the previous report [28]. For the cases of tri-1 and tri-2, two evident peaks emerge at about $E_f - 0.1eV$ and $E_f + 0.1eV$ in the calculated DOS respectively. With increasing the size of the defects, the peak under the Fermi level becomes broad and evolutes to be two peaks eventually. This is shown obviously in Fig. 2-(d) for the case of tri-4. The further analysis reveals that these peaks near the Fermi level are mainly contributed from the carbon atoms in the outmost row at the defect edges which bond to hydrogen. For convenience, we refer these atoms as the edge atoms. Moreover, we find that most carbon atoms which bond with three other carbon atoms contribute little to these peaks, whether these three coordinated carbon atoms are close to or far away from the edges.

Comparing the DOS of tri-1, tri-2, tri-3 and tri-4 (Fig. 2-(a-d)) with each other, one can find that the peaks near the Fermi level become stronger, with increasing the defect size. This is rationalized: the large defect introduces more carbon atoms at the edges which contribute to these peaks. Moreover, as the defect size increases, the interaction of the carbon atoms at the edges with their images become stronger. As a result, such interaction
induces another peak under the Fermi level. Examination the configurations reveals that the C-C bonds at the edges are shortened to be $1.404 \pm 0.04 \text{
Å}$, and the C-C bonds between the first shell and the second shell near the edges are enlarged to be $1.433 \pm 0.04 \text{
Å}$. Such distortion becomes weak in the third shell (C-C bond length is about $1.419 \text{
Å}$), indicating that the defect is localized in space.

To study the hexagonal holes in a graphene sheet, we first remove the carbon atoms in a hexagon ring, leaving a hexagonal void in the sheet. Then the shells of carbon atoms marked with $S_1$, $S_2$, and $S_3$ in Fig. 1-(b) are removed one by one. We name these systems as hex-1, hex-2, hex-3 and hex-4 respectively. The calculated DOS for these four systems are shown in Fig. 2-(e-h), in which the DOS of the perfect graphene is also plotted in Fig. 2-(h) for comparison. Different from that of triangular arrangements, the majority spin states and the minority spin states are degenerate for the graphene sheet with hexagonal holes. Such difference in spin polarization can be understood with Lieb’s theorem [32]. As shown by Lieb in graphene with a bipartite lattice in the nearest neighbor approximation, the angle between the zigzag edges of triangular arrangements is $60^\circ$, and the system is spin splitting. In contrast, the angle between the zigzag edges of hexagonal arrangements is $120^\circ$, and the system is spin degenerate [28, 32].

As shown in Fig. 2-(e), the DOS of hex-1 change little with respect to that of the perfect graphene, which implies that absence of a hexagonal ring in graphene sheet perturbs the electronic structures of the system slightly. In the case of hex-2, a weak peak at about $E_f + 1.2 \text{ eV}$ appears (Fig. 2-(f)). The local density of state analysis reveals that this peak is contributed from the edge atoms. With increasing the defect size, the interaction between the defect edges becomes stronger, and the electronic structures of the systems change largely, especially near the Fermi level. This is clearly displayed in Fig. 2-(g-h). The calculated local density of states reveal that the states near the Fermi level dominantly come from the edge atoms of the defect. After checking the geometries of the systems, we find that for hex-1, the fluctuation of the C-C bonds, either shortened to be $1.393 \text{ Å}$ or enlarged to be $1.442 \text{ Å}$, happens within the first shell around the edge. With increasing the size of the defect, such distortion of the geometry becomes heavier. For the case of hex-4, the structural distortion almost extends to the whole system, which eventually gives rise to a small band gap. This is consistent with the previous works [25, 27].

