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Published in:
Physical Review B Condensed Matter

Link to article, DOI:
10.1103/PhysRevB.86.245410

Publication date:
2012

Document Version
Publisher's PDF, also known as Version of record

Link back to DTU Orbit

Citation (APA):

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Graphene antidot lattice waveguides

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(Rceived 8 October 2012; published 11 December 2012)

We introduce graphene antidot lattice waveguides: nanostructured graphene where a region of pristine graphene is sandwiched between regions of graphene antidot lattices. The band gaps in surrounding antidot lattices enable localized states to emerge in the central waveguide region. We model the waveguides via a position-dependent mass term in the Dirac approximation of graphene and arrive at analytical results for the dispersion relation and spinor eigenstates of the localized waveguide modes. To include atomistic details we also use a tight-binding model, which is in excellent agreement with the analytical results. The waveguides resemble graphene nanoribbons, but without the particular properties of ribbons that emerge due to the details of the edge. We show that electrons can be guided through kinks without additional resistance and that transport through the waveguides is robust against structural disorder.

DOI: 10.1103/PhysRevB.86.245410  PACS number(s): 72.80.Vp, 73.22.–f, 73.21.Hb, 73.21.Cd

I. INTRODUCTION

Graphene, the two-dimensional allotrope of carbon first isolated in 2004,1,2 has emerged as a highly attractive material for future electronic devices. Graphene has exceptional electronic properties, as demonstrated in its extremely high carrier mobility,3 which even at room temperature is limited predominantly by impurity scattering.4 Already, extremely fast graphene-based transistors have been realized,5 and fabrication methods have emerged which allow for large-scale production of single-layered graphene.6 One key element of future graphene-based electronics is the ability to localize carriers in graphene wires, in order to facilitate electronic graphene networks. The most immediate way of realizing such wire structures is by cutting graphene into so-called graphene nanoribbons (GNRs).7 Quantum confinement will in general induce a band gap in such structures, the magnitude of which scales with the inverse of the width of the GNR. However, the exact atomistic configuration of the edge of the ribbon greatly influences the magnitude of this gap, with particular edge configurations resulting in vanishing gaps or localized edge states.8,9 An alternative to this is to rely on controlled generation of extended defects in graphene to act as metallic wires.10 Relying on results from two-dimensional electron gases (2DEGs) formed at the interface of semiconductor heterostructures, one could also imagine applying electrostatic gating to define wire geometries. Graphene presents an interesting challenge in this regard via the phenomenon of Klein tunneling,11 which makes it difficult to achieve carrier localization in graphene via ordinary gating. Electrons impinging on a potential barrier at close to normal incidence are transmitted with vanishing reflection, regardless of the height of the potential barrier. In spite of this, guiding of electrons via electrostatic gating has been demonstrated experimentally,12 albeit with the caveat that guiding is restricted to a specific range of wave vectors for which Klein tunneling is negligible. In contrast, a so-called mass term in graphene provides confinement that is a close analog of gate-defined localization in an ordinary 2DEG. This term originates from a Dirac description of graphene, which emerges as a low-energy approximation of a tight-binding (TB) model on the honeycomb lattice.13 Adding a diagonal term ±Δ to the resulting Hamiltonian, with the sign alternating between the two sublattices of graphene, the spectrum becomes that of gapped graphene, a semiconductor with an energy gap of twice the mass term. This staggered potential is commonly denoted a mass term due to the analogy of the low-energy carriers in graphene with massless Dirac fermions, which acquire a mass via such a term in the Hamiltonian.

In this paper, we propose realizing a waveguiding structure in graphene via graphene antidot lattices (GALs). GALs are nanostructured graphenes, which in their simplest description take the form of periodically perforated graphene structures.14 GALs have recently been produced experimentally with both electron-beam lithography15,16 and block copolymer lithography.17,18 The periodic perforation induces a band gap in graphene, rendering it semiconducting, and the resulting band structure closely resembles that of gapped graphene in the low-energy limit. GALs thus allow for the realization of a position-dependent mass term, as illustrated in Fig. I. The idea is to sandwich a region of pristine graphene between two GAL regions, the band gaps of which define an energy range for which localized guided modes are expected to emerge in the central region. This idea is closely analogous to how photonic waveguiding is realized in photonic crystal structures.19 We note that other methods beside GALs have been proposed for achieving gaps in graphene via superlattices, such as, e.g., patterned hydrogenation20 or superlattices of boron nitride islands embedded in graphene.21 Another alternative is to sandwich graphene between hexagonal boron nitride, which is predicted to induce significant band gaps in graphene.22

