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Tunable quantum dots in monolayer graphene

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We examine a graphene quantum dot formed by combining an electric and a uniform magnetic field. The electric field creates a smooth quantum well potential while the magnetic field induces an exponential tail to the dot states. The states peak in the well and the electrostatic barrier region as a result of the Klein tunneling effect. Coupling between dot states which peak in different regions can be achieved with the electric and magnetic fields. The tunability of this dot with moderate external fields could be used for designing quantum devices in monolayer graphene.

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\section{I. INTRODUCTION}

The electronic properties of monolayer graphene are derived from a two-dimensional Dirac equation. This makes graphene suitable to explore quasi-relativistic effects in condensed matter physics as well as attractive for quantum devices.\textsuperscript{1} For these two reasons various graphene-based devices are under investigation. Lithographic confinement leading to quantum dot behaviour has been achieved in nano-crystals of graphene with a diameter of some tens of nanometers. Coulomb blockade, single-electron transport, charge and spin spectroscopy have been demonstrated in single and coupled dots.\textsuperscript{1,4}

Theoretical studies have shown that the electronic properties of the dots formed in nano-crystals depend crucially on the type of edges.\textsuperscript{1} However, the edges cannot be routinely controlled experimentally, because edge disorder seems to be unavoidable and different types of edges may coexist in one sample. Forming dots in monolayer graphene is necessary in order to spatially isolate the dot states from the edges. In this case the fabrication of the dot involves the application of external fields. However, electrons in graphene are massless and via Klein tunneling can penetrate any electrostatic potential barrier.\textsuperscript{1,5} Consequently, pure electrostatic confinement in graphene is not possible\textsuperscript{6–9} and a uniform magnetic field has to be applied.\textsuperscript{9} The usual case in tuning a dot electrically cannot be exploited in graphene, though an exception may occur for zero-energy states.\textsuperscript{10}

As shown in this work, the physics of a graphene dot formed by combining an electric and a uniform magnetic field is rich and has great potential for manipulation and control over the dot system. The electric field creates a quantum well while the magnetic field provides an exponential tail to the quantum states which is needed for confinement. The electric field modifies the Landau level spectrum by inducing energy levels within the Landau gaps. These levels lie in a relatively low density of states. Here we demonstrate that the tunability of these levels can be achieved with both electric and magnetic fields.

The dot states exhibit features which arise due to the relativistic character of graphene. When the magnetic field is low, there is a class of states which peak in the quantum well region but have also a large oscillatory amplitude in the electrostatic barrier region due to Klein tunneling. As the field increases, the amplitude in the barrier decreases exponentially and the states become confined very close to the centre of the dot. The suppression of the Klein tunneling with the magnetic field may be employed in coupled dots. The interdot coupling is expected to be strong (weak) in the low (high) magnetic field regime. Strong interdot coupling seems to be possible even when the dots are separated by a long distance. There is also another class of states which peak mainly in the barrier region. These states can couple to states which have a large amplitude in the well, with a coupling strength that depends on the fields. Numerical calculations show that it should be experimentally possible to probe the dot properties in clean sheets of graphene.

The dot system studied here could allow the investigation of the Klein tunnelling effect in a well-defined two-dimensional system. In nano-crystals of graphene, the geometry is not circularly symmetric and a spectral gap opening may prevent Klein tunnelling from taking place. Moreover, the system that we study enables the formation and coupling of multiple graphene dots in various two-dimensional configurations. This may be more difficult to achieve using nano-crystals. The proposed dot could even be formed in suspended sheets of graphene in order to minimize interaction and disorder effects due to the substrate.

