Splitting of critical energies in the $n = 0$ Landau level of graphene

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New Journal of Physics 11 (2009) 095019 (13pp)
Received 4 July 2009
Published 30 September 2009
Online at http://www.njp.org/
doi:10.1088/1367-2630/11/9/095019

Abstract. The lifting of the degeneracy of states from the graphene $n = 0$ Landau level (LL) is investigated through a non-interacting tight-binding model with random hoppings. A disorder-driven splitting of two bands and of two critical energies is observed by means of density of states and participation ratio calculations. The analysis of the probability densities of the states within the $n = 0$ LL provides some insights into the interplay of lattice and disorder effects on the splitting process. An uneven spatial distribution of the wavefunction amplitudes between the two graphene sublattices is found for the states in between the two split peaks. It is shown that as the splitting is increased (linear increasing with disorder and square root increasing with magnetic field), the two split levels also get increasingly broadened, in such a way that the proportion of overlapped states remains approximately constant for a wide range of disorder or magnetic field variation.
1. Introduction

Since the first observation of the anomalous integer quantum Hall effect in graphene [1, 2], there has been an enormous interest in the quantum Hall regime of graphene. The sequence of quantized Hall plateaux in graphene is shifted by half-integer if compared to the well-known quantum Hall effect in usual two-dimensional electron gases. This is understood to be due to the unique features of the Dirac-like band structure of graphene around the Fermi energy [3], which gives rise to the following Landau level (LL) quantization: $E_n = \pm v_F \sqrt{\frac{2\hbar B}{e}|n|}$, where $n$ is the LL index and $v_F$ is the Fermi velocity. In this way, the energy dependence for LLs with magnetic field $B$ is no longer linear, but goes with the square root of $B$ (positive energies for electrons and negative energies for holes), and there is also an LL at zero energy (the $n = 0$ LL), which is shared by electrons and holes. In addition to the spin degree of freedom, graphene LLs exhibit a valley (sublattice) degree of freedom, giving a fourfold degeneracy to each LL.

Of special interest in the recent literature is the question of the lifting of the $n = 0$ LL degeneracies, observed experimentally [4]–[8] and discussed in many theoretical works [9]–[22]. Among these works, there are various different proposed explanations for the origins of the observed splittings; however, a consensus is still missing. Two of the experimental works [4, 5] show a complete lifting of the $n = 0$ fourfold degeneracy occurring for high magnetic fields up to 45 T and for low temperatures, and there are evidences that the extra quantum Hall plateaux observed at filling factors $\nu = 0$ and $\nu = \pm 1$ should be due to the lifting of the spin and sublattice degeneracy, respectively [5]. The other three experiments [6]–[8] show the lifting of only one of the degeneracies. Giesbers et al [8] suggest that the opening of the gap in $n = 0$ observed by them might be spin related (Zeeman splitting), because of a linear variation of the gap with $B$; however, in [7] the authors believe that the gap they measure should be driven by a broken sublattice symmetry. The physical mechanisms leading to sublattice symmetry breaking are still not completely understood.

In this work, a splitting of two critical energies in the $n = 0$ LL is observed within a nearest-neighbor tight-binding model for the graphene lattice, considering random hoppings as the only source of disorder. The splitting is inferred by means of participation ratio (PR) calculations, which show clearly the presence of two split peaks indicating two critical energies inside the $n = 0$ Landau band, with localized states in between the peaks, around $E = 0$. The splitting is found to have a linear dependence with disorder strength and a square root dependence with...
magnetic field, in agreement with the numerical calculations from [17] and with the magnetic field dependence observed for the gap in [5]. As the model used in this work considers a single-particle picture (no electron–electron interaction is taken into account) and there are no spin-dependent terms in the Hamiltonian, the splitting observed is related only to the effects of random hopping disorder, which promotes intervalley mixing. To help in the understanding of these effects, I look here to the wavefunctions (probability densities) of the states within the $n = 0$ LL, inspecting in particular the distribution of amplitudes in each of the sublattices, which gives interesting information about the process of disorder-driven sublattice mixing.