It is noted that for both triangular and hexagonal arrangements mentioned above, each
TABLE I: The band gap (in eV) for each case of rectangular arrays of \((N_a, N_z)\). \(N_a\) and \(N_z\) are the width of a-GNR and z-GNR respectively, which are described in the text.

| \((N_a, N_z)\) | (4,4) | (5,4) | (6,4) | (7,4) | (8,4) | (9,4) | (10,4) | (11,4) | (12,4) |
|----------------|------|------|------|------|------|------|------|------|------|
| band gap       | 0.48 | 0.14 | 0.68 | 0.29 | 0.18 | 0.75 | 0.25 | 0.10 | 0.71 |

defect has zigzag edges. For comparison, the cases with armchair edges are taken into account. A triangular hole with armchair edges, whose size is similar to that of tri-2, is concerned as a typical example. Different from that of tri-2, the majority spin and the minority spin states are degenerate. The energy gap is about 0.10 eV, and the calculated DOS is like to that of hex-2 somewhat. Moreover, we also investigate the electronic structures of a typical hexagonal array with armchair edges, the defect size and the feature of the DOS are both similar to that of hex-3, with a gap of about 0.15 eV.

In addition to the triangular and hexagonal arrays, we also arrange the rectangular holes in the graphene sheet. A crossed GNR is chosen as supercell, which is shown in Fig. 1-(c), and the network of GNRs is shown in Fig. 1-(d). Following the previous notation \[20, 22, 33\], the width of a-GNR is notated with \(N_a\) and that of z-GNR with \(N_z\), as marked in Fig. 1-(c). For convenience, we classify the graphene containing the rectangular holes with using two indexes of \(N_a\) and \(N_z\), notated as \((N_a, N_z)\). For such structural patterns, we consider two series of systems: \(N_a\) varies from 4 to 12, with keeping \(N_z\) to be 4; and \(N_z\) varies from 5 to 8, with keeping \(N_a\) to be 6. By calculations, we find that all the concerned systems exhibit semiconducting behaviors. Moreover, the band gap depends evidently on the width of a-GNR, but weakly on that of z-GNR. The band gaps for the concerned cases with varying \(N_a\) are listed in Table I. From the Table I, one can find that the gaps for (6,4), (9,4) and (12,4) are close and much larger, while the gaps for (5,4), (8,4) and (11,4) are close and smaller. With increasing the width of a-GNR, the gap variation trend seems follow the three-family behavior reported in the infinite a-GNR \[22\]. In contrast, for the cases with varying \(N_z\), the band gaps change slightly. The band gaps are 0.22, 0.31, 0.09 and 0.14 eV for the cases of (6,5), (6,6), (6,7) and (6,8) respectively.

For the rectangular cases, the majority spin and the minority spin states are degenerate. Fig. 3-(a-c) show the DOS of a typical three-family system of (7,4),(8,4) and (9,4) respectively. Evidently, the band gap is about 0.29 eV for \(N_a=7\), and it is narrowed to be 0.18
eV for \( N_a = 8 \), then broadened to be about 0.75 eV for \( N_a = 9 \). The similar variation trend is found for the other two families. In addition, the electronic structures near the Fermi level exhibit different features for each case in this family. Typically, in the case of \((7,4)\) (Fig. 3-(a)), two peaks emerge at about \( E_f \pm 0.6 \) eV, which are mainly contributed from the edge atoms of the z-GNRs (z-edge); The peaks at about \( E_f \pm 0.3 \) eV are ascribed to the atoms which bond to hydrogen at the crossing site of a-GNRs and z-GNRs (corner). Moreover, the edge atoms of the a-GNRs (a-edge) contribute a little to these peaks. For the case \((8,4)\), the peaks at about \( E_f \pm 0.6 \) eV remain, while the peaks at about \( E_f \pm 0.3 \) eV become flat, as shown in Fig. 3-(b). The atoms at the z-edge and a-edge, as well as at the corner all make contribution to these peaks, of which the former is dominant. As for the case of \((9,4)\) (Fig. 3-(c)), the concerned peaks move to \( E_f \pm 0.5 \) eV and \( E_f \pm 0.9 \) eV respectively, broadening the gap to be 0.75 eV. The former are contributed from the corner atoms, and the latter from the atoms at the z-edges. While those at a-edges contribute little to these peaks. Fig. 3-(d-f) shows the DOS for the cases of \((6,6),(6,7)\) and \((6,8)\) respectively. Similar electronic behaviors near the Fermi level are found in these systems: two flat peaks at about \( E_f \pm 0.25 \) eV and two peaks at about \( E_f \pm 0.8 \) eV. The former is mainly contributed from the atoms at the z-edges, and the latter is attributed to those at the corner and the a-edge.

