Marginal topological properties of graphene: a comparison with topological insulators

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Abstract
The electronic structures of graphene systems and topological insulators have closely related features, such as a quantized Berry phase and zero-energy edge states. The reason for these similarities is that in both systems there are two relevant orbital bands, which generate the pseudo-spin degree of freedom and, less obviously, there is a correspondence between the valley degree of freedom in graphene and electron spin in topological insulators. Despite the similarities, there are also several important differences, both for the bulk topological properties and for their implications for the edge states—primarily due to the fundamental difference between valley and spin. In view of their peculiar band structure features, gapped graphene systems should be properly characterized as marginal topological insulators, distinct from either the trivial insulators or the true topological insulators.

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(Some figures may appear in colour only in the online journal)

1. Introduction
Condensed matter physics is witnessing increasingly rapid advances marked by a sequence of surprising discoveries. Two good recent examples are the realization of graphene [1]—a plane of carbon atoms forming a honeycomb lattice—and topological insulators [2, 3]—materials that are insulating in the bulk but conducting at the surface owing to topological reasons. In fact, the two examples were linked since the beginning: quickly after the initial experiments on graphene [4, 5], it was suggested by Kane and Mele that graphene may be gapped due to the intrinsic spin–orbit interaction and provide a prototype of a novel class of time-reversal-invariant topological insulators [6–8].

This idea has since inspired numerous works that have surprisingly, yet significantly, complemented our knowledge of some seemingly well-developed fields of solid state physics. The original prediction by Kane and Mele, however, remains unfulfilled in graphene because of the very weak spin–orbit interaction in realistic graphene samples.

In fact, in graphene, spin–orbit interaction is so weak that in most existing experiments without a magnetic field, the spin degree of freedom hardly plays a discernible role except for contributing an additional degeneracy. This certainly prevents graphene from being an ideal representative of (time-reversal-invariant) topological insulators for which strong spin–orbit interaction is normally a crucial ingredient. Despite this, graphene and topological insulators share a number of closely related features such as a quantized Berry phase [4, 5, 9] and zero-energy edge states [10–13]. Both in graphene and topological insulators, one can isolate two relevant bands with the opposite orbital symmetry, which in the band structure are connected by the matrix elements linear in momentum, leading to the Dirac cones. Further, in both systems, a gap in the Dirac dispersion can be opened, by means of spin–orbit interaction in topological insulators, and simply by creating a sublattice on-site energy imbalance in graphene. The analogy between the two systems is completed by associating electron spin in topological insulators with the valley degree of freedom (also time-reversal odd) in
2. Analogy between graphene systems and topological insulators

In this section, we demonstrate the analogy between graphene systems and 2D topological insulators by comparing two examples: gapped BLG and the BHZ topological insulator. In both examples, we consider for the moment the ideal cases—meaning that, for gapped BLG, we ignore coupling between valleys, and for the BHZ topological insulator we assume spin is a good quantum number.

2.1. Effective Hamiltonians

We start by writing down the low-energy effective Hamiltonian for each system. For gapped BLG, this reads

\[ H_{\text{BLG}}(k) = \begin{pmatrix} H_K(k) & 0 \\ 0 & H_{K'}(k) \end{pmatrix}, \]  

(1)

\[ H_K(k) = -\frac{\Delta}{(k_x + ik_y)^2} \begin{pmatrix} (k_x - ik_y)^2 \\ -\Delta \end{pmatrix}, \]  

(2)

\[ H_{K'}(k) = H_{K'}^*(-k), \]  

(3)

where \( K \) and \( K' \) denote the two valleys, \( k \) is the wave vector relative to each valley, and \( \Delta \) is the electrostatically tunable gap; for the BHZ topological insulator, it reads (with immaterial simplifications of the original model)

\[ H_{\text{BHZ}}(k) = \begin{pmatrix} H_S(k) & 0 \\ 0 & H_{S'}(k) \end{pmatrix}, \]  

(4)

\[ H_S(k) = \begin{pmatrix} m - k^2 & a(k_x - ik_y) \\ a(k_x + ik_y) & -(m - k^2) \end{pmatrix}, \]  

(5)

\[ H_{S'}(k) = H_{S'}^*(-k), \]  

(6)

where \( S \) and \( S' \) denote the two spin sectors, \( m > 0 \) and \( a \not= 0 \) are two real parameters. Note that, for simplicity, both Hamiltonians above have been made dimensionless by choosing fixed length and energy scales in each model.