We denote a waveguide geometry as \((L, R)_{ac}^{W/Ac}\), where \(L\) is the sidelength of the hexagonal unit cell of the surrounding GAL, while \(R\) is the radius of the perforations, both in units of the graphene lattice constant, \(a = 2.46\ \text{Å}\). The width \(W\) of the waveguide is defined via \(N = W/w\), where \(w\) is the width of the enlarged GAL unit cell, as illustrated in the lower panel of...
Fig. 1. (Color online) Upper panel: conceptual illustration of a graphene antidot lattice (GAL) waveguide. A central region of pristine graphene is surrounded by GAL regions, the band gaps of which confine states to the waveguide region. Translational symmetry is assumed in the $y$ direction, which is along the longitudinal direction of the waveguide. Lower panel: geometry of a $[7,3]_2^2$ GAL waveguide. Black dots show the location of carbon atoms. Bloch boundary conditions are imposed on all boundaries. The dashed lines illustrate the enlarged GAL unit cell, the width of which we denote by $w$. The lowest-energy waveguide mode at the $\Gamma$ point, calculated via the tight-binding model, is illustrated with circles, the size of which shows the absolute value $|\psi(x,y)|$ of the (real-valued) eigenstate, while the color indicates the sign.

Fig. 1. The width of the waveguide is of course somewhat more ambiguous than in the case of GNRs, as we have no sharp edge defining the precise boundary between the region of pristine graphene and the surrounding antidot lattice. For denoting the geometries we simply take $W$ as the distance between the nearest edges of the two bordering antidot lattice unit cells. However, as we discuss below, the effective waveguide width is slightly larger than this. The width $w$ of the enlarged GAL unit cell depends on the orientation of the waveguide with respect to the graphene lattice, which we indicate with the superscript, with $zz$ (ac) denoting a waveguide with the longitudinal direction along the zigzag (armchair) orientation of the graphene lattice. Note that in both cases we choose the orientation of the GAL such that the superlattice basis vectors lie parallel to carbon-carbon bonds, to ensure that a band gap always exists for the GAL. The lower panel of Fig. 1 illustrates the geometry of a $[7,3]_2^2$ waveguide. For simplicity we restrict the width of the waveguide to be the same on both sides of the central waveguiding region. For $|\chi| > 1$ and zero otherwise. This equation has been derived from the Dirac Hamiltonian $H_K$, which emerges as a linearization of a TB model of graphene near the $K$ point. We will discuss the differences between the two inequivalent $K$ and $K'$ valleys of graphene below. Note that the sign of the mass term is arbitrary and that changing it has no physical significance but merely results in an interchange of the two spinor components. We stress, however, that the sign should be the same on both sides of the central waveguiding region. The Hamiltonian commutes with the $y$ component of the momentum operator, and we thus take spinor components of the form $\phi_A(\chi, \gamma) = f(\chi)e^{i\gamma}$ and $\phi_B(\chi, \gamma) = g(\chi)e^{i\gamma}$, where $k = kW/2$ is the dimensionless Bloch wave vector along the longitudinal direction of the waveguide. As the mass term is piecewise constant, the equations for the spinor components can be decoupled in each region of the waveguide structure. We look for bound states and thus take $e^2 < \delta^2$. The normalizable solutions for the first spinor component thus read $f(\chi) = A_\delta e^{i\beta\chi}$ for $|\chi| > 1$ and $f(\chi) = B\cos(\alpha\chi) + C\sin(\alpha\chi)$ for $|\chi| < 1$ while the second component is given via $g(\chi) = i(k - \delta_\gamma)f(\gamma)/[\epsilon + m(\chi)]$. Here, we have defined $\alpha = \sqrt{\epsilon^2 - k^2}$ and $\beta = \sqrt{\epsilon^2 + k^2 - e^2}$. The requirement of continuity of both spinor components at the boundaries of the central waveguide region leads to a transcendental equation for the energies, $\sqrt{\delta^2 - \alpha^2} = \alpha \tan 2\alpha$, regardless of which valley is considered (see below). The solution of the problem is thus closely reminiscent of that of an ordinary one-dimensional square-well potential, albeit with the crucial difference that for graphene $E \propto \alpha$ rather than $E \propto \alpha^2$, as a consequence of the linear dispersion relation of graphene. Also, we note that, contrary to the case of the Schrödinger equation, the derivative of the eigenstate spinors need not be continuous at the boundary.
B. Infinite mass limit

