Localized states at zigzag edges of bilayer graphene

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We report the existence of zero energy surface states localized at zigzag edges of bilayer graphene. Working within the tight-binding approximation we derive the analytic solution for the wavefunctions of these peculiar surface states. It is shown that zero energy edge states in bilayer graphene can be divided into two families: (i) states living only on a single plane, equivalent to surface states in monolayer graphene; (ii) states with finite amplitude over the two layers, with an enhanced penetration into the bulk. The bulk and surface (edge) electronic structure of bilayer graphene nanoribbons is also studied, both in the absence and in the presence of a bias voltage between planes.

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Introduction: The quest for new materials and material properties has recently led to graphene, the missing two-dimensional (2D) allotrope of carbon\textsuperscript{1}. Stability and ballistic transport on the submicrometre scale, even at room-temperature, make graphene based electronics a promising possibility\textsuperscript{2}. Indeed, with Si-based technology approaching its limits, a truly 2D material with unconventional electronic properties is regarded with great expectations.

Graphene is a zero-gap semiconductor, and this prevents standard logic applications where the presence of a finite gap is paramount. Band gaps can still be engineered by confining graphene electrons in narrow ribbons\textsuperscript{1, 3, 4}. However, the lateral confinement brings about the presence of edges, which in graphene can have profound consequences on electronics. This is essentially due to the rather different behavior of the two possible (perfect) terminations in graphene: zigzag and armchair. While zigzag edges support localized states, armchair edges do not\textsuperscript{3, 5, 6}. These edge states occur at zero energy, the same as the Fermi level of undoped graphene, meaning that low energy properties may be substantially altered by their presence. The self-doping phenomenon\textsuperscript{8} and the edge magnetization with consequent gap opening\textsuperscript{9} are among edge states driven effects.

Bilayer graphene, as its single single layer counterpart, is also a zero gap semiconductor\textsuperscript{10}, but only in the absence of an external electric field: the electronic gap can be tuned externally\textsuperscript{11}. Nevertheless, the question regarding the presence of edge states in bilayer graphene is pertinent. Firstly, zigzag edges are among the possible terminations in bilayer graphene, and secondly, the presence of edges is unavoidable in tiny devices.

In the present paper we show that zero energy edge states do exist at zigzag edges of bilayer graphene. An analytic solution for the wavefunction is given assuming a semi-infinite system and a first nearest-neighbor tight-binding model. The analytic solution we have found defines two types of edge states: monolayer edge states, with finite amplitude on a single plane; and bilayer edge states, with finite amplitude on both planes, and with an enhanced penetration into the bulk. A schematic representation of the two families of edge states can be seen in Fig. 1. We also show that bilayer graphene nanoribbons with zigzag edges have four flat bands occurring at zero energy, consequence of the two families of edge states localized on each edge. In the case of a biased ribbon, where the two planes are at different electrostatic potential and a band gap develops for the bulk electronic structure, the spectrum still shows two flat bands while the other two give rise to level crossing inside the gap.

Surface states in semi-infinite bilayer graphene: The study of edge states in $AB$—stacked bilayer graphene given here is based on the ribbon geometry with zigzag edges shown in Fig. 2. We use labels 1 and 2 for the top and the bottom layers, respectively, and labels $A_i$ and $B_i$ for each of the two sublattices in layer $i$. Each four-atom unit cell (parallelograms in Fig. 2) has integer indices $m$ (longitudinal) and $n$ (transverse) such that $m a_1 + n a_2$ is its position vector, where $a_1 = a(1,0)$ and $a_2 = a(1,−\sqrt{3})/2$ are the basis vectors and $a ≈ 2.46\text{ Å}$ is the lattice constant. The simplest model one can write to describe non-interacting electrons in $AB$-stacked bilayer is the first nearest-neighbor tight-binding model given by,

\begin{equation}
H = - t \sum_{m,n} \sum_{i=1}^{2} \sum_{m,n} a_{i;m,n}^{\dagger} (b_{i;m,n} + b_{i;m-1,n} + b_{i;m,n-1}) - t \sum_{m,n} a_{1;m,n}^{\dagger} b_{2;m,n} + \text{h.c.},
\end{equation}