One immediate observation can be made regarding the similar roles played by valley and spin. In the BHZ topological insulator, time-reversal symmetry is respected and imposes a constraint on the two spin sectors given by equation (6). The same constraint, given by equation (3), governs the two valley sectors of gapped BLG also as a consequence of time-reversal invariance (as mentioned above, due to lack of any significant spin–orbit interaction spin degree of freedom is decoupled from the orbital ones; therefore one can treat electrons as spinless particles in all calculations, reinstating the spin at the end). These constraints, generally obeyed in the two systems, have crucial implications for the overall bulk and edge properties as we will see in due course.

The two effective Hamiltonians, when characterized using the usual topological measures, seem to show an even closer analogy. For this purpose we use the common expression for the topological invariant associated with a fully gapped two-component Hamiltonian \( H(k) = g(k) \cdot \sigma \). Here \( \sigma \) is the vector of Pauli matrices and \( g(k) \) is a real vector. In terms of the normalized vector \( \tilde{g} = g/|g| \), the expression for the topological invariant is [26]

\[ c = \frac{1}{4\pi} \int d^2k \ \tilde{g} \cdot \left( \partial_k \tilde{g} \times \partial_{k'} \tilde{g} \right). \]  

(7)

The number \( c \) characterizes the mapping from the 2D \( k \)-space to the 2D parametric sphere subtended by \( \tilde{g} \) (see figure 1; for details see section 3). It is straightforward to find for individual valley/spin sectors (assuming \( \Delta < 0 \); \( c_{K/K'} \) changes sign when \( \Delta \) changes sign)

\[ c_K = -c_{K'} = 1, \]  

(8)

\[ c_S = -c_S = 1. \]  

(9)

Namely, the signs of \( cs \) reverse with respect to valleys/spins, echoing the underlying symmetry; the magnitudes of \( cs \) appear to be the same integer, 1, in both gapped BLG and the BHZ topological insulator!

2.2. Band inversion

The nontrivial bulk properties characterized by equations (8) and (9) are closely related to the occurrence of band inversion.

4 Note that the spinless time-reversal symmetry is not respected by the Haldane model [6], or one spin sector of the Kane–Mele model (spin being a good quantum number) [7].
Here we present a simple account of the band inversion in each system, bearing in mind that such accounts are by no means descriptions of the microscopic mechanisms, but rather phenomenological illustrations based on the effective models.

We start with the BHZ topological insulator. Due to the symmetry, it is sufficient to look at one spin sector, say $H_S$. Let us first compare two cases, with $m < 0$ and $m > 0$, respectively, and $a = 0$. It is obvious that the eigenstates in both cases are completely polarized in terms of the two-component pseudo-spin. Meanwhile, the crucial difference is that in the latter case ($m > 0$), the two parabolic bands intersect each other, while in the former case ($m < 0$) they are fully separated. In a purely phenomenological sense, we call such a reversal of the order of bands at low energy ‘band inversion’. But the real excitement about band inversion comes only after we turn on $a$, which leads to an avoided crossing between the two bands resulting in the opening of a gap (which makes the bulk of the system insulating). Figure 2 shows how the profile of the inverted bands changes with increasing magnitude of $a$. Clearly enough, at small $k$ (where only $m$ plays an important role), the electronic eigenstates tend to preserve their inverted character—states in the upper/lower band are largely polarized (in terms of pseudo-spin, see also figure 1) the same way as the original downward/upward parabolic band; at large $k$ (where both $m$ and $a$ become irrelevant), the order restores and the eigenstates in the upper/lower band deviate little from the original upward/downward parabolic bands; in between, as long as $a \neq 0$, the eigenstates continuously evolve such that the pseudo-spin polarization is gradually reversed. Hence, the band inversion has a direct consequence on the nontrivial evolution of the pseudo-spin polarization that is characterized by the number $c$. More specifically, the occurrence of band inversion leads to the occurrence of a topological phase transition with $c_{S/P}$ changing from 0 to ±1.