The standard textbook graphical solution suggests that there are $N$ bound states in the waveguide if $(N-1)\pi/2 < \delta < N\pi/2$. In general, the energies and spinor components of these states will need to be determined by numerical solution of the transcendental equation. However, in the limit of an infinite mass term, $\delta \to \infty$, the problem admits an analytical solution for the energies:

$$E_n^\infty(k) = s_n \left( k^2 + \frac{\pi^2}{4} \left( n + \frac{1}{2} \right)^2 \right)^{1/2}, \quad n = 0, 1, 2, \ldots$$

or, reverting to ordinary units,

$$E_n^\infty(k) = s_n \left( E^2_1(k) + \frac{\hbar^2 v_F^2 \pi^2}{W^2} \left( n + \frac{1}{2} \right)^2 \right)^{1/2}, \quad n = 0, 1, 2, \ldots$$

where $E_1(k) = \hbar v_F k$ is the bulk graphene dispersion relation and $s = \pm 1$. The dispersion relations of the waveguide modes thus resemble those of gapped graphene, with a mass term of $\Delta_{\text{eff}} = (\hbar v_F \pi / W k n + \frac{1}{2})$. Series expansion of the transcendental equation for the energies in the case of a finite mass term reveals that $\alpha = \pi / 2(n + 1/2)(1 + 1/(2b))^{-1}$, so the results in the infinite mass limit are expected to be valid when $E_1 > 2b \hbar v_F / W \simeq 1.278$ eV nm$^2$W$^{-1}$. Including the leading-order correction, the energies are given as

$$E_n^{(1)}(k) = E_n^\infty(k) - s_n \frac{2\hbar^2 v_F^2 \pi^2 (n + \frac{1}{2})^2}{W^2 \sqrt{k^2 W^2 + \pi^2 (n + \frac{1}{2})^2}} \times \frac{1}{E_1}.$$  

The eigenstate spinors in the infinite mass limit are most easily determined by using the boundary conditions derived by Berry, which set a phase relationship between the spinor components at the edge of the waveguide. The spinor components for $\kappa = 0$ and $\epsilon > 0$ can then be derived as

$$f_\alpha(\chi) = \begin{cases} \cos(\pi/2[n + \frac{1}{2}]\chi) & \text{for } n \text{ zero or even} \\ \sin(\pi/2[n + \frac{1}{2}]\chi) & \text{for } n \text{ odd} \end{cases},$$

while

$$g_\alpha(\chi) = \begin{cases} i \sin(\pi/2[n + \frac{1}{2}]\chi) & \text{for } n \text{ zero or even} \\ -i \cos(\pi/2[n + \frac{1}{2}]\chi) & \text{for } n \text{ odd} \end{cases},$$

for $|\chi| < 1$. Here, we have omitted normalization constants. We note that, because of the particle-hole symmetry of graphene, the eigenstates should be normalized separately on each sublattice. The spinors for the $\epsilon < 0$ eigenstates are given by exchanging $f_\alpha$ and $g_\alpha$. Interestingly, and in stark contrast to an ordinary infinite square-well potential, this shows that in the limit of infinite mass the charge density is evenly distributed within the waveguide, for states with vanishing wave vectors. We find that this holds true also in the more general case of a finite mass term, the main difference being that in this case the spinors extend slightly into the mass regions. Including a nonzero wave vector, the charge density is localized predominantly on either the edges or the middle of the waveguide.