A unit cell of the graphene hexagonal lattice contains two carbon atoms, defining two sublattices, $A$ and $B$. For the perfect graphene lattice (in the absence of any kind of disorder), the wavefunctions of all the states within the $n = 0$ LL have nonzero amplitudes in only one of the sublattices. ‘How does disorder affect this picture?’ was the question addressed in [23], where it is shown how the increasing inclusion of a diagonal disorder (both white-noise or a Gaussian-correlated disorder model) increasingly promotes the spread of wavefunctions to both sublattices in the $n = 0$ LL, an effect that was revealed to be directly connected to an anomalous localization: an enhancement of the PR of the states with increasing disorder [23, 24]. However, for diagonal disorder models (on-site energy fluctuations) the splitting of critical energies does not occur. Here, the same question is addressed for an off-diagonal disorder model, for which a completely different picture emerges. The probability densities of the states from the $n = 0$ LL are not anymore concentrated predominantly in one of the sublattices, even when small disorders are considered. However, the interesting find is that for the states closest to $E = 0$ (which get localized for the present disorder model, contributing to the critical energies splitting process), the spatial distribution of the wavefunctions in one sublattice is completely different and independent of the distribution in the other sublattice. These are states corresponding to the region where the two split bands have an important overlapping.

2. Model: tight-binding Hamiltonian with random hopping

The tight-binding Hamiltonian model for the honeycomb lattice used in this work is defined by

$$H = \sum_{\langle i,j \rangle} (t_{ij} e^{i\phi_{ij}} c_i^\dagger c_j + t_{ij} e^{-i\phi_{ij}} c_j^\dagger c_i),$$

where $c_i$ is the fermionic operator on site $i$ and the sum runs over nearest-neighbor bonds. The hopping parameters $t_{ij}$ randomly fluctuate (white noise) from bond to bond around the average value $t$:

$$t - \frac{W}{2} \leq t_{ij} \leq t + \frac{W}{2}.$$  

Thus, $W$ represents the width of the uniform off-diagonal disorder distribution, $t \approx 2.7$ eV for graphene; however, note that the results shown here are parameterized by $t$. The graphene lattices considered have $M \times N$ atoms ($M$ zigzag chains, each containing $N$ atoms), which are repeated by means of periodic boundary conditions. In this way, the dimensions of the lattices that are periodically repeated are given by

$$L_x = (M - 1)a\frac{\sqrt{3}}{2}, \quad L_y = (N - 1)a\frac{a}{2},$$

where $a = 2.46$ Å is the lattice constant for graphene.
Figure 1. (a) DOS showing five LLs (from \( n = 0 \) to 4), broadened by a random hopping disorder with \( W/t = 0.1 \), averaged over 360 disorder realizations. The size of the lattice considered is 19.8 nm \( \times \) 11.4 nm \( (M \times N = 94 \times 94) \), with magnetic flux \( \Phi/\Phi_0 = 2/94 \approx 0.021 \). (b) Zoom showing details of the shape of the DOS for the \( n = 0 \) LL, including the small peak at \( E = 0 \). (c) Corresponding PR of the states from the \( n = 0 \) LL, calculated for the same parameters, also averaged over 360 disorder realizations. A clear splitting in two critical energies can be observed.

The perpendicular magnetic field \( B \) is included by means of a Peierls substitution (a complex phase factor in the hopping parameter): \( \phi_{ij} = 2\pi(e/h) \int_j^i A \cdot dl \). In the Landau gauge, \( \phi_{ij} = 0 \) along the \( x \)-direction and \( \phi_{ij} = \pm \pi(x/a)\Phi/\Phi_0 \) along the \( \mp y \)-direction, with \( \Phi/\Phi_0 = B a^2 \sqrt{3} e/(2h) \). The focus is on the low-flux limit (\( \Phi/\Phi_0 < 0.05 \)), where the graphene LLs are well defined in this model.