In Sec. II we show that some of the basic properties of the dot system can be extracted directly from the classical energy-momentum relation of massless particles. We also present the quantum mechanical model of the dot which is based on the Dirac equation for the envelope functions of graphene. Furthermore, in Sec. II a semi-analytical model is solved for a piece-wise constant quantum well potential. Calculations are presented in Sec. III, where the tunability of the dot with electric and magnetic fields is demonstrated. The conclusions of this work are given in Sec. IV.
II. GRAPHENE DOT FORMED BY ELECTRIC AND MAGNETIC FIELDS

A. Dot properties derived from the classical motion of massless particles

Consider a graphene sheet in a vector potential $\mathbf{A} = (0, A_\theta, 0)$ and an electrostatic potential $V$, and for simplicity choose $A_\theta$ and $V$ to have cylindrical symmetry. Here $V$ defines a quantum well potential centered at $r = 0$ which is modelled as $V(r) = -V_0 \exp(-r^2/d^2)$, with $V_0 \geq 0$. Also, $A_\theta$ generates a perpendicular magnetic field to the graphene sheet which, in this work, is chosen to be uniform, $\mathbf{B} = B\mathbf{\hat{z}} = \nabla \times \mathbf{A}$, thus $A_\theta = Br/2$. The classical energy-momentum relation of a massless particle moving in the fields $\mathbf{A}$ and $V$, with energy $E$ and radial momentum $p$, is

$$v_F^2 p^2 = (E - V)^2 - v_F^2 \left(\frac{M}{r} + eA_\theta\right)^2,$$  \hspace{1cm} (1)

where the term $M/r$ is due to the angular motion and $v_F = 10^6 \text{ m s}^{-1}$ is the Fermi velocity in graphene. The classical motion is restricted to the region where $p^2 > 0$, and $p^2 = 0$ defines the classical turning points. The ability of the magnetic field to induce confined states with an exponential tail, regardless of momentum and energy, is based on the observation that if $A_\theta \neq 0$ then $p^2 < 0$ in the asymptotic regime of large $r (r \rightarrow \infty)$. Therefore, classical motion is not allowed in this regime. But if $A_\theta = 0$, then asymptotically $p^2 > 0$ and classical motion is allowed, resulting in deconfined states with an oscillatory tail. In this work we are interested in the energy range $-V_0 < E < 0$ and confined states which are needed for quantum dots, henceforth $A_\theta \neq 0$.

Equation (1) shows that $p^2$ can take positive values even within the barrier region before it eventually becomes negative asymptotically. For instance this can happen when $|E - V_0|$ is large and $B$ is low. This regime can always be arranged by tuning the electric and magnetic fields. If $p^2 > 0$ in the barrier, then the quantum states are expected to have an oscillatory spatial dependence; a property which is related to the Klein tunneling effect for massless particles. In contrast, the states of a particle with mass described by a Schrödinger equation decay in the barrier.

Figure 1 shows $v_F^2 p^2$ together with the potential well. When $B = 0.3$ T, then $p^2 > 0$ almost entirely in the barrier (well) region for $E = -30 \text{ meV} (E = -8 \text{ meV})$, and hence the state is expected to be confined in the barrier (well). When $B = 0.05$ T and $E = -12 \text{ meV}$, then $p^2 > 0$ both in the well and barrier regions. In this case the state can have a large amplitude both in the well and the barrier. It can also be derived from Eq. (1) that if $B$ is high enough and/or the angular momentum term is large ($M \gg 0$) then there exists an energy range for which $p^2 < 0$ for all $r$. So classical motion is not allowed, suggesting that quantum states do not exist in this range.

Therefore, a gap, proportional to $B$ and $M$, is formed in the energy spectrum. Then as the depth of the potential well increases the gap closes and states are formed in the well and/or barrier depending on the choice of energy.

Equation (1) provides some insight into the properties of the quantum dot and shows that different classes of states can be defined according to the region which the states tend to occupy. It also demonstrates the ability to tune at will the dot states with electric and magnetic fields. However, it cannot predict the energy levels of the dot and the exact pattern of the states. For this reason we present below the exact model of the quantum dot and identify its connection to the classical energy-momentum relation in Eq. (1).