Overall, for the rectangular arrangements, the peaks in DOS near the Fermi level are dominantly contributed from the atoms at the edges including a-edges, z-edges and the corners. This is essentially ascribed to the large structural distortion near the defect edges: the C-C bonds within two neighbouring are enlarged (shortened) by about 0.03 Å (0.05 Å). Moreover, the width of the a-GNR has significant influence on the electronic structures of the system, whereas the width of z-GNRs has little. This can be well understood from the DOS analysis: the states near the Fermi level dominantly come from the contribution of the carbon atoms at the z-edges, but the number of such atoms decrease with increasing the width of a-GNR. Therefore, the width of the a-GNR have a crucial role on the electronic structures, and this agrees well with that of infinite GNRs, in which the energy gap of a a-GNR significantly dependents on its width [22].

To better understand the nature of the states in the vicinity of the Fermi level, we further explore the charge density for the concerned systems. Some typical cases of the triangular and the hexagonal arrangements are given in Fig. 4 where the highest occupied molecular orbital (HOMO), the lowest unoccupied molecular orbital (LUMO) and the level under the
HOMO (HOMO-1) at Γ point are plotted. For the graphene with zigzag-edged triangular arrangements, the majority spin states for the HOMO-1 and the HOMO are of clear localized nature. The former is mainly contributed from the atoms at the edge along $y$ axis, and the latter is from those at the other edges. In contrast, the minority spin states for the HOMO-1 and the HOMO are delocalized. The majority spin state and the minority spin state for the LUMO are mostly confined in three edges. This is obviously shown in Fig. 4(a-b) (tri-2 is a representative). For the triangular arrangements with armchair edges, the HOMO-1, the HOMO and the LUMO states are all of extensive nature, especially for the HOMO and the LUMO states, as shown in Fig. 4(d). These states, with features of local $\pi$-bond, are contributed from the atoms in the strips along $x$ axis which contain the defect arrays or not.

For the graphene containing the hexagonal arrangements with either zigzag edges or armchair edges, these concerned orbitals are also localized in a small region near the defect edges: the HOMO-1 state is from the atoms at the edges along $x$ axis, and the HOMO and the LUMO states are from that along $y$ axis. The typical case of hex-3 and the hexagonal holes with armchair edges are shown in Fig. 4(c) and (e) respectively.

From above, we find that for the graphene with hexagonal arrangements, zigzag edges and armchair edges make little difference on the electronic structures of the systems. In contrast, for the cases with triangular arrangements, edge features influence the electronic structures significantly. This is rational with considering the symmetry and Lieb’s Theorem [32]. For such different patterns, the size of the defects plays a critical role in modulating the electronic structures of the systems.

On the other hand, Fig. 4 shows some mixture of a triangularly or hexagonally symmetric edges for the HOMO-1 and the HOMO states, corresponding to the triangular and the hexagonal arrangements. In fact, the case is the same for the LUMO and the level above the LUMO (LUMO+1). This is different from the previous literatures [27, 28], where triangular or hexagonal symmetry exists in the HOMO or LUMO states. This difference is ascribed to the symmetry of a system. In the previous literatures, the triangular or hexagonal holes were arranged in a hexagonal unit cell, which exhibited high symmetry. In contrast, in our concerned cases, the triangular or hexagonal holes are arranged in a rectangular unit cell, which makes the symmetry of the bipartite lattice low. For the cases of the triangular arrangements, only a mirror symmetry along $x$ axis exists in each system. While for the concerned hexagonal arrangements, there are two mirror symmetries along both $x$ axis and $y$
axis. This is displayed evidently in the isosurface of the charge density in Fig. 4. Therefore, it can be concluded that, in addition to the features and the size of the defect patterns, the symmetry of the patterns is another important factor to influence the electronic structures of the systems.