3. Density of states (DOS) and the \( n = 0 \) LL splitting

In figure 1(a), one can see the DOS with the formation of five LLs, \( n = 0 \) to 4, corresponding to a magnetic flux \( \Phi/\Phi_0 = 2/94 \approx 0.021 \). The broadening of the levels is due to a random hopping disorder strength of \( W/t = 0.1 \). The lattice considered has \( M \times N = 94 \times 94 \), which means dimensions of \( L_x = 19.8 \) nm along the armchair boundary and \( L_y = 11.4 \) nm along the zigzag boundary, defining the disordered unit cell that is periodically repeated (periodic boundary conditions). The spectrum is symmetric about zero energy; thus the \( n < 0 \) LLs are not shown here. The DOS is an energy histogram accumulated after the Hamiltonian diagonalization for hundreds of disorder realizations—in the case of figure 1, 360 realizations were taken.
Here, we see that the $n = 0$ LL has about the same broadening width of the higher levels, which is different from the result obtained in [17] for a random magnetic flux disorder. However, the dependence of the $n = 0$ LL broadening with the correlation length of the random hopping was already elucidated in [20], which shows a DOS with broadenings in accordance with the results shown here for spatially uncorrelated random bonds.

Figure 1(b) shows a zoom of the DOS for the region of the $n = 0$ LL, allowing the observation of the shape of this central band in more detail, which has many differences when compared to the $n = 0$ Landau band shape obtained for a diagonal disorder model [23, 24] and even when compared to the higher levels. One can see in figure 1(b) that the broadening of this central LL does not resemble the usual Gaussian or semi-elliptic LL line shape, but instead has a lowering in the DOS around the center (apart from the presence of a small peak observed in the DOS for states closer to $E = 0$). It is important to note that these characteristics for the $n = 0$ LL are kept for many other values of disorder and magnetic fluxes investigated in this work.

The localization properties of the states within the $n = 0$ LL are inspected, as shown in figure 1(c), throughout the calculation of the PR [25], defined by

$$\text{PR} = \frac{1}{N'} \sum_{i=1}^{N'} |\psi_i|^4,$$

where $\psi_i$ is the amplitude of the normalized wavefunction on site $i$, and $N' = M \times N$ is the total number of lattice sites. The PR gives therefore the proportion of the lattice sites over which the wavefunction is spread. In this way, the PR for a localized state vanishes in the thermodynamic limit, while peaks in the PR indicate the presence of extended states (critical energies).

Figure 1(c) allows the observation of two peaks in the PR, clearly indicating the splitting of two critical energies. Note that this splitting occurs exclusively for the $n = 0$ LL in the considered model: the PR calculated for the higher levels (not shown here) shows always only one critical energy (one PR peak) per Landau band. It is also important to point out that the splitting of the two critical energies is well defined even considering that there is no clear opening of an energy gap in the DOS (see section 6 for further discussion on this). As observed in the comparison between figures 1(b) and (c), there are states contributing to the DOS in the energy range between the two PR peaks; however, the closer these states are to $E = 0$, the more localized they are, i.e. the smaller is their localization length.

Throughout this work, various PR calculations are shown, corresponding to different system sizes, different values of magnetic flux and different disorder strengths. In order to examine the question of possible finite lattice-size effects on the PR calculations and get a better understanding of the PR peaks, in figure 2 the PR is shown for three different lattice sizes. The dimension $L_x = 19.8$ nm is kept constant while $L_y$ varies from $L_y = 3.3$ nm, to 4.8 nm and to 11.4 nm (more details are specified in the figure caption). Disorder and magnetic flux values are the same for the three cases: $W/t = 0.1$ and $\Phi/\Phi_0 = 2/94 \approx 0.021$. The magnetic length for this flux is $l_B = \sqrt{\hbar/(eB)} = 6.3$ Å. Figure 2 shows how the peaks in the PR are better defined as the system size is increased. As expected, the PR values corresponding to the localized states (at the LL tails and LL center) decrease with increasing system size [25].