In gapped BLG, a similar occurrence of band inversion can be best seen at the level of the single-valley four-component Hamiltonian $H_{K}^{(4\times 4)}$—corresponding to two layers each with two sublattices—from which the two-component one in equation (2) is derived [27]. Again, the symmetry makes it sufficient to consider one of the two valleys. Explicitly, we have

$$H_{K}^{(4\times 4)}(k) = \begin{pmatrix}
V & hv_{F}k_{-} & t_{\perp} & -V \\
-hv_{F}k_{+} & V & t_{\perp} & -V \\
t_{\perp} & -V & -V & -V \\
-hv_{F}k_{+} & -V & -V & -V
\end{pmatrix}, \quad (10)$$

where $\pm V$ are the electrostatic potential on two graphene layers, $t_{\perp}$ is the interlayer nearest-neighbor hopping energy (assuming Bernal stacking of BLG), $k_{\pm} = k_{x} \pm ik_{y}$, $v_{F} = 3ta_{0}/2\hbar$ with $a_{0}$ and $t$ being the intralayer nearest-neighbor distance and hopping energy. Note that, unlike equation (2), this Hamiltonian is not dimensionless, and the parameters here are related to those in equation (2) by defining $\Delta = V/t_{\perp}$ and redefining $\ell_{0}k \Rightarrow k$ with $\ell_{0} = hv_{F}/t_{\perp}$.

We imagine first that the coupling between the two graphene layers is turned off ($t_{\perp} = 0$) such that, due to different electrostatic potentials, the electronic band structure...
at one single valley consists of two shifted Dirac cones as illustrated in the left panel of figure 3. For clarity we mark with two different colors the initial conduction bands (blue) and valence bands (red) in figure 3. Then we gradually turn on $t_\perp$. Analogous to the turning on of $a$ in the BHZ model, a finite $t_\perp$ opens a gap in the bulk and turns the system into an insulator (figure 3). As a result, at low $k$, a small part of the valence band of one layer is now ‘glued’ with the conduction band of the other layer to compose the new low-energy conduction band. The opposite happens to the new low-energy valence band (see text for details).

To summarize, the band inversion in gapped BLG, as illustrated above, happens as long as $V \neq 0$, and results in a finite $c_{K/S}$ that changes sign $\pm 1$ when $V$ changes sign. This is similar to the topological phase transition exhibited by the BHZ model, but not identical, since in the BHZ model $c_{K/S}$ changes from 0 to 1 as parameters are varied, and not from $-1$ to $+1$.

3. Limitations of the analogy

One fundamental limitation of the analogy between graphene and topological insulators, with valley playing the role of spin, is that valleys, which correspond to only parts of the Brillouin zone, are only defined in an energy window much smaller than the electron bandwidth (fraction of eV). That is in contrast to electron spin, which is well defined throughout the Brillouin zone. In most cases, due to the symmetry, the two valleys contribute oppositely to the topological property of the entire Brillouin zone—equation (8) for gapped BLG is an example. This results in overall topologically trivial graphene systems (according to the definition of a topological insulator; in the present context, this means that the integral in equation (7) needs to be calculated over the entire Brillouin zone and not only around one valley). This is certainly important since gapless edge states, if they exist at all, are not protected against intervalley scattering. More importantly, even in the complete absence of intervalley scattering (in the bulk and at the edges) crucial differences exist between the two systems at the level of single-valley models. These differences are the subject of this section.

3.1. Characterization of bulk properties

In the previous section we have employed equation (7) for the two models that we are considering, suggesting that the numbers $c$ characterize the topological properties associated with bulk bands. We now go back to examine the validity of such characterizations, based on a geometric interpretation of equation (7).

To begin with, we parameterize the general two-band Hamiltonian by setting $\hat{g} = (\sin \theta \cos \varphi, \sin \theta \sin \varphi, \cos \theta)$, where $\theta$ and $\varphi$ are the angles defining the direction of $k$. It is straightforward to show that the integrand of equation (7) is equal to $\sin \theta (\partial_\theta \varphi - \partial_\varphi \theta)$. Recognizing that the factor in parentheses is but the Jacobian $|\partial(\theta, \varphi)/\partial(k_\perp, k_\parallel)|$, equation (7) can be rewritten as

$$c = \frac{1}{4\pi} \int \int \sin \theta \, d\theta \, d\varphi. \quad (11)$$

It is immediately clear that geometrically $c$ measures the solid angle (divided by $4\pi$) covered by $\hat{g}$ when $k$ runs over the domain of interest.