B. Quantum mechanical dot model

In monolayer graphene there are two valleys that have to be considered, at the Dirac points $K$ and $K'$, respectively, of the Brillouin zone. In this work we assume that the two valleys are uncoupled, as it has been demonstrated in many experimental studies. The physics is the same for both valleys, thus we examine the dot properties only for one valley. In the continuum approximation and for energies near the Dirac points, the two-component envelope functions $\Psi$ satisfy a two-dimensional Dirac equation

$$[v_F \sigma \cdot (p + eA) + V] \Psi = E \Psi,$$  \hspace{1cm} (2)

where $\sigma$ are the Pauli matrices, $p$ is the momentum operator and $E$ the energy. For a circularly symmetric system, defined by the vector potential $\mathbf{A} = (0, A_\theta, 0)$ and the quantum well potential $V$ of the previous subsection,
we can write for the wavefunctions
\[ \Psi = \frac{1}{\sqrt{r}} \left( f_1(r) \exp[i(m-1)\theta] + i f_2(r) \exp[im\theta] \right), \tag{3} \]
where \( m = 0, \pm 1, \ldots \) is the angular momentum number and \( r, \theta \) is the radial distance and the azimuthal angle respectively. The radial functions satisfy the equations
\[ V f_1 + \left( U + \gamma \frac{d}{dr} \right) f_2 = Ef_1, \tag{4a} \]
\[ \left( U - \gamma \frac{d}{dr} \right) f_1 + V f_2 = Ef_2, \tag{4b} \]
with \( v_F = \gamma/\hbar \) and
\[ U = \gamma \frac{2m-1}{2r} + \gamma \frac{eA_\theta}{\hbar}. \tag{5} \]
This eigenvalue problem can be diagonalised numerically to give the eigenenergy \( E \) and the corresponding two-component eigenstate \( (f_1, f_2) \). Here \( f_i \) gives the probability amplitude of finding an electron on one of the two sublattices of graphene.

Following the same method as that developed in Ref. 9, it can be shown that the state \( f_i \) satisfies the second order differential equation
\[ \frac{d^2 f_i}{dr^2} + s(r) \frac{df_i}{dr} + w_i(r) f_i = 0, \tag{6} \]
with \( s = V’/(E-V) \) and the prime denotes differentiation with respect to \( r \). Also
\[ w_i = -\frac{U^2}{\gamma^2} \pm \frac{U'}{\gamma} \pm \frac{U}{\gamma} + \frac{(E-V)^2}{\gamma^2}, \tag{7} \]
where the minus (plus) sign is for \( i = 1 \) (\( i = 2 \)). To derive a more familiar Schrödinger-like equation we eliminate the first derivative term. This can be done by writing \( f_i \) in the form
\[ f_i = u_i \exp \left( -\frac{1}{2} \int s(r) dr \right), \tag{8} \]
and substituting Eq. (8) into Eq. (6). This procedure shows that the function \( u_i \) is a solution to the equation
\[ \frac{d^2 u_i}{dr^2} + k_i^2(r) u_i = 0, \tag{9} \]
with
\[ k_i^2 = \frac{\pm U'}{\gamma} \pm \frac{U}{\gamma} - \frac{s'}{2} - \frac{s^2}{4} + \frac{v_F^2 p^2}{\gamma^2}, \tag{10} \]
where the minus (plus) sign is for \( i = 1 \) (\( i = 2 \)). Unlike the coupled equations for \( f_i \), the single equation for \( u_i \) has a more convenient form. The radial momentum \( p^2 \) in Eq. (1) can be directly identified in Eq. (10) with \( M \) replaced by \((m-1/2)\hbar \). The quantum model reveals that the region where the states are localised is defined not only by the \( v_F^2 p^2 \) term but also by the additional terms which appear in Eq. (10). Confined states, i.e., states that decay asymptotically, occur when \( k_i^2(r \rightarrow \infty) \sim -v_F^2 \gamma c^2 A_\theta^2 < 0 \), which is satisfied for \( A_\theta \neq 0 \). This condition agrees with that derived from Eq. (1).