The periodical boundary conditions impose restrictions over the values of magnetic flux in order to guarantee a commensurability between the wavefunction periodicity (the phases in the hopping) and the periodicity of the lattice considered. For a system with $M \times N$ sites, and considering the Landau gauge, the possible sequence of flux values is $\Phi/\Phi_0 = 2/M, 4/M, 6/M, \ldots$. Consequently, to consider small values of magnetic flux it is necessary...
to increase \( M \) (i.e. increase the \( L_x \) dimension); however, computational limitations restrict the maximum matrix sizes to be diagonalized. To analyze small fluxes it is then usually necessary to consider rectangular lattices, which are bigger in \( L_x \) than in \( L_y \). However, we have to keep in mind that \( L_y \) should still be many times greater than \( l_B \) in order to avoid finite size effects and for the PR peak to be well defined, as figure 2 suggests. It is worth noting here that the increase of \( N \) in figure 2 also determines the increase of the number of states per LL. The degeneracy is given by \( d = M \times N \times \Phi/\Phi_0 \) and, due to the flux restrictions discussed above, for a flux \( \Phi/\Phi_0 = 2/M \), we have \( d = 2N \) states per LL.

4. Probability densities in each of the sublattices

In order to try to get more information about the states within the \( n = 0 \) LL and, consequently, about the origin of the critical energies splitting observed, the wavefunctions of these states are now examined. Special attention is paid to how the wavefunction amplitudes are distributed in each of the two sublattices; once in the absence of disorder all the eigenstates within the \( n = 0 \) LL have nonzero amplitudes only on one of the sublattices [9].

Figure 3, shows the PR calculated for a graphene lattice with \( M \times N = 60 \times 108 \) atoms \((L_x \times L_y = 12.6 \text{ nm} \times 13.2 \text{ nm})\), considering a magnetic flux \( \Phi/\Phi_0 = 2/60 \approx 0.033 \) and a random hopping amplitude \( W/t = 0.1 \) averaged over 500 disorder realizations. For three selected states indicated by arrows, the wavefunction probability densities, \(|\Psi|^2\), plotted over all the \( 60 \times 108 \) lattice sites are also shown, and then, below, the probability densities over each sublattice are plotted separately. After calculating the eigenvectors from the exact Hamiltonian diagonalization, we have the probability density \(|\Psi_{i,j}|^2\) for all the lattice sites \((i, j)\). Then, the
Figure 3. Top: PR of the \( n = 0 \) LL states from a graphene lattice with size \( L_x \times L_y = 12.6 \text{ nm} \times 13.2 \text{ nm} \) \( (M \times N = 60 \times 108 \text{ atoms}) \), \( \Phi / \Phi_0 = 2/60 \approx 0.033 \), \( W/t = 0.1 \) averaged over 500 disorder realizations. Bottom: probability densities \((|\Psi|^2)\) and their decompositions, showing separately the contributions from each sublattice \((|\Psi_A|^2\) and \(|\Psi_B|^2\)) for three selected states indicated by arrows. While for states 2 and 3, the probability density has the very same spatial distribution in each of the two sublattices, state 1 is peculiar because it has completely different and independent probability densities in each sublattice (all the states close to \( E = 0 \), from the region inside the circle, show this behavior).

The most interesting observation refers to state 1, for which the spatial distribution of the probability density on one sublattice is completely different from that on the other one.

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Observing $|\Psi_A|^2$ and $|\Psi_B|^2$ separately, after sublattice decomposition, one can see that for state 1 the distribution of maxima and minima amplitudes on one sublattice occurs in different positions and is completely independent of the distribution on the other. All the states in the region delimited by the dashed circle, states close to $E = 0$, have this absolutely uneven spatial distribution of amplitudes between the sublattices. Observe that this is the region where the two Landau bands (split after the symmetry breaking) have a more significant overlap. For states in the region right after the limits of the circle, some position correlation between the two sublattices starts to appear in the probability density (not shown here). But when reaching the proximities of the PR peak, and after that until the band tail, we observe that all the states show a symmetric distribution between the sublattices, as can be observed from the decompositions of states 2 and 3: for these states, the amplitudes on one sublattice look pretty much like the amplitudes on the other.