Keeping this in mind, we can easily understand, by referring back to figure 1, that $c = \pm 1$ for the BHZ topological insulator because $\hat{g}$ covers the full sphere once, while $c = \pm 1$ for the gapped BLG because $\hat{g}$ covers a hemisphere (the upper or lower hemisphere, depending on the sign of $\Delta$) exactly twice. This fact reflects the crucial difference in how the vectors $\hat{g}$ behave at $k \rightarrow \infty$ in the two models. In the BHZ topoloyical insulator case, $\hat{g}(k \rightarrow \infty)$ converges identically to the south pole regardless of the direction along which infinity is approached, which allows for the compactification of the infinite $k$-plane as a Riemann sphere. In this case, a topologically nontrivial mapping from the compactified $k$-plane to the sphere subtended by $\hat{g}$ is well defined and $c$ is the topological invariant associated with this mapping. In the gapped BLG case, however, we have a different behavior: $\hat{g}(k \rightarrow \infty)$ sits on the equator and varies with the polar angle of $k$, preventing a proper compactification of the $k$-plane. Consequently, the numbers $c$ obtained, albeit nonzero, cannot be identified as topological invariants\(^5\) as in the previous case.

Despite this important difference, an alternative interpretation of equation (7) in terms of Berry phase and Hall conductivity remains valid, as we proceed to discuss. This indicates that the number $c$ has a well-defined physical (albeit not topological) meaning associated with the properties of the bulk. The Berry phase acquired by an electron transported adiabatically along a closed trajectory $C$ in the momentum space is given by

$$\gamma = \int_C \hat{g}(k) \cdot d\hat{C}, \quad (12)$$

\(^5\) The mapping $\hat{g}(k)$ based on a noncompact $k$-space is always topologically trivial, and the associated topological invariants are by definition zero.
where $u(k)$ is the periodic part of the Bloch wave function and can be obtained by taking the eigenstate of the Bloch Hamiltonian $H(k)$. Using Stokes’ theorem it is easy to show that the above formula is equivalent to
\[
\gamma = \frac{1}{2} \int_{\mathcal{C}} \partial^2 k \cdot 4 \Im (\langle \partial_k u(k) | \partial_k u(k) \rangle),
\]
with $\mathcal{C}$ the area enclosed by $\mathcal{C}$. For $H(k) = g(k) \cdot \sigma$ with $\hat{g}(k) = (\sin \theta \cos \varphi, \sin \theta \sin \varphi, \cos \theta)$ (the same parameterization as that used above; we consider a fully gapped system where $|g(k)| \neq 0$ for all $k$), the lower band 

eigenstate
\text{can be written as } u(k) = (\sin \frac{\pi}{2} e^{-i \varphi}, -\cos \frac{\pi}{2})^T. \text{ Then it is straightforward to show that the integrand in equation (13) is again equal to } \sin \theta |\partial(\theta, \varphi)/\partial(k_1, k_z)|; \text{ hence}
\[
\gamma = 2 \pi c. \quad (14)
\]
It follows from equation (13) that $\gamma$ can also be identified as the Hall conductivity (in units of $e^2/2\pi h$) contributed by all the occupied Bloch states [28]. Therefore, $c_{K_K'}$ in equation (8), despite not representing a true topological invariant, still corresponds to the valley-specific Hall conductivity [29, 30, 31] (in units of $e^2/h$) when the Fermi energy lies in the bulk band gap (i.e. $c$ is the contribution of the occupied states in a given valley to the bulk Hall conductivity).

Similarly, of course, $c_{S/S'}$ in the BHZ model represent the spin-specific Hall conductivity, next to being well-defined topological invariants.