FIG. 1: Charge density representation for the two families of edge states in bilayer graphene at $ka/2\pi ≈ 0.35$: monolayer (a), given by Eq. (12); and bilayer (b), given by Eq. (13).
where \(a_{i,m,n}(b_{i,m,n})\) is the annihilation operator for the state in sublattice \(A_i (B_i), i = 1, 2,\) at position \((m, n)\). The first term in Eq. (1) describes in-plane hopping while the second term parametrizes the inter-layer coupling \((t_\perp/t \ll 1)\). Without loss of generality we assume that the ribbon has \(N\) unit cells in the transverse cross section \((y)\) direction with \(n \in \{0, \ldots, N - 1\}\), and we use periodic boundary conditions along the longitudinal direction \((x)\) direction. This last simplification enables the diagonalization of Hamiltonian (1) with respect to the \(m\) index just by Fourier transform along the longitudinal direction, \(H = \sum_k H_k\), with \(H_k\) given by,

\[
H_k = -t \sum_{i=1}^2 \sum_n a_i^{\dagger} e^{i ka} b_{i,k,n} + h.c., \quad (2)
\]

In order to search for zero-energy edge states we solve the Schrödinger equation, \(H_k |\mu, k\rangle = E_{\mu,k} |\mu, k\rangle\), for \(E_{\mu,k} = 0\). First we note that Hamiltonian \(H_k\) in Eq. (2) effectively defines a 1D problem in the transverse direction of the ribbon. It is then possible to write any eigenstate \(|\mu, k\rangle\) as a linear combination of the site amplitudes across the cross section,

\[
|\mu, k\rangle = \sum_{n} \sum_{i=1}^2 \left[ a_{i,k,n} |a_i, k, n\rangle + \beta_{i,k,n} |b_i, k, n\rangle \right], \quad (3)
\]

where the four terms per \(n\) refer to the four atoms per unit cell, to which we associate the one-particle states \(|a_i, k, n\rangle = e^{i k a} |0\rangle\), with \(c_i = a_i, b_i,\) and \(i = 1, 2\). In addition we require the following boundary conditions,

\[
\alpha_1(k,N) = \alpha_2(k,N) = \beta_1(k,-1) = \beta_2(k,-1) = 0, \quad (4)
\]

accounting for the finite width of the ribbon. Then, analogously to the single layer case, it can be straightforwardly shown that if Eq. (3) is a zero energy solution of the Schrödinger equation, the coefficients satisfy the following matrix equations:

\[
\begin{bmatrix}
\alpha_1(k, n+1) \\
\alpha_2(k, n+1)
\end{bmatrix} = e^{-i k a} 
\begin{bmatrix}
D_k & 0 \\
-\frac{1}{t_\perp} e^{-i k a} D_k
\end{bmatrix} 
\begin{bmatrix}
\alpha_1(k, n) \\
\alpha_2(k, n)
\end{bmatrix}, \quad (5)
\]

\[
\begin{bmatrix}
\beta_2(k, n-1) \\
\beta_1(k, n-1)
\end{bmatrix} = e^{i k a} 
\begin{bmatrix}
D_k & 0 \\
-\frac{1}{t_\perp} e^{i k a} D_k
\end{bmatrix} 
\begin{bmatrix}
\beta_2(k, n) \\
\beta_1(k, n)
\end{bmatrix}, \quad (6)
\]

where \(D_k = -2 \cos(ka)/2\). As the \(2 \times 2\) matrix in Eqs. (5) and (6) has the following property for any complex \(p_k\),

\[
\begin{bmatrix}
D_k & 0 \\
p_k & D_k
\end{bmatrix}^n = 
\begin{bmatrix}
D_k^n & 0 \\
p_k D_k^{n-1} D_k^n & D_k^n
\end{bmatrix}, \quad (7)
\]

we conclude by induction that the general solution of Eqs. (5) and (6) has the form:

\[
\begin{bmatrix}
\alpha_1(k, n) \\
\alpha_2(k, n)
\end{bmatrix} = e^{-i \frac{ka}{t_\perp} n} T_n \begin{bmatrix}
\alpha_1(k, 0) \\
\alpha_2(k, 0)
\end{bmatrix}, \quad (8)
\]