The derivation above takes as its starting point a linearization of graphene near the $K$ valley. The Dirac Hamiltonian near the inequivalent $K'$ valley is such that the energy spectrum of the waveguide is the same for both valleys, while the eigenstate spinors are related by an interchange of the spinor components, i.e., $\phi_A^{K'} = \phi_B^K$ and $\phi_B^{K'} = \phi_A^K$. We also note that to obtain the full wave function, a Bloch phase factor of $e^{iK' \cdot r}$ or $e^{iK \cdot r}$ should be added to the spinors, with $K$ and $K'$ depending on the orientation of the graphene lattice. We will return to this point below, when we compare the results of the Dirac approximation with those obtained with the tight-binding model.

III. TIGHT-BINDING MODEL

To include the atomistic details of the waveguide structures, and to clarify the validity of the results derived in the Dirac approximation, we employ a nearest-neighbor tight-binding approximation. This will also allow us to assess the significance of the orientation of the waveguide with respect to the graphene lattice. The TB model is parametrized via a hopping term $t = -3$ eV between $\pi$ orbitals, the on-site energy of which we set to zero. We ignore nonorthogonality of the $\pi$ orbitals. Figure 1 illustrates the geometry used for the TB model in the case of a $[7,3]^{\infty}$ GAL waveguide. We use periodic boundary conditions along the $x$ axis as well as the $y$ axis. We have ensured that the results are converged with respect to the number of GAL unit cells included around the waveguide. The band gap of the GAL is quite well developed even with just a few rows of antidots, so including three GAL unit cells on each side of the waveguide usually yields converged results.

IV. RESULTS

A. Dispersion relations

In Fig. 2, we show the band structure of a $[7,3]^{\infty}$ GAL waveguide calculated using the TB model as well as the Dirac
approximation. Only electron \((E > 0)\) bands are shown. Both the TB and the Dirac model exhibit perfect electron-hole symmetry, so hole \((E < 0)\) bands simply follow from a sign change. For the Dirac results we take the effective width of the waveguide to be \(W_{\text{eff}} = (N + \frac{1}{2})w_u\), slightly larger than the definition used for denoting the waveguide geometries. Note that the wave vector is shown relative to the lattice constant of the GAL waveguide, which is \(\Lambda = 3La\) for the zigzag orientation and \(\Lambda = \sqrt{3}La\) for the armchair orientation. In the figure, the shaded gray region illustrates the projected bands of the GAL, which define the region below which localized waveguide states are expected to appear. This particular waveguide structure supports several localized states. Higher-lying band gaps also appear in the GAL, and we have confirmed that localized waveguide modes are also supported in these gaps. It is worth stressing that localized waveguide modes exist for all wave vectors in the first Brillouin zone. This is in contrast to the case of waveguides defined via electrostatic gating, where guided modes generally exist only for a limited range of wave vectors. The dispersions of the waveguide states agree very well between the TB and the Dirac model, as long as the wave vector is not too near the Brillouin zone edges. The largest deviations between the two models occur for energies close to the projected bands of the GAL, where coupling between the waveguide and the GAL states is pronounced. We note that the analytical result obtained in the infinite mass limit \(E_{\text{in}}^\infty\) is a very good approximation of the lowest waveguide mode. Including the first-order correction to \(E_{\text{in}}^\infty\) leads to near-perfect agreement with the full solution of the transcendental equation. Note that, as illustrated in the derivation of the Dirac result, the waveguide dispersion relation quite closely resembles that of gapped graphene, which in turn is approximately the same as bulk graphene for wave vectors \(k \gg \Delta_{\text{eff}}/(\hbar v_F)\). This is evident in the figure, where for comparison we also include the bulk graphene dispersion relation. The density of states (DOS) calculated using the TB model is shown in the right panel of Fig. 2. As expected from the Dirac approximation, the DOS closely resembles one-dimensional gapped graphene, i.e., \(g(E) \propto \Theta(E - \Delta_{\text{eff}})/(\hbar v_F (E^2 - \Delta_{\text{eff}})^2)\) with the Van Hove singularities characteristic of one-dimensionality clearly evident in the figure.