It is worth noting here that even when distributions are asymmetric between the sublattices, it is not the case that the wavefunction is more concentrated in one sublattice than in the other. The behavior observed here is different from that observed when disorder is introduced through on-site energy fluctuations (diagonal disorder models), for which the amplitudes over one sublattice are more significant than over the other [23, 24]. In fact, both sublattices have equally significant amplitudes (note that the vertical scales of the graphics are kept the same in the decomposition): the sum of all amplitudes over sublattice $A$ was calculated and is practically equal (with differences smaller than 5%) to the sum over sublattice $B$.

Observing the wavefunction amplitudes in figure 3, one sees that the localization length of state 1 is larger than that of state 3. Nevertheless, this difference reflects exactly the difference in the PR values of these states: for this system size, the PR for state 1 is not as small as for state 3 (remember, however, that the lattice size analysis shown in figure 2 indicates that both state 1 and state 3 are truly localized states). The reason for the larger localization length for state 1 compared with state 3 is most probably due to the mixing of states in this region between the PR peaks, where the split LLs are overlapped (see section 6).

Another point to be noted is that, for the disorder model considered (only off-diagonal, with random nearest-neighbor hoppings), there is perfect symmetry around $E = 0$: the probability density of the state at a specific energy $E$ is exactly equal to that of the state at $-E$. One can also observe this through the perfect symmetry of the PR curve around $E = 0$.

5. Splitting as a function of disorder and magnetic field

The evolution of critical energies splitting in the $n = 0$ LL with increasing disorder strength and magnetic field is investigated here; the results are shown in figures 4 and 5, respectively.

Figure 4(a) shows the PR of the states from the $n = 0$ LL for three different disorder strengths: $W/t = 0.2, 0.4$ and 0.6. One can see that the energy splitting $\Delta E/t$ between the two critical energies (between the two PR peaks) is increased as the disorder increases the LL broadening. In figure 4(b), the splitting $\Delta E/t$ is plotted as a function of disorder for several values of $W/t$. The linear regression fits well the data for the wide interval of disorder considered: the red line passes through the origin and has an angular coefficient of $0.1306 \pm 0.0006$, which is valid for the specific magnetic flux considered here ($\Phi/\Phi_0 = 2/66 \approx 0.030$):

$$\frac{\Delta E}{t} = (0.1306 \pm 0.0006) \frac{W}{t}. \quad (5)$$
Figure 4. (a) PR for the states from the $n = 0$ LL, for three different disorder strengths. The energy splitting between the two peaks is observed to increase with disorder. Lattice size is 13.9 nm $\times$ 9.2 nm ($M \times N = 66 \times 76$) and magnetic flux $\Phi/\Phi_0 = 2/66 \approx 0.030$. (b) Energy splitting of the two critical energies versus $W/t$, the random hopping disorder strength, showing linear fit.

The results shown in figure 4 are calculated for lattices having sizes 13.9 nm $\times$ 9.2 nm ($M \times N = 66 \times 76$) and for a magnetic flux $\Phi/\Phi_0 = 2/66 \approx 0.030$, averaging the results for each $W/t$ over 400 disorder realizations.

In figure 5, the dependence of the splitting with magnetic field is examined. Figure 5(a) shows four examples of PR calculations within the $n = 0$ LL for fixed disorder ($W/t = 0.1$) and different magnetic fluxes. An increasing energy splitting of the two PR peaks is observed with increasing magnetic flux. In figure 5(b), this splitting is plotted as a function of magnetic flux for several magnetic fluxes, and we observe that the data are well adjusted by a square root fit (the specific coefficient value is valid for the specific disorder $W/t = 0.1$ considered):

$$\frac{\Delta E}{t} = (0.0736 \pm 0.0005) \sqrt{\frac{\Phi}{\Phi_0}}.$$  \hfill (6)

The dimensions of the lattices considered for each flux, as well as the magnetic lengths and number of disorder realizations undertaken are listed in table 1.