### 3.2. Bulk–edge correspondence

For the BHZ topological insulators $c_{S/S'}$ are topological invariants. According to the principle of bulk–edge correspondence, this implies the presence of one pair of spin-helical gapless edge modes at each edge [18, 32]—their helicity follows from the opposite signs of $c_S$ and $c_S$. In contrast, a similar conclusion cannot be drawn for the valley-specific $c_{K/K'}$, since these quantities are not topological invariants. In general, therefore, we cannot expect bulk–edge correspondence in this case. In the particular example of gapped BLG, this leads, as we will show, to the possibility that the number of gapless edge modes is dependent on boundary conditions.

To analyze the presence of edge states in gapped BLG we solve the wave equation for different edge structures (that do not couple states in different valleys), with which corresponding appropriate boundary conditions are associated. Possible boundary conditions for the single-valley models can be derived from the constraint of vanishing probability current across the boundary (we will comment on their physical realizations later). Again we use one single-valley Hamiltonian, described by equation (2), as our example. Let us write the two-component wavefunction as $\Psi(r) = (\psi_1(r), \psi_2(r))^T$. From the single-valley continuity equation $\frac{\partial}{\partial r} \Psi^T + \nabla \cdot j = 0$, we obtain the probability current density
\[
j(r) = [i(\psi_1^* \partial_- \psi_2 + \psi_2^* \partial_+ \psi_1) + \text{c.c.}] \hat{x} + [(\psi_1^* \partial_- \psi_2 - \psi_2^* \partial_+ \psi_1) + \text{c.c.}] \hat{y}, \quad (15)
\]
where $\partial_\pm = \partial_x \pm i \partial_y$ and c.c. stands for complex conjugation.

Without loss of generality, let us suppose we have a semi-infinite sample occupying the upper half-plane with its edge along $y = 0$. The requirement that no current should flow through the boundary implies
\[
j_y(y = 0) = 0 \quad \text{for } y > 0. \quad (16)
\]
This equation can be satisfied in various settings, each accounting for one possible boundary condition. For example, we can easily see that any of the following conditions satisfy the above equation:
\[
\begin{align*}
\psi_1 |_{y=0} &= (\partial_y \psi_1) |_{y=0} = 0, \\
\psi_2 |_{y=0} &= (\partial_y \psi_2) |_{y=0} = 0, \\
\psi_1 |_{y=0} &= \psi_2 |_{y=0} = 0, \\
(\partial_y \psi_1) |_{y=0} &= (\partial_y \psi_2) |_{y=0} = 0.
\end{align*}
\]
In principle, the boundary conditions obtained this way should be verified physically by finding corresponding terminations of the BLG lattice, which is in general a formidable job. As a matter of fact, the examples given in equations (17)–(20) can be derived alternatively from the tight-binding models by considering specific sublattices that are missing at a sample edge. This latter approach follows in spirit Brey and Fertig [33] who dealt with single-layer graphene nanoribbons, and has been discussed in detail for the case of BLG in [24]. At this point, without entering into details, we simply list the physical realizations of the boundary conditions (17)–(20) in the same order in the left panels of figure 4. Note in particular that all the edges shown are parallel to a zigzag configuration, which leaves the valleys uncoupled and hence allows the application of the single-valley models.

Also illustrated in figure 4 are the bulk band profiles (blue areas) and the dispersion relations of edge states (red and green curves for two opposite edges) corresponding to the specific type of edges shown in the same row of the figure. These spectra of electronic states can be calculated either by solving the single-valley models with boundary conditions discussed above or by exact diagonalization of tight-binding models including edges—the results of the two approaches agree quantitatively [24]. One can immediately see that, as the most important fact, the number of subgap edge modes per valley per edge explicitly depends on the edge configurations, and can be 1, 2 or 0. Therefore, a naive extension of bulk–edge correspondence to the individual valleys, which would predict the above number to be 1 independent of the specific boundary, does not hold for gapped BLG.