### B. Band gaps and effective masses

In Fig. 3(a), we show the energy of the lowest localized waveguide state at the \(\Gamma\) point as a function of the width of a \([7,3]_N^{\text{zz, ac}}\) waveguide. Results are shown for the TB model as well as the analytical results obtained in the Dirac equation approach. Note that, due to electron-hole symmetry, the band gap is twice this value. While the solution derived from the Dirac equation does not distinguish between zigzag and armchair orientation of the waveguide, the TB model predicts that there are some differences between the two cases. To illustrate this, we show results obtained for waveguides oriented along the zigzag (ZZ) and armchair (AC) directions, respectively. The inset of the figure illustrates that both ZZ and AC orientations exhibit a clear \(1/W\) dependence of the energies, as predicted from the Dirac approximation, provided the waveguide is sufficiently wide. We note that while differences do exist between AC and ZZ orientations these are rather small and could be attributed to a slightly different effective width of the waveguides in the two cases, due to the \(\pi/6\) difference in the orientation of the surrounding GAL with respect to the waveguide. Indeed, results of the Dirac approximation fit both orientations quite well, especially when including the first-order correction. The results of the Dirac model can be made to fit even better if we take into account the fact that the effective width of the waveguide is likely to be somewhat larger than the definition we have used (see Fig. 1). Indeed, introducing the same effective width in the Dirac model as we did for Fig. 2 results in even better agreement with TB results.

The absence of a well-defined edge means that the dependence of the properties of the guided modes on the width of the waveguide is much simpler than is the case for GNRs. In particular, GAL waveguides are always semiconducting, whereas in a nearest-neighbor TB model armchair GNRs alternate between metallic and semiconducting behavior depending on the exact width of the ribbon, while zigzag GNRs display dispersionless midgap states, localized on the edges. In Fig. 3(a), we show the energies for ZZ and AC GNRs, calculated via the TB model. We stress that the ZZ GNRs also contain dispersionless edge states at the Dirac point energy. Furthermore, AC GNRs are metallic for widths...
$W = (3p - 1)a$, with $p$ being an integer. To compare with the waveguide results, we have included only semiconducting AC GNRs in the figure. With these modifications, there is quite good agreement between the energies of GNRs and the GAL waveguide structures. GAL waveguides thus resemble ribbons without the particulars resulting from edge effects. As such, we speculate that the electronic properties of GAL waveguides may be easier to control experimentally.

In Fig. 3(b), we show the effective mass of the lowest waveguide state at the $\Gamma$ point as a function of the width. From the analytical Dirac results in the limit of an infinite mass term, we find an effective mass $m^{\text{eff}}_W = \hbar \pi (n + \frac{1}{2})/(\nu_F W)$. Again, we note that there is excellent agreement between the results obtained in the Dirac approximation and those from a TB model, for both orientations of the waveguide. Including the first-order correction, results from the Dirac model are in near-perfect agreement with the TB model. While electrons in pristine graphene have vanishing effective masses, the appearance of an effective mass term in the waveguide structures results in nonzero albeit still very small effective electron masses, which tend to zero in the limit of infinitely wide waveguides.

**C. Eigenstates**

To further compare the AC and ZZ waveguide orientations, we show in Fig. 4(a) the eigenstates corresponding to the lowest (positive) energy of $\{7,3\}^{(zz)}_5$ and $\{7,3\}^{(ac)}_3$ GAL waveguides, calculated at the $\Gamma$ point. Note that these have approximately the same waveguide widths. The lower panels in the figure show the absolute square of the wave function, with the color indicating the sublattice. These results demonstrate a crucial difference between the AC and ZZ orientations, namely, that while for the AC waveguide the wave function is distributed even across the two sublattices the ZZ waveguide exhibits pseudo-spin-polarization, with the wave functions of the two sublattices localized predominantly on opposite edges of the waveguide. We note that the lowest energy is doubly degenerate and that the second eigenstate (not shown) has the opposite pseudo-spin-distribution. The upper panels in the figure show the integrated probability density, $\rho(x) \equiv \int |\psi(x,y)|^2 dy$, along the transversal direction of the waveguide, with color indicating the sublattice. Note that, despite the lack of a clearly defined edge, the probability densities very closely resembles those of GNRs. The black lines in the upper panels show the total probability density, if a broadening of the order of the graphene lattice constant is included in order to account for the spatial extent of the $\pi$ orbitals. As predicted from the Dirac equation approach above, these results illustrate how the charge density is nearly uniformly distributed across the entire waveguide also for the higher-lying states shown in Fig. 4(b). This is in stark contrast to gate defined waveguides, which have wave functions more reminiscent of ordinary standing wave solutions.