Regarding the square root dependence of the splitting with magnetic field, observe that the critical energies from the $n = 0$ LL follow then the same dependence with $B$ presented by the higher LLs.

It is important to note that these results are in agreement with [17], where a linear dependence of the energy splitting with the disorder and a square root dependence with the magnetic field were observed for a random magnetic-field disorder model, and where the splitting was determined by means of the two-terminal conductance peaks.
Figure 5. (a) PR for the states from the $n = 0$ LL, for four different magnetic fluxes. The energy splitting between the two peaks is observed to increase with magnetic flux. (b) Energy splitting of the two critical energies versus magnetic flux $\Phi/\Phi_0$, showing a square root dependence. The lattice dimensions vary for each flux considered and are listed in table 1. The disorder strength is $W/t = 0.1$ for all of the fluxes.

Table 1. Lattice dimensions, magnetic length $l_B$ and number of disorder realizations, for each of the magnetic fluxes considered to define the data points in figure 5.

| $\Phi/\Phi_0$ | $M \times N$ | Lattice sites | $L_x (\text{Å}) \times L_y (\text{Å})$ | $l_B (\text{Å})$ | Realizations |
|---------------|---------------|---------------|---------------------------------|-----------------|--------------|
| 0.0114        | 176 $\times$ 50 | 8.800         | 372.7 $\times$ 60.3              | 8.57            | 350          |
| 0.0118        | 170 $\times$ 60 | 10.200        | 360.0 $\times$ 72.6              | 8.42            | 210          |
| 0.0133        | 150 $\times$ 62 | 9.300         | 317.4 $\times$ 75.0              | 7.91            | 410          |
| 0.0161        | 124 $\times$ 40 | 4.960         | 262.0 $\times$ 48.0              | 7.19            | 390          |
| 0.0179        | 112 $\times$ 44 | 4.928         | 236.4 $\times$ 52.9              | 6.83            | 360          |
| 0.0213        | 94 $\times$ 94  | 8.836         | 198.1 $\times$ 114.4             | 6.26            | 360          |
| 0.0227        | 88 $\times$ 56  | 4.928         | 185.3 $\times$ 67.6              | 6.06            | 280          |
| 0.0270        | 74 $\times$ 58  | 4.292         | 155.5 $\times$ 70.1              | 5.55            | 400          |
| 0.0303        | 66 $\times$ 76  | 5.016         | 138.5 $\times$ 92.2              | 5.25            | 400          |
| 0.0333        | 60 $\times$ 108 | 6.480         | 125.7 $\times$ 131.6             | 5.00            | 500          |
| 0.0370        | 54 $\times$ 110 | 5.940         | 112.9 $\times$ 134.1             | 4.74            | 600          |
| 0.0400        | 50 $\times$ 90  | 4.500         | 104.4 $\times$ 109.5             | 4.56            | 280          |
| 0.0435        | 46 $\times$ 130 | 5.980         | 95.8 $\times$ 158.7              | 4.38            | 500          |
| 0.0465        | 86 $\times$ 60  | 5.160         | 181.1 $\times$ 72.6              | 4.23            | 400          |

6. Interplay between splitting and LL broadening

It is observed in figure 1 that the splitting is much better defined through the localization properties of the states (definition of two clearly split PR peaks) than through the DOS. Nevertheless, the DOS shape observed in figure 1(b) suggests a superposition of two split bands.

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Indeed, using a multi-peak fitting procedure, shown in figures 6(a) and (b), it is found that the DOS obtained are reasonably well fitted by two overlapping Gaussian curves (apart from the small peak at $E = 0$, whose origin is not understood). In figure 6(c) it is possible to compare overall broadenings of the two $n = 0$ LL DOS, which are calculated for two different values of disorder, $W/t = 0.05$ and 0.6, keeping all the other parameters constant: lattice size is $M \times N = 66 \times 76$ and the magnetic flux considered is $\Phi/\Phi_0 = 2/66 = 0.0303$. Figure 6(d) shows that two Gaussian peaks fitted to the DOS have a good coincidence with the calculated PR peaks positions.