One consequence of the Dirac equation is that the energy spectrum of the graphene dot is unbound. Depending on the external fields, the energy levels and the quantum states may exhibit simple patterns. For instance, when \( V = 0 \) and \( m \geq 1 \) the energy spectrum consists of two ladders (sets) of Landau levels (LLs) which can be determined analytically. The ladders are separated by a gap which is proportional to the field \( B \) and angular number \( m \). The LLs are formed symmetrically with respect to \( E = 0 \) and the number of nodes in the corresponding Landau states increases successively by one for each new level. For \( m \leq 0 \) one additional LL is formed at \( E = 0 \) and one of the two components \( f_i \) is zero.

The inclusion of a potential term \( V \neq 0 \) modifies the spatial region where the Landau states are localised. In addition energy levels are formed within the Landau gaps. The simplest regime was studied in detail in Ref. 15 within an approximate model. It was shown there that for \( m \geq 1 \) the dot states are approximately localised in either of the two regions defined by the two terms \( E - V \pm U \), which create two (independent) effective quantum wells. Each well contributes one ladder of energy levels to the dot spectrum. For \( m \leq 0 \) the two wells can no longer be defined because the two curves \( V = \pm U \) cross. Nevertheless, exact numerical calculations confirm the formation of the two ladders for \( m \leq 0 \) as well.

The model developed in Ref. 15 is valid provided the two effective wells have no common energy range. This can be achieved when there is an energy gap between the minimum of \( E - V + U \) and the maximum of \( E - V - U \). If this condition is not satisfied the states of the two ladders can form a richer pattern. Specifically, when \( m \) is small the states of the upper ladder tend to couple to the states of the lower ladder. This regime is the main concern of the present work and it typically occurs when the potential depth \( V_0 \) is large and the field \( B \) is low. As shown below, the induced coupled states can have a large amplitude both near the centre of the quantum well and in the barrier region. The coupling strength can be controlled at will by tuning the external fields and hence different classes of states can be probed. If \( m \) is large the states are not affected by the quantum well since they are localised in the asymptotic region where the potential is constant. As a result the states for large \( m \) behave as Landau states.

The formation of the two ladders and the coupling between the corresponding states have also been predicted theoretically in a graphene dot system at zero magnetic field but with a spatially modulated spectral (Dirac) gap in the energy dispersion. In this system the states are confined provided their energies lie inside the gap. The coupling strength can be tuned with the potential depth and the induced coupled states can be probed inside the
Spatial region where the spectral gap is zero.\textsuperscript{13}

\section*{C. Dot formed by a piecewise-constant potential well and uniform magnetic field}

This work is concerned with quantum dots formed in monolayer graphene by combining electric and magnetic fields. In this context, the simplest dot model that can be studied is when the magnetic field is uniform and the induced potential well is piecewise-constant, e.g., \( V(r) = -|v_0|\Theta(r-R) \), where \( |v_0| \) is the depth of the well, \( R \) its radius and \( \Theta \) is the step function. For completeness the method to solve this problem is outlined here, though the reader can proceed directly to Sec. III. As in Sec. II B Eqs. (4a) and (4b) are uncoupled and a second-order differential equation is derived for each radial state \( f_i \). Equation (6) for \( s = 0 \) gives

\begin{equation}
\frac{d^2 f_i}{dr^2} + \frac{(E-V)^2}{\gamma^2} f_i - \frac{U^2}{\gamma^2} f_i \pm \frac{1}{\gamma} \frac{dU}{dr} f_i = 0. \tag{11}
\end{equation}