For the graphene with rectangular holes, the charge density displays some interesting features. As typical examples, the isosurface charge density for the cases of (8,4), (9,4) and (6,8) are plotted in Fig. 5(a-c) respectively. For the case of (8,4), the HOMO-1 orbital is localized in the separated regions which only belong to z-GNRs. This indicates that the electrons are confined in the matrix of z-NGRs. In contrast, either the HOMO or the LUMO presents a delocalized state with a major portion in a-GNRs. Such a nature implies that a-GNRs in the networks act as some special channels for electronic transport. Nevertheless, both the HOMO and the LUMO states exhibit local \( \pi \)-bond, of which the former is from two carbon dimer tilted to \( x \)-axis (or \( y \)-axis), while the latter is from that along \( x \)-axis (Fig. 5(a)). Differently, for the case of (9,4), the HOMO-1 orbital is localized in a-GNRs, and the HOMO and the LUMO states are localized in the crossing areas of z-GNRs and a-GNRs, as shown in Fig. 5(b). Meanwhile, both the HOMO and the LUMO also exhibit local \( \pi \)-bond, but the former is from two carbon dimer along \( x \)-axis, while the latter is from that tilted to \( x \)-axis (or \( y \)-axis). Fig. 5(c) is the isosurface charge density for the case of (6,8), which exhibits some new features. the HOMO-1 state having delocalized nature dominantly comes from the a-GNRs, which also acts as a channel for electronic transport. In contrast, the HOMO and LUMO states are all contributed from the atoms at the z-edges.

From above, we find that due to the quantum confinement effect, the electronic states near the Fermi level exhibit either localized or delocalized features, which is tightly related to the width of GNRs in the networks. As a result, tuning the electronic structures as well as wave function of electron may be realized by patterning the graphene sheet.

IV. CONCLUSION

Triangular, hexagonal and rectangular holes are arranged in a graphene sheet respectively. For these patterned graphene, we systematically investigate their electronic structures, based on the first principles calculations. We find that defect patterns with different features can modulate the electronic structures of the graphene significantly. For the cases of the trian-
gular holes with zigzag edges, the remarkable peaks appear near the Fermi level, with spin splitting, and the peak under the Fermi level converts to be two peaks with increasing the defect size. In contrast, for the triangular holes with armchair edges and the hexagonal holes, the majority and the minority spin states are degenerate. The graphene with hexagonal arrangements give rise to a narrow band gap. For the graphene sheet with rectangular holes, they all exhibit semiconducting behaviors. The band gap significantly depends on the width of a-GNRs, but weakly on that of z-GNRs. Moreover, such structural patterns can induce either localized or delocalized states near the Fermi level, and the latter gives rise to the special transport channels along the a-GNR or z-GNR in the networks.

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FIG. 1: (Color online) The geometries of the supercell for (a) triangular, (b) hexagonal, (c) rectangular arrays in graphene. (d) is geometry of the networks of graphene nanoribbons of (e).

FIG. 2: (Color online) The density of states (DOS) for the patterned graphene of (a) tri-1, (b) tri-2, (c) tri-3, (d) tri-4, (e) hex-1, (f) hex-2, (g) hex-3 and (h) hex-4. The black solid lines and the blue dot lines stand for total density of states of the patterned graphene and the corresponding perfect graphene respectively. The red dash lines refer to local density of states for the atoms at the defect edges. For (a-d), the upper and below curves in each figure stand for the majority spin and the minority spin states respectively. The Fermi level in each figure is shifted to zero.

FIG. 3: (Color online) The density of states (DOS) for the patterned graphene of (a) (7,4), (b) (8,4), (c) (9,4), (d) (6,6), (e) (6,7) and (f) (6,8). The black solid lines and the violet dot lines stand for total density of states of the patterned graphene and the corresponding perfect graphene respectively. The short red dash lines, the long green dash lines and the blue dash-dot lines refer to local density of states for the atoms at the z-edges, in the corner and at the a-edges respectively. The Fermi level in each figure is shifted to zero.
FIG. 4: (Color online) The isosurface charge density of HOMO-1, HOMO and LUMO for the patterned graphene of (a) the majority spin state and (b) the majority spin state of tri-2, (c) hex-3, (d) triangular arrangement with armchair edges, and (e) hexagonal arrangement with armchair edges respectively. The isosurface value is $\pm 0.05 \text{ eV/Å}^3$ (distinguished by orange and blue surfaces).

FIG. 5: (Color online) The isosurface charge density of HOMO-1, HOMO and LUMO for the patterned graphene of (a) (8,4), (b) (9,4) and (c) (6,8) respectively. The isosurface value is $\pm 0.05 \text{ eV/Å}^3$ (distinguished by orange and blue surfaces).