The splitting is observed in the previous section, through the separations of the two PR peaks, to have a square root dependence with magnetic field and to increase linearly with the disorder. However, what is important to discuss now is that, even when the splitting increases, the overlapping between the two degeneracy broken levels is pretty much the same, for a wide
range of parameters observed, as the comparison between figures 6(a) and (b) indicates. This is because the same increasing disorder or magnetic field that causes the increasing of critical energies splitting also causes the increasing broadening of each Landau band. In other words, we would require that the rate at which the splitting increases with disorder and with magnetic field is much higher than the rate at which the LLs get broadened by them, to be able to observe an increasingly well-defined separation of the DOS in two independent (not overlapped) bands. The interplay between both these effects (splitting and broadening of the bands) produces in the end an overall DOS for the \( n = 0 \) LL having always the same appearance (the difference is only the broadening, but the main features are kept the same). This indicates a direct connection between the effects of LL broadening and LL splitting. In the experiments, one consequence of this behavior is that it might be equally difficult to observe a real energy gap throughout the DOS measurements, independent of the amount of disorder or of the intensity of the magnetic field.

The random hoppings considered here are spatially uncorrelated. A further desirable extension for this work is to consider a spatial correlation for the hoppings, a situation that better emulates the ripples in real graphene sheets. In [20] it is observed that the introduction of an increasingly higher correlation length for the random hoppings in graphene results in an increasingly thinner \( n = 0 \) LL compared to the broadening of the higher LLs. On the other hand, the splitting of two critical energies within the \( n = 0 \) LL is observed in [17] for a random flux model, for which the \( n = 0 \) LL is much thinner than the higher levels (even being thinner, the DOS shows an \( n = 0 \) LL shape very similar to that observed here). The results of these two works [17, 20], together with the results shown here, suggest that the splittings in the central LL should survive when a finite correlation length for the hoppings is taken into account in the model, however with the energy splittings being scaled down with the width of the Landau band, in a fashion similar to that observed in figure 6.

7. Conclusions

The breaking of the degeneracy of the graphene \( n = 0 \) LL is investigated in this work through a simple numerical model, considering a non-interacting tight-binding model for a hexagonal lattice with random nearest-neighbor hoppings. Inferring the localization properties of the states by means of PR calculations, two clearly split critical energies are observed. The origin for this splitting has to be intrinsic to the disorder model considered, which introduces inter-valley mixing to the system.

The energy splitting is observed to have a linear dependence with disorder strength and a square root dependence with magnetic field, confirming the results obtained in [17]. The analysis of the probability densities of the wavefunctions within the \( n = 0 \) LL shows that there is a region, for the states closer to \( E = 0 \), where there is an important asymmetry in the distribution of the wavefunction amplitudes between the two sublattices. In this region, although there are amplitudes over both sublattices, there is no matching of the spatial positions of the amplitudes over each sublattice. It is as though each sublattice has its own probability density, completely independent from the other. This may lead to a reduction in the state percolation over the lattice, increasing even further the localized character of these states and also influencing anomalous localization properties. It is observed that this region of states with special wavefunction distribution between the sublattices coincides with the region where the two degeneracy broken LLs have a more important overlapping. Another interesting observation
is that this overlapping remains approximately constant for different disorder strengths, due to
the interplay between LL broadenings and LL splittings with increasing disorder or magnetic
flux. These results may help in elucidating the physics involved in the splitting of the $n = 0$ LL
due to valley/sublattice symmetry breaking.

Acknowledgments

I acknowledge fruitful discussions with Peter A Schulz, Yakov Kopelevich, Caio H
Lewenkopf and Eduardo R Mucciolo. This work is supported by FAPESP. Some of the
numerical simulations were performed on the clusters from CENAPAD/UNICAMP and
IFGW/UNICAMP.

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