This equation is written in the more convenient form

\begin{equation}
\frac{d^2 f_i}{dr^2} - \frac{a^2 r^4 + b_1 r^2 + c_i}{r^2} f_i = 0, \tag{12}
\end{equation}

with the coefficients \( a = eB/2\hbar \) and

\begin{align}
b_1 &= \frac{meB}{\hbar} - \frac{(E-V)^2}{\gamma^2}, \quad c_1 = (m-1)^2 - \frac{1}{4} \tag{13a} \\
b_2 &= \frac{(m-1)eB}{\hbar} - \frac{(E-V)^2}{\gamma^2}, \quad c_2 = m^2 - \frac{1}{4} \tag{13b}
\end{align}

The form of Eq. (12) reveals that the states \( f_i \) can be expressed with the help of the confluent hypergeometric functions \( U \) and \( M \) (Ref. 17). For example, when \( m \geq 1 \) the two-component state which is regular at the origin and decays asymptotically has for \( r \leq R \) the following form

\begin{equation}
\begin{pmatrix} f_1 \\ f_2 \end{pmatrix} = \eta e^{-ar^2/2} \begin{pmatrix} r^{d_1} M(A_1,B_1,ar^2) \\ c_0 r^{d_2} M(A_2,B_2,ar^2) \end{pmatrix}, \tag{14}
\end{equation}

with

\begin{equation}
c_0 = \frac{2\gamma a}{E + |v_0|} \left( 1 - \frac{A_1}{B_1} \right). \tag{15}
\end{equation}

For \( R \leq r \) the two-component state has the form

\begin{equation}
\begin{pmatrix} f_1 \\ f_2 \end{pmatrix} = \beta e^{-ar^2/2} \begin{pmatrix} r^{d_1} U(A_1,B_1,ar^2) \\ g_0 r^{d_2} U(A_2,B_2,ar^2) \end{pmatrix}, \tag{16}
\end{equation}

with

\begin{equation}
g_0 = \frac{2\gamma a}{E}. \tag{17}
\end{equation}

The auxiliary parameters are

\begin{align}
d_i &= 1 + \frac{\sqrt{4c_i + 1}}{2}, \tag{18a} \\
A_i &= \frac{b_1}{4a} + \frac{1}{2} \left( \frac{1}{2} + d_i \right), \quad B_i = \frac{1}{2} + d_i. \tag{18b}
\end{align}

The corresponding energies can be obtained from a standard matching condition which requires that both \( f_1 \) and \( f_2 \) to be continuous at \( r = R \). This condition leads to an algebraic equation for the energies which is solved numerically. Then having calculated the energies and taking also into account the normalization condition the ratio \( \beta/\eta \) is determined. Some quantum states for \( m = +1 \) are plotted in Fig. 2. The region in which the states peak depends on the magnetic field and energy. This behaviour is consistent with that derived from the energy-momentum relation in Eq. (1).
III. TUNING THE GRAPHENE DOT WITH ELECTRIC AND MAGNETIC FIELDS

In any real sample of graphene a piecewise-constant potential well cannot be generated. So a more realistic choice is made for the potential well and, specifically, the potential well is chosen to have a Gaussian form as in Sec. II A with \( d = 55 \) nm. Numerical calculations of the electrostatic potential in the graphene sheet confirm that the Gaussian potential is a very good approximation to the potential that is generated by gate electrodes or a scanning tunneling microscope tip.\(^9\) Also, the Gaussian potential produces a smooth spatial variation on the scale of the lattice constant and hence intervalley scattering can be ignored.

The results presented in this work are for the \( m = +1 \) states which can peak very close to the centre of the potential well and thus can be considered as quantum dot states. States with small values of \( m \) show similar trends to those with \( m = +1 \). As \( m \) increases, the states become localised in the asymptotic region where the potential is constant and hence the effect of the quantum well on the states becomes negligible. This property also occurs for a repulsive potential profile.\(^{18}\) Therefore, in the large \( m \) regime the corresponding quantum states are described approximately by Landau states since only the vector potential and angular momentum are important.