\[
\begin{bmatrix}
\beta_2(k, N - n - 1) \\
\beta_1(k, N - n - 1)
\end{bmatrix} = e^{i \frac{ka}{t_\perp} n} T_n \begin{bmatrix}
\beta_2(k, N - 1) \\
\beta_1(k, N - 1)
\end{bmatrix}, \quad (9)
\]

for \(n \geq 1\), where the matrix \(T_n\) is given by,

\[
T_n = 
\begin{bmatrix}
D_k^n & 0 \\
-n^2 D_k^{n-1} & e^{i \frac{ka}{t_\perp} n} D_k^n
\end{bmatrix}, \quad (10)
\]

and \(T_n^*\) is the matrix whose elements are the complex conjugate of \(T_n\). One also requires the convergence condition \(|-2 \cos(ka)/2| < 1\), which guarantees that Eq. (4) is satisfied for semi-infinite systems. It is easy to see that the semi-infinite bilayer sheet has edge states for \(k\) in the region \(2\pi/3 < ka < 4\pi/3\), which corresponds to \(1/3\) of the possible \(k\)'s, as in the graphene sheet. The next question concerns the number of edge states. As any initialization vector is a linear combination of only two linearly independent vectors there are only two states per edge (per \(k\)). Moreover, Eqs (8) and (9) are edge states solutions on different sides of the ribbon. When the semi-infinite system is considered only one of them survives. In particular, taking the limit \(N \rightarrow \infty\), and choosing the simplest linear independent initialization vectors \(|\alpha_1(k, 0), 0\rangle\) and \(|0, \alpha_2(k, 0)\rangle\), the two possible edge states are,

\[
\begin{bmatrix}
\alpha_1(k, n) \\
\alpha_2(k, n)
\end{bmatrix} = \alpha_1(k, 0) D_k^n e^{-i \frac{ka}{t_\perp} n}, \quad (11)
\]

and

\[
\begin{bmatrix}
\alpha_1(k, n) \\
\alpha_2(k, n)
\end{bmatrix} = \alpha_2(k, 0) D_k^n e^{-i \frac{ka}{t_\perp} n}, \quad (12)
\]

Although linearly independent, it is clear that the edge states (11) and (12) are not orthogonal, except for \(ka = \pi\). It is convenient to orthogonalize (11) with respect to (12) so that we finally obtain,

\[
\begin{bmatrix}
\alpha_1(k, n) \\
\alpha_2(k, n)
\end{bmatrix} = \alpha_1(k, 0) D_k^n e^{-i \frac{ka}{t_\perp} n}, \quad (11)
\]

\[
\begin{bmatrix}
\alpha_2(k, n) \\
\alpha_2(k, n)
\end{bmatrix} = -\alpha_1(k, 0) D_k^n e^{-i \frac{ka}{t_\perp} n} \left(n - \frac{D_k}{1 + D_k^2} \right), \quad (13)
\]

which, together with Eq. (12), represent all possible orthonormalized zero-energy edge states for bilayer graphene. The normalization constants in Eqs. (12) and (13) are given by,

\[
|\alpha_1(0, k)|^2 = \frac{(1 - 2 D_k^3)^3}{(1 - D_k^2)^2 + \frac{t_\perp^2}{t_\perp^2}}, \quad (14)
\]

\[
|\alpha_2(0, k)|^2 = 1 - D_k^2, \quad (15)
\]
An example of the charge density associated with Eq. (13) is shown in Fig. 3 for $t_\perp/t = 0.2$, where the $|\alpha_1(k, n)|^2$ dependence can also be seen as the solution given by Eq. (12) for $|\alpha_2(k, n)|^2$, apart from a normalization factor.

The solution given by Eq. (12) is exactly the same as that found for a single graphene layer [5, 6], where the only sites with non-zero amplitude belong to the $A$ sublattice of layer 2, the one disconnected from the other layer. Solution (13), on the other hand, is an edge state that can only be found in bilayer graphene. The sites of non-vanishing amplitude for this edge state occur at sublattice $A$ of layer 2, and at sublattice $A$ of layer 1, which is connected to the other layer through $t_\perp$ (see Fig. 1). Had we increased the semi-infinite sheet from the other side of the ribbon, and two similar edge states would have appeared in the opposite edge with non-zero amplitudes at sites of the $B$ sublattices. In regards to the penetration depth, $\lambda$, it is easily seen from Eqs. (12) and (13) that both solutions have the same value: $\lambda = -1/\ln |D_k|$. Nevertheless, the solution given by Eq. (13) has a linear dependence in $n$ which enhances its penetration into the bulk. We expect these states to contribute more to self doping than the usual single layer edge states [8], as the induced Hartree potential which limits the charge transfer between the bulk and the edge will be weaker. Note that the key to self doping is the presence of both an electron-hole asymmetry and extended defects. Electron-hole asymmetry may be due to inplane next nearest-neighbor hopping (NNN) $t'$, while edges play the role of extended defects. The finite $t'$ shifts the energy of edge states, leading to charge transfer between clean regions and defects. The energy shift for the single layer is given by $E_k \approx -t'(D_2^2 - 1)$ to first order in $t'$, apart from a global factor of $-3t'$ [12]. This is exactly the energy shift we get (away from the Dirac points) for bilayer graphene with inplane NNN hopping, if we neglect terms of the order $t' t_\perp/t$ and higher.

**FIG. 3:** (Color online) (a) - Charge density for bilayer edge states at $k_a/2\pi = 0.36$. (b) - The same as in (a) at $k_a/2\pi = 0.34$.

**FIG. 4:** (Color online) (a) - Energy spectrum for a graphene bilayer ribbon with zigzag edges for $N = 400$. (b) - Zoom of panel (a). The inter-layer coupling was set to $t_\perp/t = 0.2$.

Eqs. (12) and (13) [and the other two resulting from Eq. (9)] are eigenstates of the semi-infinite system only. In the ribbon the overlapping of four edge states leads to a slight dispersion and non-degeneracy. However, as long as the ribbon width is sufficiently large, this effect is only important at $ka \approx 2\pi/3$ and $ka \approx 4\pi/3$ where $\lambda$ is large enough for the overlapping to be appreciable [7]. As Eq. (13) has a deeper penetration into the bulk, its degeneracy is lifted first, as can be seen in Fig. 4(b). We may then conclude that edge states do exist in bilayer graphene ribbons. We expect band gaps to open due to magnetic instabilities induced by electron-electron interactions, similarly to graphene single layer [5, 9]. Actually, the edge states we have found live only on a single sublattice: $A$ or $B$ depending on the edge they are localized. This kind of localized states favor a ferromagnetic arrangement along the edge and antiferromagnetic between edges [13], consistent with what is found by first principles for stacked graphitic strips [14]. Also half-metallicity should occur in graphene bilayer nanoribbons as a consequence of edge states, analogously to the single layer [13].

From the point of view of scanning tunneling microscopy (STM) we notice that bilayer edge states give rise to different intensities depending on the ribbon edge. As an example we consider the ribbon shown in Fig. 2 and assume that the STM signal is essentially proportional to the local density of states of the upper layer. At the top zigzag edge the STM signal is due to edge states of the bilayer type, the only ones with finite amplitude on the upper layer [Eq. (13)]. On the other hand, at the bottom zigzag edge both bilayer and monolayer families have finite amplitude on the upper layer, and a higher STM intensity is expected therefrom.

**Nanoribbons of bilayer graphene (biased):** So far we studied localized states at the semi-infinite bilayer graphene. Experimentally, however, the relevant situation is a bilayer ribbon. The band structure of a bilayer ribbon with zigzag edges is shown in Fig. 4(a) for $N = 400$, obtained by numerically solving Eq. (4). We can see the partly flat bands at $E = 0$ for $k$ in the range $2\pi/3 \leq ka \leq 4\pi/3$, corresponding to four edge states, two per edge. The zoom shown in Fig. 4(b) for $ka \approx 2\pi/3$ clearly shows that there are four flat bands. Strictly speaking, the edge states given by

$$
\sum_{m,n}(n_1+m,n-n_2+m,n), \quad \text{with} \quad n_1+m,n = a_{i,m,n} a^{\dagger}_{i,m,n} + b^{\dagger}_{i,m,n} b_{i,m,n}. 
$$

Edge states are strongly affected by the bias. The semi-infinite biased system has only one edge state given by Eq. (12), as the edge state having finite amplitudes at both
layers [Eq. (13)] is no longer an eigenstate. In Fig. 5 we show the band structure of a bilayer ribbon for different values of the bias. Two partially flat bands for \( k \) in the range \( 2\pi/3 \leq ka \leq 4\pi/3 \) are clearly seen at \( E = \pm V/2 \). These are bands of edge states localized at opposite ribbon sides, with finite amplitudes on a single layer [Eq. (12) and its counterpart for the other edge]. Also evident is the presence of two dispersive bands crossing the gap. Both the closeness of these dispersive bands to \( E \approx \pm V/2 \) for \( ka \approx \pi \) and their crossing at \( E = 0 \) near the Dirac points can be understood using perturbation theory in \( V/t \). As surface states living at opposite edges have an exponentially small overlapping, and those belonging to the same edge are orthogonal, we can treat the solution given by Eq. (13) and its counterpart for the other edge separately. Starting with Eq. (13), the first order energy shift induced by the applied bias is \( E_k = V/2(\langle n_k^1 \rangle - \langle n_k^2 \rangle) \), where \( \langle n_k^1 \rangle \) and \( \langle n_k^2 \rangle \) give the probability of finding the localized electron in layer 1 and 2, respectively. The value of these quantities is easily obtained from Eq. (13) through a real space summation,

\[
\langle n_k^1 \rangle = \frac{(1 - D_k^1)^2}{(1 - D_k^2)^2 + t_{\perp}^2 / t^2}, \tag{16}
\]

\[
\langle n_k^2 \rangle = \frac{t_{\perp}^2 / t^2}{(1 - D_k^2)^2 + t_{\perp}^2 / t^2}. \tag{17}
\]

The band dispersion is thus given by,

\[
E_k^\pm = \frac{V}{2} \left( \frac{1 - D_k^2)^2 - t_{\perp}^2 / t^2}{2 (1 - D_k^2)^2 + t_{\perp}^2 / t^2} \right), \tag{18}
\]

where the minus sign stands for the band of states localized at the opposite edge. The result of Eq. (18) is shown in Fig. 5 as a dashed line which is hardly distinguishable from the numerical result. Note that for \( ka \approx \pi \) we have \( D_k \to 0 \), so that \( E_k^\pm \approx \pm V/2 \). This means that for \( ka \approx \pi \) the edge state given by Eq. (13) is essentially localized at layer 1, which is clearly seen from Eqs. (16) and (17) as long as \( t_{\perp} / t \ll 1 \). For \( 1 - D_k^2 = t_{\perp} / t \) the energy shift [Eq. (13)] is zero, which leads to band crossing. For \( t_{\perp} \ll t \) we can expand around the Dirac points, \( k_{\perp} a = 2\pi/3, 4\pi/3 \). If \( k = k_0 + \delta k \), the crossing takes place for \( \delta k a = \pm t_{\perp} / (t\sqrt{3}) \), each sign being assigned to different Dirac points. Note that \( \delta k \) is \( V \) independent, and its value compares fairly well with the numerical results shown in Fig. 5. Indeed, the approximate result given by Eq. (18) only fails at the Dirac points, where the edge states localization length diverges and their overlap has to be considered. Increasing the bias makes first order perturbation theory to break down. We have found numerically that the gap opens for \( V \gtrsim t \).

**Conclusions:** We have shown that zero energy edge states do exist at zigzag edges of bilayer graphene. We have derived an analytic solution for the wavefunction assuming a semi-infinite system and a first nearest-neighbor tight-biding model. This analytic solution defines two types of edge states: monolayer edge states, with finite amplitude over a single plane; and bilayer edge states, with finite amplitude over the two planes, and with an enhanced penetration into the bulk. Edge states are present even in bilayer graphene nanoribbons, where edge magnetization as well as half-metallicity are expected to show up in analogy with single layer graphene. We have also shown the robustness of bilayer graphene edge states to the presence of an electrostatic potential difference between planes.

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