In agreement with the Dirac results, we find that the uniform distribution only occurs at the $\Gamma$ point. For nonzero wave vectors the density becomes localized predominantly at the edges of the waveguide for the lowest state, as illustrated with dashed lines in the figure for $k\Lambda = \pi/2$. In contrast to this, the densities of the second-lowest states tend to localize in the center of the waveguide as the wave vector is increased.

To compare the wave functions with the spinor components derived via the Dirac equation, we first note that both the $K$ and the $K'$ points of graphene are folded onto the $\Gamma$ point of the waveguide structure. We thus expect the eigenstates to resemble linear combinations of the eigenstates in both valleys. The differences between the zigzag and armchair waveguides emerge due to the Bloch phase factors $e^{iKx}$ and $e^{iKr}$, which differ depending on the orientation of the graphene lattice. For the zigzag orientation, the integrated probability density $\rho(x)$ becomes a simple linear combination of the eigenstates belonging to each valley, $\rho(x) \propto |f_K(x) + f_{K'}(x)|^2$ and $\rho(x) \propto |g_K(x) + g_{K'}(x)|^2$. Using the expressions for the spinor components derived above, we find $\rho(x) \propto 1 \pm \sin(2\pi |n + \frac{1}{2}|x/W)$, with the sign depending on the sublattice, which is in excellent agreement with the TB results. In contrast to this, because of the mixing of the valleys the probability densities of the armchair orientation exhibit a rapidly oscillating term, with a period $2\pi/\Delta K$. These rapid oscillations are clearly evident in the right panels of Fig. 4. As shown in the figure, these rapid oscillations are quickly washed out if a small amount of broadening is included. In this case, we recover the nearly uniform charge distribution within the waveguide predicted from the Dirac results. Finally, we note that the differences between armchair and zigzag oriented GAL waveguides are very similar to those seen between GNRs with armchair and zigzag edges.

In the case of GNRs, though, the difference emerges due to different boundary conditions at the edge of the ribbon, which are not present in the case of GAL waveguides.
Because the GAL waveguides have no clearly defined edge, one might wonder whether the guiding properties of the waveguides would be relatively robust to disorder. Indeed, the crucial ingredient is the existence of a band gap in the surrounding GAL. As this gap essentially occurs due to interference between transmitted and reflected waves at the boundaries between the GAL waveguide and the graphene leads, we calculate the conductance at a finite temperature of $T = 100$ K. We consider a relatively high level of disorder, for which edge atoms are removed with a 5% probability. In Fig. 5(a), we show the conductance of disordered ($7,3$)$_{zz,ac}$ GAL waveguides of length $L = 89$ nm (dashed lines). For comparison, we also include the conductance of the pristine waveguides (solid lines). The shaded area indicates the energy range for the confined waveguide mode. In Fig. 5(b), we show the corresponding results for ZZ and AC GNRs.

Focusing first on the conductances for the pristine systems (solid lines), we observe that the ZZ and AC oriented GAL waveguides have a similar conductance in the energy range of the waveguide mode. The high conductances $G \sim 1.5 G_0$ show that there is a relatively good electronic coupling between the GAL waveguide and the graphene leads. The metallic ZZ GNR is conducting at all energies, but the conductance saturates at a value close to unity. In the energy range of the waveguide mode, the GAL waveguides thus have a higher conductance than both of the GNRs. Turning to the results for the disordered systems we observe that the GAL waveguides retain relatively high conductances. The ZZ and AC orientations show comparable reductions in the conductance due to disorder. The ZZ GNR is less affected by disorder, while the AC GNR conductance is significantly reduced except at the lowest energies. We conclude that the GAL waveguides appear to be relatively robust against structural disorder and in general have higher conductances than the corresponding GNR systems.