The dot states of interest have to be energetically isolated from other states, that is, they have to lie in a region of low density of states (DOS), in order to be detectable. If this condition is satisfied then scanning tunneling microscopy (STM) could be used to probe the local density of states and features of individual quantum states. When there is no potential variation and a perpendicular magnetic field is applied to a graphene sheet confirm that the potential produces a smooth spatial variation on the scale of the potential and angular momentum are important.

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We assume that for the low magnetic fields considered in this work there is no overlap between the LLs. Any LL broadening, for example, due to disorder and interaction effects\(^{19,20}\) is small and it is typically less than \( \sim 2 \) meV. It may be possible to achieve this condition in clean samples of graphene. In particular, experiments in suspended sheets of graphene have measured very large mobilities indicating that these sheets are less sensitive to disorder and impurities introduced by the substrate.\(^{21,22}\)

![FIG. 3: Energy levels (dashed curves) as a function of the potential depth \( V_0 \), for \( m = +1 \) and a magnetic field \( B = 0.1 \) T. Some Landau levels are also shown (horizontal solid lines). The quantum states for the marked energies are shown in Fig. 4.](image)

A. Electric field effect on the dot

As discussed in Sec. II B, in the graphene dot system formed by electric and magnetic fields, the energy levels for a fixed value of \( m \) can form two energy ladders separated by a gap.\(^{16}\) This configuration can be achieved for a high magnetic field and a small well depth. When the well depth increases, energy levels from the upper energy ladder fall in the gap, while the corresponding states develop a large amplitude in the well region. Eventually, when the well depth is large, states which peak in the well couple to states of the lower energy ladder, and this behaviour is manifested in the energy diagram by the formation of anticrossing points.\(^{23}\)

To demonstrate these effects, we show in Fig. 3 the energy diagram as a function of the potential depth \( V_0 \) for a magnetic field \( B = 0.1 \) T. For \( V_0 = 0 \) the energy levels correspond to the LLs of a graphene sheet. For a small \( V_0 \), the energy levels which define the two energy ladders merge into the Landau gaps. When \( V_0 \sim 0 \) these levels are separated by a gap which is approximately equal to the energy splitting between the LLs \(+1\) and \(-1\) (Ref. 24). As \( V_0 \) increases, the gap closes and eventually energy levels of the upper ladder anticross with energy levels of the lower ladder. The general trend is that if the magnetic field is low the gap between the two energy ladders is small and therefore the anticrossing points appear for a small well depth \( V_0 \). For instance, the first anticrossing point at \( B = 0.1 \) T is formed for \( V_0 \approx 60 \) meV (see Fig. 3), while at \( B = 0.2 \) T for \( V_0 \approx 90 \) meV. With a further increase of \( V_0 \), a state of the lower energy ladder couples successively to states with higher energy in the well, forming a series of anticrossing points. Thus the required variation of \( V_0 \) to probe two successive anticrossing points becomes smaller as the typical energy...
splitting in the well decreases.

Here we focus on the state with the highest energy in the lower ladder and its coupling to quantum well states. This state corresponds for $V_0 = 0$ to the LL $-1$ and then for $V_0 \neq 0$ the coupling leads to a “hybridised” state with energy between the LLs $-1$ and $-2$. Some examples of this hybridised state are shown in Fig. 4. Comparison of the states in the left panels with those in the right panels shows that the number of nodes for each radial component increases by one in the well region for $V_0 = 0$ to the LL $-1$ and $-2$. Some examples of this hybridised state are shown in Fig. 4. Comparison of the states in the left panels with those in the right panels shows that the number of nodes for each radial component increases by one in the well region for $V_0 = 0$ to the LL $-1$ and $-2$. Some examples of this hybridised state are shown in Fig. 4. Comparison of the states in the left panels with those in the right panels shows that the number of nodes for each radial component increases by one in the well region. Near the anticrossing points the coupling is strong and hence the amplitude of each state in the well decreases, e.g., for $V_0 = 40$ meV and $90$ meV.

As seen in Fig. 4, the hybridised state has a peak in the barrier region which, to a good approximation, is insensitive to $V_0$. This happens because the charge carriers in graphene are massless and they exhibit Klein tunneling. This allows the states to develop an oscillatory amplitude in the barrier. In conventional dots obeying Schrödinger’s equation the quantum states decay in the barrier. Klein tunneling takes place in the two-dimensional graphene dot when within a barrier region $k^2 > 0$ in Eq. (10). This condition can be arranged easily for the dot system by tuning the external fields. Furthermore, the exact pattern of the state in the barrier depends on the choice of energy. Specifically, the number of peaks within the barrier increases for states which lie between higher excited LLs and thus they have the dependence $E_N \propto -\sqrt{B/N}$. Between the LLs there are regions where only a few levels lie, suggesting that the corresponding states could be probed with STM measurements. For this reason the energy of the hybridised state in Fig. 4 is adjusted between the LLs $-1$ and $-2$ where the DOS is relatively low. Inspection of Fig. 5 shows that there are ranges of $V_0$ in which the energy of this state is at least 0.5 meV away from any other energy levels. Thus it should be experimentally possible to probe the pattern of the state in Fig. 4 in clean samples of graphene with a small broadening of the LLs. The local DOS near the centre of the dot is expected to show a low-high-low variation as $V_0$ sweeps through an anticrossing point.

When the broadening of the LLs is large, the regions of low DOS which occur between the LLs shrink making the detection of individual dot states difficult; the states have to lie in a very narrow energy range (window) in order that to be detectable. A high magnetic field $B$ may increase this energy range provided that any broadening of the LLs increases much slower than the Landau gaps. However, when $B$ is high a large $V_0$ is needed for anticrossing points to appear and the states to follow the

FIG. 4: (Color online) Quantum states for the marked energies shown in Fig. 3. The potential depth is $V_0$. Solid (dashed) curves show $|f_i|^2$ ($|f_2|^2$). The potential well (in arbitrary units) is also shown.
B. Magnetic field effect on the dot

The effect of a magnetic field on the dot is now examined when the potential well is fixed. In Sec. III A it was shown that, when the electric field increases, states belonging to different energy ladders become coupled. Moreover, the coupling is strong near the anticrossing points. In this section it is demonstrated that when the magnetic field increases the coupling is suppressed resulting in states which correspond approximately to individual states in each ladder.

In Fig. 6 the energy diagram is plotted as a function of the magnetic field when the depth of the potential well is $V_0 = 51$ meV. We are interested in the two highest energy levels below zero (for $m = +1$) which are denoted by $E_1$ and $E_2$. In the field range $0.04 < B < 0.5$ T, the $E_1$ level lies between the LLs 0 and −1, while for $0.04 < B < 0.35$ T the $E_2$ level lies between the LLs −1 and −2. Thus quantum states can be arranged to lie between successive LLs in a wide field range. Calculations show that this effect is robust and can be achieved for different well depths and $m$ values. In addition, as shown below within a specific magnetic field range the spatial region in which the states are localised changes drastically, confirming that the tunability of the dot is feasible with a uniform magnetic field.

Figure 7(a) shows the quantum state corresponding to the energy level $E_2$ in Fig. 6. This “hybridised” state is formed due to the coupling between the states of the two energy ladders. In particular, the peak in the well (barrier) region is due to the state in the upper (lower) ladder. When $B = 0.04$ T the coupling is strong and the hybridised state peaks both in the well region, that is $r \lesssim 80$ nm, and the barrier. As $B$ increases, the coupling is suppressed and the amplitude of the hybridised state very close to the centre of the well decreases. Consequently, the hybridised state takes approximately the form of the state in the lower energy ladder. Also, when the field increases the magnetic confinement becomes stronger, therefore the state in the lower ladder shifts...
because there is only a small region with involved in the low field regime but is not so pronounced but has also a large amplitude in the barrier region before the barrier. Then as $B$ increases the amplitude of the states in each region decreases. This produces a function of the magnetic field $B$.

The magnetic-field-dependent coupling is also responsible for the behaviour of the “hybridised” state corresponding to the energy level $E_1$ [see Fig. 7(b)]. At $B = 0.04$ T the state peaks in the well region $r \lesssim 80$ nm, but has also a large amplitude in the barrier region before it eventually decays asymptotically. Klein tunneling is involved in the low field regime but is not so pronounced because there is only a small region with $k_f^2 > 0$ within the barrier. Then as $B$ increases the amplitude of the two components in the barrier is gradually suppressed and the state becomes localised near the centre of the well. For $B = 0.5$ T the hybridised state has approximately the form of the state in the upper energy ladder and $k_f^2 < 0$ everywhere in the barrier. Thus no Klein tunneling occurs at high $B$. The regime of high magnetic field as well as the symmetry between a dot and an antidot system have been investigated theoretically in Ref. 27. A graphene dot system especially designed to probe the Klein tunneling effect in a two-dimensional geometry has been studied in Ref. 9.

Figure 7 demonstrates that the magnetic field-induced suppression of the amplitude of the states in the barrier is a strong effect. This property may be useful for a system of two neighboring graphene dots separated by a long distance. In this system the interdot coupling can be tuned with the magnetic field instead of using gate electrodes as in conventional dots. The coupling is expected to be strong in the low field regime, provided Klein tunneling is involved. The coupling can be strong even when the two dots are far apart, a situation that cannot be arranged easily in conventional semiconductor dots.

In Fig. 8 we show the energy diagram of the dot as a function of the magnetic field $B$. All values of $m$ are taken into account, but as explained above, when $m$ is large the energy levels have the LL dependence $E_N \propto -\sqrt{BN}$. As seen in Fig. 8, regions of very low DOS are formed between the LLs. This condition can be arranged when the quantum well is narrow enough so that only a limited number of $m$ values contributes energy levels between the LLs. The energy level $E_1$ in Fig. 6 crosses some other levels for a few values of $B$ but it is isolated from any other levels in a wide field range. This property makes possible the resolution of the corresponding state and its variation with the magnetic field [Fig. 7(b)]. Investigation of the energy diagram in Fig. 8 shows that for $0.04 < B < 0.3$ T the energy level $E_2$ in Fig. 6 is at least 0.4 meV away from any other levels. This suggests that the corresponding quantum state [Fig. 7(a)] could be detected using similar STM measurements, for example, as those in Ref. 25.

### IV. Discussion and Conclusion

This work investigated the properties of a graphene quantum dot formed by the combination of electric and magnetic fields. The electric field creates a smooth quantum well potential which could be generated using gate electrodes. Because of the Klein tunneling effect in graphene the well cannot confine electrons. However, when a uniform magnetic field is applied perpendicular to the graphene sheet the states decay asymptotically as needed for confinement. The electric field modifies the Landau level spectrum by inducing levels within the Landau gaps. It was demonstrated that the states which correspond to these levels are tunable with the electric and magnetic fields. This property has also been considered in one-dimensional systems of graphene such as wave-guides and wires.28,29

Some of the basic properties of the dot system were qualitatively extracted from the classical energy-momentum relation of massless particles. These properties were then confirmed and quantified with exact numerical calculations. It was found that some states can peak both in the potential well and within the barrier because of the Klein tunneling effect in graphene. This behaviour cannot be observed in a non-relativistic dot described by a Schrödinger equation. States which peak in the barrier region occur also in one-dimensional systems of graphene and have been examined in Refs. 30 and 31.

The relative amplitude of the states in each region depends on the values of the electric and magnetic fields and thus it can be tuned at will. For example, at a high magnetic field there are states which peak very close to the centre of the well and decay exponentially in the barrier. For low enough fields, Klein tunneling is involved.
and the amplitude of the states in the barrier increases drastically. Coupling between states which peak in different regions results in the formation of anticrossing points in the energy spectrum of the dot. