Band gaps in graphene via periodic electrostatic gating

Jesper Goor Pedersen\textsuperscript{1,2} and Thomas Garm Pedersen\textsuperscript{1,2}
\textsuperscript{1}Department of Physics and Nanotechnology, Aalborg University, Skjernvej 4A DK-9220 Aalborg East, Denmark
\textsuperscript{2}Center for Nanostructured Graphene (CNG), Aalborg University, DK-9220 Aalborg East, Denmark

(Received 14 March 2012; revised manuscript received 24 April 2012; published 14 June 2012)

Much attention has been focused on ways of rendering graphene semiconducting. We study periodically gated graphene in a tight-binding model and find that, contrary to predictions based on the Dirac equation, it is possible to open a band gap at the Fermi level using electrostatic gating of graphene. However, comparing to other methods of periodically modulating graphene, namely, perforated graphene structures, we find that the resulting band gap is significantly smaller. We discuss the intricate dependence of the band gap on the magnitude of the gate potential as well as the exact geometry of the edge of the gate region. The role of the overlap of the eigenstates with the gate region is elucidated. Considering more realistic gate potentials, we find that introducing smoothing in the potential distribution, even over a range of little more than a single carbon atom, reduces the attainable band gap significantly. This represents a serious challenge to achieving gapped graphene via periodic gating.

DOI: 10.1103/PhysRevB.85.235432

I. INTRODUCTION

While graphene\textsuperscript{1} has proven to be a remarkable material, with electronic properties that are interesting from a fundamental\textsuperscript{2,3} as well as a technological viewpoint,\textsuperscript{4,5} the absence of a band gap severely limits its possible applications. Several methods have been proposed for opening a gap in graphene. Relying on quantum confinement effects, the most immediate way of making graphene semiconducting is by reducing the dimensionality by cutting graphene into narrow ribbons. Such so-called graphene nanoribbons (GNRs) have band gaps that in general scale inversely with the width of the GNR, but which are very sensitive to the exact geometry of the edge of the ribbon.\textsuperscript{5-8} Related to these ideas, periodically perforated graphene, termed graphene antidot lattices, effectively results in a network of ribbons and has been shown to be an efficient way of inducing an appreciable band gap in graphene.\textsuperscript{9} This idea has been successfully applied to fabricate simple graphene-based semiconductor devices.\textsuperscript{10,11} Modifying graphene via adsorption of hydrogen presents another route towards opening a gap in graphene, with fully hydrogenated graphene exhibiting a band gap of several electron volts,\textsuperscript{12,13} while patterned hydrogen adsorption yields band structures resembling those of graphene antidot lattices, with reported band gaps of at least 450 meV.\textsuperscript{14}

The prospect of opening a band gap in graphene via electrostatic gating is intriguing since it would allow for switching between semimetallic and semiconducting behavior and to dynamically alter the band gap to fit specific applications. This makes it significantly more flexible than proposals relying on structural modification of graphene. However, a linearization of the tight-binding Hamiltonian of graphene, resulting in the now widely studied Dirac equation (DE) of graphene,\textsuperscript{1,15} suggests that while magnetic confinement of the Dirac fermions of graphene is possible,\textsuperscript{16,17} they can not be confined by electrostatic gating, due to the phenomenon of Klein tunneling.\textsuperscript{18,19} Thus, while periodic gating of usual semiconductor heterostructures such as, e.g., GaAs quantum wells, does induce gaps in the dispersion relation,\textsuperscript{20} previous theoretical studies have indicated that band gaps are induced for neither one-dimensional\textsuperscript{21,22} nor two-dimensional\textsuperscript{23,24} periodic gating of graphene.

These studies have taken as their starting point the Dirac model of graphene, which is a low-energy continuum model, ignoring atomistic details. Here, we instead use a more accurate tight-binding (TB) model to study periodically gated graphene. For sufficiently smooth potentials, the low-energy electronic properties are expected to be well described by both the DE and TB models. However, a vital difference is that atomistic details are included in TB, whereas DE represents a continuum approximation. Consequently, in contrast to continuum (Dirac) models, we find that the TB model suggests that it is indeed possible to open a band gap in graphene via periodic gating, provided the potential is sufficiently abrupt. The aim of this paper is twofold: (i) to compare periodically gated graphene with graphene antidot lattices; in doing so, we will illustrate that, contrary to what may be expected from the Dirac equation, a sufficiently large scalar potential, i.e., not necessarily a mass term, yields a band structure that is highly similar to that of perforated graphene structures; (ii) to serve as a feasibility study of periodic gating as a means of inducing a band gap in graphene. To this end, we will illustrate and discuss the nontrivial dependence of the band gap on the gate potential, as well as the intricate relation between band gap and the edge geometry of the gated region. These results will also serve to illustrate some of the key differences between graphene and ordinary two-dimensional electron gases. While, initially, the potential will be modeled as a simple step function, we will show in the following that introducing smoothing in the potential distribution severely reduces the attainable band gap.

Continuum and atomistic models of periodically gated graphene have previously been compared in Ref. 25. That study, however, focused on a single value of the potential strength and only considered structures that are rotated 30° compared to those of this work and, therefore, do not necessarily display any band gap even for perforated structures.\textsuperscript{26} Moreover, in this work, we examine in detail the nontrivial dependence of the band gap on the magnitude of the potential and we consider more realistic, smooth potential profiles. Finally, we elucidate the intricate dependence on the precise
FIG. 1. (Color online) Unit cells used in the calculations for the \([12,5]\) lattice. (a) Perforated graphene sheet, with carbon atoms removed in the region of the antidot. (b) Staggered potential (mass term) in the antidot region. The color indicates the sign of the onsite energies. (c) Constant gate potential in the antidot region. (d) Gate potential modeled via Eq. (2), assuming the gate is directly below the graphene sheet, with no insulating layer in-between. The lower panel illustrates the potential of each model on the separate \(A\) and \(B\) sublattices.

edge geometry and show how the energy gap correlates with the gate region overlap of electron and hole states.

II. MODELS

In Fig. 1, we illustrate the graphene structures that we will consider in this article. We consider only superlattices with triangular symmetry, as shown in the figure. An important decision lies in the choice of the angle between the basis vectors of the superlattice and the carbon-carbon bonds in graphene. In particular, if the superlattice basis vectors are rotated \(30^\circ\) compared to the carbon-carbon bonds (such as in Ref. 25), Clar sextet theory predicts that perforated graphene structures only exhibit significant band gaps for every third value of the side length of the hexagonal unit cell.\(^{26}\) In contrast to this, perforated graphene structures with basis vectors parallel to the carbon-carbon bonds always have band gaps. We choose to focus in this paper on the latter geometries, in order to ensure that the superlattice symmetry in itself does not prohibit the emergence of a band gap.

We characterize a given structure by \(\{L,R\}\), where \(L\) denotes the side length of the hexagonal unit cell, while \(R\) is the radius of the central region, both in units of the graphene lattice constant, as illustrated in Fig. 1. In these units, \(L\) also corresponds to the number of benzene rings along each edge of the unit cell. Note that the exact geometry of the edge of the central region differs greatly depending on the radius \(R\). In the following, we discuss in detail the crucial dependence of the results on the edge geometry. We will consider four distinct ways of periodically modifying graphene: (a) perforated graphene (graphene antidot lattices), with carbon atoms removed from the central region, (b) a periodic mass term, nonzero only in the central region, and (c) periodically gated graphene, with a constant gate potential within the central region and a vanishing potential outside. Furthermore, to discuss the feasibility of realizing gapped graphene via periodic gating, we will also consider (d) periodically gated graphene, with a more realistic model of the spatial dependence of the gate potential, obtained from a solution to the Laplace equation. Focus will be on periodically gated graphene, with the other forms of modulation included for comparison only.

To illustrate the dependence of the results on the exact edge of the gate or mass region, we will use a Dirac model as well as a more accurate tight-binding treatment, in which the atomistic details of the structures are included. We find significant discrepancies between these two methods, quantitatively as well as qualitatively. In particular, we will show that the DE does not predict a band-gap opening for periodic gating, which is present in the TB results. In what follows, we briefly describe the two models. In the continuum model of the problem, we employ the Dirac Hamiltonian

\[
H_0 = \begin{bmatrix}
\Delta(x,y) & v_F (\hat{p}_x - i \hat{p}_y) \\
v_F (\hat{p}_x + i \hat{p}_y) & \pm \Delta(x,y)
\end{bmatrix},
\]

where \(v_F \approx 10^6 \text{ m/s}\) is the Fermi velocity, while \(\Delta(x,y)\) denotes the gate potential or mass term. Here, the \(+\) \((-\) is used when modeling a gate potential (mass term). Imposing periodic Bloch boundary conditions at the edge of the unit cell, we solve the problem in a plane-wave spinor basis \(\langle \mathbf{r} | A_G \rangle = (\frac{\gamma}{G_1}) e^{i(G+\mathbf{k}) \mathbf{r}}\) and \(\langle \mathbf{r} | B_G \rangle = (\frac{\gamma}{G_1}) e^{i(G+\mathbf{k}) \mathbf{r}}\), with \(\mathbf{k}\) the Bloch wave vector and \(G\) the reciprocal lattice vectors. We take \(\Delta(x) = V_0 \Theta(R - r)\), with \(\Theta(r)\) the Heaviside step function, yielding \(\Delta(G) = 2\pi R V_0 J_1(G R)/(G A)\), where \(A\) is the unit-cell area, while \(J_1(x)\) is the Bessel function of the first kind. A total of 1058 plane-wave spinors were included in the calculations to ensure convergence of the results.

In the tight-binding model, we include only nearest-neighbor coupling between \(\pi\) orbitals, parametrized via the hopping term \(-t\), with \(t = 3\) eV. We ignore the overlap between neighboring \(\pi\) orbitals, assuming that our basis is orthogonal, and set the onsite energy of the \(\pi\) orbitals to zero. This parametrization accurately reproduces the Fermi velocity of graphene, and is also in quantitative agreement with density functional theory when applied to perforated graphene structures.\(^{27}\) For periodically gated graphene, we set the diagonal terms of the Hamiltonian equal to the gate potential. In the case of a mass term, the diagonal terms become \(\pm V_0\), with the sign depending on which sublattice the carbon atom resides on. For perforated graphene, atoms are removed entirely in the region of the hole, ensuring that no dangling bonds are created. While including next-nearest-neighbor coupling, as well as taking into account the nonorthogonality of the basis set, will change our results quantitatively, we expect the overall trends and the conclusions to remain the same in more accurate models.

III. BAND STRUCTURES

In Fig. 2, we show the band structure for a \([12,5]\) graphene antidot lattice, i.e., periodically perforated graphene, and compare to the case of a periodic mass term, modeled using either the TB or the DE approach. A sufficiently large mass term should ensure that electrons are excluded entirely from the region of the mass term, and we thus expect relatively good correspondence with perforated graphene. In the figure, we consider the case where the mass term is equal in size to the TB hopping term \(V_0 = t\). As expected, we find quite
A periodic mass term is expected to induce a gap in graphene due to the fact that it explicitly breaks sublattice symmetry via the \( \sigma_z \) operator in the continuum model or, similarly, through the staggered onsite potential in the TB approach. Contrary to this, analysis of periodic potentials in a DE model of graphene suggests that periodic gating does not induce a gap in graphene around the Fermi level,\(^{21,22} \) but rather leads to the generation of new Dirac points near the superlattice Brillouin zone boundaries.\(^{23} \) Superlattices lacking inversion symmetry have been suggested as a means of achieving tunable band gaps in graphene, based on results using a DE model.\(^{24} \) However, these results were recently found to be based on numerical errors.\(^{25} \) Indeed, based on the DE model, a gap can not be produced by any Hamiltonian that preserves time-reversal symmetry, i.e., \( H = \sigma_z H^* \sigma_z \), where \( \sigma_z \) is the Pauli spin matrix while \( H^* \) denotes the complex conjugate (not the Hermitian conjugate) of the Hamiltonian.\(^{26} \) A pure scalar potential, such as the one we consider for periodically gated graphene, see Eq. (1), preserves this symmetry and the DE model thus suggests that periodic gating does not open a band gap. Instead, a combination of a scalar as well as a vector potential is needed,\(^{27} \) such as, e.g., in gated strained graphene structures, where the vector potential emerges from the pseudomagnetic field created by the strain.\(^{30} \)

In Fig. 3, we show the band structure of a periodically gated [12,5] graphene structure, with a gate potential of half the TB hopping term \( V_0 = t/2 \). Results are shown for TB and DE models, respectively. Contrary to a periodic mass term, we see that, as could be expected, periodic gating breaks electron-hole symmetry and shifts the Fermi level to higher energies. Comparing DE and TB results, we note that there is quite good agreement overall between the two methods. However, a crucial difference emerges when considering the bands in close vicinity of the Fermi level, as illustrated in the inset: while the DE results suggest that periodic gating does not open a band gap, TB results demonstrate that a band gap does occur right at the Fermi level. We attribute this to a local sublattice symmetry breaking at the edge of the gate region and substantiate this claim below. We note that while a band gap appears, the magnitude of the band gap is of the order of tens of meV, an order of magnitude smaller than that of the corresponding perforated graphene structure. This dramatic qualitative difference between TB and DE modeling agrees with previous results\(^{25} \) comparing density functional theory and Dirac models for rotated triangular geometries.

Above, we illustrated how a sufficiently large mass term serves as an excellent model of a hole in graphene (see Fig. 2). Because a simple scalar potential can not confine Dirac electrons,\(^{18,19} \) one would expect that modeling the hole via a large gate potential would be inaccurate. In Fig. 4, we show the band structure of periodically gated graphene, with a very large gate potential of \( V_0 = 10t \). While this is a very large value of the gate potential, our point here is not whether such a periodic gate can be realized, but rather that there is theoretical agreement between this model and that of perforated graphene. To demonstrate this, we also show the corresponding perforated graphene structure. Contrary to the aforementioned expectations, we see that the periodically gated graphene structure is an excellent model of perforated graphene with a periodic mass term of \( V_0 = t \), calculated using the TB (DE) model. The thick, red line shows the location of the Fermi level. Note the nearly dispersionless band near \(-0.2 \) eV. Inset: A zoom of the band structure near the \( \Gamma \) point, illustrating the emergence of a band gap in the TB results and the absence of such a gap in the DE model.

FIG. 2. (Color online) Band structures of the [12,5] lattice. The solid, black lines show results for perforated graphene, calculated using a TB model. The blue, dashed (red, dotted) lines correspond to graphene with a periodic mass term of \( V_0 = t \), calculated using the TB (DE) model. The thick, red line shows the location of the Fermi level. Note the perfect electron-hole symmetry in this case, and the agreement on the magnitude of the band gap between all three methods.

FIG. 3. (Color online) Band structures of a periodically gated [12,5] lattice. The solid, black (blue, dashed) lines show results for periodically gated graphene, calculated using a TB (DE) model. The gate potential is \( V_0 = t/2 \). The thick, red line shows the location of the Fermi level. Note the nearly dispersionless band near \(-0.2 \) eV. Inset: A zoom of the band structure near the \( \Gamma \) point, illustrating the emergence of a band gap in the TB results and the absence of such a gap in the DE model.
graphene. We note that increasing the gate potential further results in near-perfect agreement between the periodically gated and the perforated structures. With a gate potential of \( V = 10t \), we are way beyond the linear regime of the band structure, for which a Dirac treatment of graphene is viable, which explains why the theoretical arguments pertaining to Dirac electrons break down in this case. We stress that the Fermi energies of the two structures are very different, and that no band gap occurs at the Fermi energy of the gated structure. However, the results indicate that the bands near the Fermi energy of perforated graphene can be quite accurately modeled via a periodic gate potential, provided the magnitude of the gate potential is significantly larger than the hopping term.

A. Gate region overlap

Returning now to the band structure for the periodically gated \( \{12,5\} \) lattice, shown in Fig. 3, we note the appearance of a nearly dispersionless band near \(-0.2\) eV. This state is localized predominantly within the gate region. In the upper panel of Fig. 5, we show the overlap of all eigenstates with the gate region, i.e., \( \int_{\text{gate}} d\vec{r} |\Psi(\vec{r})|^2 \), as a function of energy, calculated at the \( \Gamma \) point. For comparison, we show the corresponding results for a periodic mass term in the lower panel. We note that several states exist, which have significant overlap with the gate region, also at energies below the Fermi level. An example of one such state is shown in the figure. As the gate potential is increased further, these states become less energetically favorable, and are eventually all situated at energies well above the Fermi level. In stark contrast to this, a periodic mass term dictates perfect electron-hole symmetry, and thus always predicts states below the Fermi level having significant overlap with the gate region. In fact, as the mass term is increased, states nearly entirely localized within the mass term region develop at both extrema of the spectrum. Below, we will illustrate how this fundamental difference between a mass term and a scalar potential manifests itself via the dependence of the band gap on the magnitude of the gate potential for periodically gated graphene.

IV. BAND GAPS IN PERIODICALLY GATED GRAPHENE

Having determined that a TB treatment of periodically gated graphene does indeed suggest the opening of a band gap at the Fermi level, we now proceed to investigate the behavior of the band-gap magnitude in more detail. From here on, all results shown have been calculated using the TB model.

In Fig. 6, the solid lines show the magnitude of the band gap at the Fermi level for three different lattices, \( \{7,3\} \), \( \{12,5\} \), and \( \{15,6,3\} \), all of which have approximately the same ratio \( R/L \) of gate radius to side length of the hexagonal unit cell. When plotted against the gate radius times the gate potential, the resulting curves emerge as simple scaled versions of each other, as seen in Fig. 6. While, initially, raising the gate potential increases the band gap, a maximum gap is reached at a certain gate potential, after which the band gap diminishes. This behavior is completely different from the case where the potential is replaced by a mass term, as illustrated in the inset of the figure. In this case, the band gap continues to increase until a saturation point is reached in the limit where the structure resembles that of perforated graphene. While the three periodic lattices indicated with solid lines in Fig. 6 result in similar dependencies of the gap on \( RV_0 \), we stress that this is not the case for all lattices, even if the ratio \( R/L \) is approximately the same. To illustrate this, we also show in Fig. 6 results for the \( \{7,2,8\} \) lattice. The dependence of the band gap on gate potential differs markedly for this lattice. This indicates that the exact geometry at the edge of the gate region plays a large role in determining the band gap, in agreement with findings in Ref. 25.
Further. Around $RV_0 \simeq 3t$, the band gap changes from direct ($\Gamma$–$\Gamma'$) to indirect ($\Gamma$–$K$) as the gate voltage is raised. The dashed lines in the figure illustrate the $\Gamma$–$\Gamma'$ gap above the direct- to indirect-gap transition. However, after a slight further increase of the gate voltage, the final closing of the band gap occurs as the energy at the $K$ point moves below that at the $\Gamma$ point, resulting in crossing bands at the Fermi level. Finally, we note that while the three lattices show similar behavior, the dependence of the band gap on the radius of the gate region is clearly not monotonic, and a larger gate region does not necessarily result in a larger band gap.

In contrast to the similarities of the other three structures, the dependence of the band gap on the gate potential for the (7,2,8) lattice differs greatly. In the upper panel of Fig. 7 we show the unit cells corresponding to the (7,R) lattices considered, with the edge geometries highlighted. The (7,2,8) lattice stands out from the rest of the geometries in that the entire edge of the gate region is made up from several pure zigzag edges. We stress that the sublattice imbalance for the entire edge is zero, while there is a local sublattice imbalance on the individual straight zigzag edges. In contrast to this, the other geometries have gate regions with zigzag as well as armchair edges. We find that the general trend is for zigzag edges to quench the band gap of the periodically gated graphene structures, which we have also verified via calculations of gate regions of hexagonal symmetry, which always have pure zigzag edges. This trend can be explained by noting that pure zigzag edges, such as, e.g., in zigzag graphene nanoribbons$^{53}$, or graphene antidot lattices with triangular holes,$^{32–35}$ lead to localized midgap states.$^{35}$ For periodically gated graphene, the edge is defined by a finite potential rather than a complete absence of carbon atoms, so we expect the tendency of electrons to localize on the edge to be less pronounced. Nevertheless, our findings suggest that local zigzag geometry still has the effect of quenching the band gap. Since, in general, larger circular holes will have longer regions of zigzag geometry at the edge of the gate region, this explains why larger gate regions will not invariably lead to larger band gaps. In the present case, we note that the (7,3,0) structure indeed has a significantly smaller band gap than the (7,2,5) structure. The (7,3,0) lattice is unique in that the equivalent of dangling bonds are present at the edge of the gate region, which further decreases the magnitude of the band gap.

### B. Dependence on gate region overlap

First-order perturbation theory suggests that the dependence of the energy of the eigenstate on the gate potential be proportional to the overlap of the state with the gate region, i.e., $\frac{\partial E}{\partial V_0} \propto \int_\text{gate} |\Psi(\mathbf{r})|^2$. We thus expect the overlap with the gate region of the two eigenstates closest to the Fermi level to be a crucial parameter in describing the opening and quenching of the band gap as the gate voltage is varied. We will also see that it illustrates the crucial differences between graphene and ordinary two-dimensional electron gases. In Fig. 8, we show the overlap of the eigenstate with the gate region as a function of the magnitude of the potential. The overlap is shown for the eigenstates at the valence- and conduction-band edges, and normalized by the ratio between gate and unit-cell areas. A value of one thus indicates that the overlap with the gate...
region is the same as if the eigenstate is evenly distributed throughout the unit cell, while a value larger (smaller) than 1 suggests that the eigenstate is localized predominantly inside (outside) the gate region. As we saw also in Fig. 5, the states near the Fermi level both have quite large overlaps with the gate region, even when the potential is of the order of the TB hopping term. Initially, for low values of the gate potential, the overlap with the gate region of the unoccupied state in the conduction band is larger than the corresponding overlap of the occupied state in the valence band. Relying on first-order perturbation theory, we thus expect the energy of the conductance-band state to increase more strongly with the gate potential than the valence-band state, resulting in a larger band gap as the gate potential is raised, as illustrated in the left inset of Fig. 8. However, contrary to what would be expected for an ordinary two-dimensional electron gas, we see that as the potential is increased further, the valence-band state also becomes localized predominantly within the gate region. Indeed, eventually the overlap of the valence-band state with the gate region becomes larger than the one of the conduction-band state, which results in a quenching of the band gap as the potential is increased further, as illustrated in the right inset of Fig. 8. We note that the point where the overlap of the two states with the gate region become equal exactly matches the point where the band gap is at a maximum. This is illustrated by the vertical, black line in the figure. The strong influence of the exact edge geometry is apparent, manifesting itself in a qualitatively different dependence of the overlap on gate voltage for the \{7, 2.8\} lattice. In particular, while the gate region overlap of the valence-band state of the \{7, 2.5\} and \{7, 3.3\} lattices initially decreases with the size of the potential, both valence- and conduction-band states immediately start localizing within the gate region for the \{7, 2.8\} structure. This leads to much faster quenching of the initial band gap.

C. Realistic potential profiles

As we have illustrated above, the band gap of periodically gated graphene depends strongly on the edge geometry at the boundary between the gated and the nongated regions. So far, we have used a simple step function to model the spatial dependence of the potential due to the gate. However, it is obvious that in actual realizations of periodically gated graphene, some form of smoothing of the potential will inevitably be present. Due to the intricate relationship between the band gap and the edge geometry, it is relevant to investigate the effect of smoothing out the potential. In particular, since the DE model predicts no gap at all, one may wonder whether smoothing will cause the gap to close entirely. Previous studies have included smoothing of the gate potential, but with a smearing distance of the order 0.1 Å,25 small enough that an atomically resolved edge can still be defined.

To model a more realistic gate potential, we use an analytical expression for the potential distribution resulting from a constant potential disk in an insulating plane, obtained by direct solution of the Laplace equation. In cylindrical coordinates, this reads as\textsuperscript{36}

\[ V(r, z) = \frac{2V_0}{\pi} \sin^{-1}\left(\frac{2R}{\sqrt{(r - R)^2 + z^2} + \sqrt{(r + R)^2 + z^2}}\right), \]

with \(z\) the distance above the gate, while \(r\) is the distance from the center of the disk. Note that for \(z = 0\), the expression simplifies to \(V(r, 0) = V_0\) for \(r \leq R\), while \(V(r, 0) = 2V_0 \sin^{-1}(R/r)/\pi\) for \(r > R\). Of course, more exact approaches such as, e.g., finite-element methods, could be used to determine the potential distribution from a realistic back gate. However, we choose to use this relatively simple analytical expression since we are mainly interested in discussing the general trends that occur as the edge of the potential region becomes less well defined. One could imagine more elaborate setups that would generate sharper potential distributions. To include such possibilities, we consider a modified potential distribution \(V(r, z) = V_0|V(r, z)/V_0|^\eta\), with the additional parameter \(\eta\), which allows us to control the smoothing of the potential further. As \(\eta \to \infty\), we approach the limit where the potential is described by a Heaviside step function, as in the results presented so far. We note that Eq. (2) is derived for an isolated constant potential disk rather than a periodic array of gates. Ignoring coupling between the different gates, one simple way of improving this model would be to add the potentials generated from the nearest-neighbor gates to account for the overlap between them. However, this would merely serve to smoothen out the potential further, as well.
The radial position of the carbon atoms.

as add a constant background potential, effectively decreasing the height of the potential barrier. Here, we are interested in illustrating the critical dependence of the band gap on smoothing out the potential, so we are adopting a “best-case” scenario, which also means that we will use \( z = 0 \) throughout, assuming that the graphene layer is deposited directly on the periodic gates, with no insulating layer in-between.

In Fig. 9, we show the band gap for a \{12,5\} lattice as a function of gate potential, for increased values of the \( \eta \) parameter. While for \( \eta \to \infty \), corresponding to a Heaviside step function distribution, the maximum band gap is about 33 meV, the band gap for \( \eta = 1 \) is drastically lower, with a maximum value of only 0.9 meV. As we artificially decrease the amount of smoothing by raising the value of \( \eta \), we slowly recover the maximum band gap attainable. However, we stress that even for \( \eta = 20 \), which as shown in the inset of the figure amounts to smoothing over a range of little more than a single carbon atom, the maximum band gap has decreased by more than 20% from the value at \( \eta \to \infty \). This suggests that the band gap does indeed critically depend on an edge effect, which is very quickly washed out as the potential step is smoothed out over several carbon atoms. This is in agreement with previous studies, which have indicated that intervalley scattering is crucial in describing the band gap of periodically gated graphene. In order for a scalar potential to induce intervalley scattering, it must vary significantly on a scale of the carbon-carbon distance, so that a local sublattice asymmetry is introduced.

\[ \text{FIG. 9. (Color online) Band gap as a function of gate potential for the \{12,5\} periodically gated graphene lattice. The potential distribution due to the periodic gates is modeled via Eq. (2). We assume the distance from the plane of the gate to the graphene layer is zero. Results are shown for increased values of the } \eta \text{ parameter, which determines the strength of the smoothing. The inset illustrates the potential distribution } V / V_0 \text{ for each case, with markers indicating the radial position of the carbon atoms.} \]

V. SUMMARY

By employing a tight-binding description of graphene, we have shown that, contrary to what is predicted on the basis of a continuum model, it is indeed possible to induce a band gap in graphene via periodic, electrostatic gating. Further, if the magnitude of the potential is made sufficiently large, periodically gated graphene is an accurate model for perforated graphene structures, with one caveat, namely, that the Fermi level is far removed from the location of the band gap. For smaller, more realistic values of the gate potential, a band gap appears right at the Fermi level. However, we find that the band gap is orders of magnitude smaller than that of the corresponding perforated graphene structure.

The dependence of the band gap on the gate potential is highly nontrivial, and entirely different from the case where graphene is modulated by a periodic mass term. In particular, a maximum magnitude of the band gap is reached, after which increasing the gate potential further quenches the gap. Also, a transition from a direct (\( \Gamma - \Gamma \)) to an indirect (\( \Gamma - K \)) semiconductor occurs for larger gate potentials. The exact magnitude and dependence of the band gap on gate potential depends critically on the precise geometry of the edge of the gate region. In particular, large regions of local zigzag geometries tend to result in significantly smaller band gaps than geometries where armchair edges dominate.

Because the emergence of a band gap relies on a local sublattice asymmetry, we find that it is extremely fragile. If smoothing is introduced in the potential distribution, such that the edge of the gate region is no longer atomically resolved, the magnitude of the band gap drops significantly. Even if the smoothing occurs over a range of little more than a single carbon atom, we find that the maximum band gap decreases to less than 80% of the value for a perfectly defined edge. This presents a serious challenge to opening a band gap in graphene via periodic gating, and suggests that periodic electrostatic gating as discussed in this paper does not currently represent a feasible method of achieving gapped graphene.

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. The Center for Nanostructured Graphene (CNG) is sponsored by the Danish National Research Foundation. We thank A.-P. Jauho for helpful comments during the development of the manuscript.