D. Conductance

Because the GAL waveguides have no clearly defined edge, one might wonder whether the guiding properties of the waveguides would be relatively robust to disorder. Indeed, the crucial ingredient is the existence of a band gap in the surrounding GAL. As this gap essentially occurs due to an averaging of the effect of the individual holes, the emergence of a gap should be relatively robust to disorder. A thorough investigation of disorder is beyond the scope of this paper, but as a preliminary study we model disorder by randomly removing atoms at the edges of the holes in the GAL. We consider a disordered GAL waveguide sandwiched between semi-infinite pristine graphene leads as illustrated in the inset of Fig. 5(a). For comparison we also consider the analogous system with the two graphene leads connected with a GNR having the same width as the corresponding waveguide ($W = 4.5$ nm for ZZ, $W = 6.0$ nm for AC), as illustrated in the inset of Fig. 5(b). We calculate the transmission through the waveguide and GNR using a recursive Green’s-function method with the lead self-energies determined using an iterative procedure. The transmission is averaged over 100 values of the transverse wave vector, and we further average over ten samples with different realizations of the random disorder. To smear out the rapid oscillations that occur due to interference between transmitted and reflected waves at the boundaries between the GAL waveguide and the graphene leads, we calculate the conductance at a finite temperature of $T = 100$ K. We consider a relatively high level of disorder, for which edge atoms are removed with a 5% probability. In Fig. 5(a), we show the conductance of disordered ($7,3$)$_{zz,ac}$ GAL waveguides of length $L = 89$ nm (dashed lines). For comparison, we also include the conductance of the pristine waveguides (solid lines). The shaded area indicates the energy range for the confined waveguide mode. In Fig. 5(b), we show the corresponding results for ZZ and AC GNRs.

E. Waveguide bends

As mentioned earlier, the waveguides introduced in this paper are closely analogous to photonic crystal waveguides. In such structures, light can be guided through bends in the waveguide with very little radiation loss. Relying on this analogy, we expect GAL waveguides to show a similar...
robustness to kinks along the waveguide. To illustrate the localization of the electronic state and the guiding properties of the GAL waveguide we show the local current through the confined waveguide mode of the straight waveguide. Note the nearly identical conductances of the two systems.

To further illustrate the strong guiding properties of GAL waveguides, we show in Fig. 7 the conductance of the waveguide bend illustrated in Fig. 6. A similar method was used as that for the results of Fig. 5. For comparison, we also show the corresponding conductance through a waveguide generated by omitting the kink in Fig. 6 and instead having the waveguide run straight through. These results show that, while there are small differences between the two structures in the oscillations of the conductance, overall the introduction of a kink has almost no consequence on the conductance through the GAL waveguide. Very low reflection loss is thus introduced by the kink, despite the fact that the waveguide alternates between zigzag and armchair orientations. We note that we have found similar results for slightly different waveguide structures.

V. DISCUSSION AND SUMMARY

The GAL waveguide systems studied in this paper represent idealized structures, which may be challenging to realize experimentally due to the small hole sizes. However, the applicability of the Dirac model allows for simulations of arbitrarily large structures. Moreover, the Dirac model can equally well be applied to other gapped graphene systems, where the band gap is not induced through periodic holes, but, e.g., via periodically absorbed hydrogen. Although the Dirac and TB models applied in this work are very simple, previous studies of pure GALs have shown that both the Dirac and TB models reproduce the trends obtained from more accurate density-functional theory calculations. In any case, the concept of a GAL waveguide depends only on the existence of a band gap in the GAL region and not on the specific details, and we believe our simplified models capture the correct physics.

In summary, we have introduced GAL waveguides. The band structures of GAL waveguides have been modeled with the Dirac model including a mass term, which is shown to be in excellent agreement with an atomistic tight-binding model. We have shown that GAL waveguides support modes which are highly confined to the waveguide region and are robust against structural disorder and kinks in the waveguide. In transport calculations, we find that GAL waveguides have higher conductances than corresponding graphene nanoribbons. A further advantage of the surrounding GAL may be that it will mechanically stabilize the structure and be able to carry some of the generated Joule heat away from the device. GAL waveguides may thus be an attractive way of realizing electronic wires in integrated graphene circuits.

ACKNOWLEDGMENTS

The work by J.G.P. is financially supported by the Danish Council for Independent Research, FTP Grants No. 11-105204 and No. 11-120941. T.M. acknowledges support from the Danish Council for Independent Research, FTP Grants No. 11-104592 and No. 11-120938. The Center for Nanostructured Graphene is sponsored by the Danish National Research Foundation.

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