We close this section with two remarks. Firstly, our focus in this paper centers on the subgap edge states, which are dispersive and may contribute significantly to the subgap conductance. Nevertheless, there can be another set of edge states (also depending on boundary conditions) that are dispersionless and form flat bands [10] (shown as gray lines in figure 4). Indeed, the presence of both types of edge states as a whole has a well-defined topological origin from the bulk [34], and can be formulated in
4. Marginal topological character of graphene systems

Although the analogy between valley-based graphene systems and spin-based topological insulators has significant limitations, the bulk topological properties of graphene systems, as we show below, nevertheless play an important role, with experimental consequences. To emphasize the connection of graphene and topological insulators, and at the same time to stress that there are important limitations to this connection, we refer to graphene as a ‘marginal topological insulator’. We borrow the term ‘marginal’ from Volkov to characterize the occurrence of noncompact momentum space in the case of massive Dirac fermions [26]. In the following section, we illustrate that such a marginal character has direct experimental implications.

4.1. Domain wall

The noncompact momentum space for single-valley models can sometimes be ‘cured’ to obtain well-defined nontrivial topological properties. One such case is a smooth topological insulator. We borrow the term ‘marginal’ from Volkov to emphasize the experimental implications.

In the case of gapped BLG, a domain wall can be realized controllably by gating two adjacent areas of a sample with opposite polarities (i.e. opposite signs of $\Delta$) [11]. The resulting nontrivial topological character that can be ascribed to an individual valley can be understood as follows. We have discussed in the previous section that the $\hat{\mathbf{g}}$ vectors—corresponding to the pseudo-spin polarization of Bloch states—cover a hemisphere twice when $k$ runs over the 2D $k$-plane in the case of a single valley of uniformly gapped BLG (see the left panel of figure 5). It is also clear that, for a specific valley, the hemisphere which $\hat{\mathbf{g}}$ covers is determined by the sign of $\Delta$. It follows that for a domain wall across which $\Delta$ changes sign, the bulk $\hat{\mathbf{g}}$ vectors of the two domains cover two opposite hemispheres (each twice) which seamlessly connect on the equator at $k \rightarrow \infty$ (where $\Delta$ in the Hamiltonians becomes irrelevant; see the right panel of figure 5). In this way the momentum space—within the same valley—is compactified as a 2D sphere and the topological invariant characterizing the mapping from such a sphere to the parametric $\hat{\mathbf{g}}$-sphere is evidently nontrivial. In fact, this topological invariant is given by $\delta c_\tau = c^+_\tau - c^-_\tau$ ($\tau = K$, $K'$), which is the difference of the number $c_\tau$’s for the two domains.

For a BLG domain wall, therefore, $\delta c_{K/K'} = \pm 2$, and the difference is a topological invariant. Moreover, it is this difference that defines the number of valley-polarized chiral modes via the spectral asymmetry theorem [26]. While the theorem readily applies to...
smooth—quasiclassical—interfaces such as domain walls, its conditions are in general violated at the sharp interface between graphene and vacuum, even when the interface does not mix valleys. The problem is that one cannot adiabatically interpolate between the spectrum of gapped BLG to the spectrum of vacuum, while one easily can in the case of an interface between the regions with $\Delta > 0$ and $\Delta < 0$. Indeed this is what has been analytically and numerically verified by Martin et al [11]. Moreover, the presence of these ‘interface’ gapless modes—unlike those discussed previously for plain BLG edges—does not depend on microscopic details such as the crystallographic configuration of the domain wall, which confirms the topological origin of these modes.

4.2. Gapped bilayer graphene with disordered edges

The marginality of single-valley models implies, on the one hand, that a small perturbation in the Hamiltonian or boundary conditions can have a substantial effect on the presence or absence of the gapless modes (see e.g., [13])—this is merely a restatement of the absence of bulk–edge correspondence for individual valleys. On the other hand, considering a realistic graphene device where the edges of the sample are inevitably under the influence of disorder, physical properties will necessarily be subject to ensemble averages, such that, at sufficiently strong disorder, universal behaviors are expected to occur. It is thus interesting to ask in this context whether, or how, the marginal bulk property can be manifested in the subgap regime.

Such a question has been addressed by Li et al [25] with the motivation to investigate the ‘extra’ contribution by the edge states to the subgap conduction in realistic experiments on gapped BLG. It is found that indeed strong edge disorder leads to universal presence of subgap edge states which are localized with fairly long localization length. To be specific, figure 6 shows how the localization length

$$\lambda_{loc}$$—extracted numerically with the aid of the scattering theory [39, 40]—depends on the strength of edge disorder ($d$ denotes edge roughness depth; solid lines represent the effect of structural disorder only and broken lines represent the effect of structural plus chemical disorder; see [25] for details), with various choices of initially ordered (e.g. zigzag and armchair) edges, and a fixed gap size at an experimentally accessible value. For the cases where the starting edges accommodate pre-existing edge states (counter-propagating for two valleys), gradually turning on disorder leads to localization and a shrinking in the transverse size of the edge states (see the solid lines in blue and green)—this is certainly expected because of the lack of protection for the edge states from back-scattering in the system. What is unexpected is that when starting from initial edges where no edge state exists even in the absence of disorder, disorder starts to introduce (localized) edge states inside the gap. Moreover, at relatively weak disorder, increasing disorder enhances their $\lambda_{loc}$ (see the solid line in red). Most interestingly, the localization length in all cases converges to the same value at sufficiently strong disorder (see the solid lines in the large-$d$ limit, and also the broken lines where dominant chemical disorder leads to very fast convergence even at small $d$). The universal value $\lambda^*$ depends only on the gap size ($\lambda^* \propto 1/\sqrt{\Delta}$; it is about 20 nm in the cases illustrated in figure 6 corresponding to a gap of approximately 100 meV). These values of the localization length can have experimental consequences in real devices [25].

Thus subgap edge states with a long localization length are generically present at disordered edges of gapped BLG (see figure 7). These states can manifest themselves in transport experiments at finite temperature. In particular, when the subgap transport is dominated by 1D variable range hopping along the edges (which is the case if the hopping through the bulk disorder is sufficiently suppressed), the temperature dependence of the conductance is of the form $G \propto \exp[-(T^*/T)^{1/2}]$, with $T^*$ being the characteristic energy scale that determines the low-temperature transport behavior. For practically relevant parameters, $T^*$ is proportional to, but approximately one order of magnitude smaller than, the size of the bulk gap [25]. Indeed in very clean gapped BLG devices, in which the bilayer is suspended and not in contact with a substrate, an excess conductance, thermally activated with a characteristic energy one order of magnitude smaller than the gap, has

![Figure 6.](image_url)

**Figure 6.** The localization length $\lambda_{loc}$ of low-energy edge states as a function of edge roughness depth $d$ (divided by $a_0$, the intralayer nearest-neighbor distance of graphene). The simulations to extract $\lambda_{loc}$ are carried out starting from different ordered edges, with or without chemical disorder. In all cases, sufficiently strong disorder leads to the same value of $\lambda_{loc}$ regardless of any detail. This universal localization length, $\lambda^*$, is about 20 nm for the current case shown here, corresponding to a gap of approximately 100 meV.

![Figure 7.](image_url)

**Figure 7.** Local density of states at zero energy (center of the gap) in a typical two-terminal device of gapped BLG with disordered edges. Edge states are universally present in the subgap regime with long localization length.
been observed experimentally [41]. These observations are in agreement with our predictions.

To verify the link between the presence of the weakly localized subgap edge states and the marginal bulk properties, we performed a comparative study between the current BLG system and a half-filled square lattice with different on-site energies on nearest neighbors. Such a square lattice is fully gapped but has a topologically trivial band structure [25]. It turns out that in the latter system no subgap edge states are present neither in the ordered nor in the disordered limit; subgap transport is solely due to weak direct tunneling between contacts, which is even further suppressed in the presence of edge disorder. The sharp contrast between the two systems strongly suggests that the universal features in terms of the (localized) subgap edge states in gapped BLG are indeed a manifestation of its nontrivial marginal topological bulk properties.

5. Conclusion

To conclude, we have carefully explored the apparent analogy between a gapped graphene system and topological insulators, with the central element of the correspondence being the valley degree of freedom in the former and electron spin in the latter. We find that the strong formal resemblance is fundamentally limited by the different topological structures in the electron spectrum underlying the two systems. We have used gapped BLG as a concrete example to facilitate the comparison and, more importantly, to demonstrate how the ‘marginal’ topological properties associated with valleys can manifest themselves significantly in experiments. Yet, such marginal characteristics are generic for graphene systems, and can be readily seen by considering other graphene multi- and single layers.

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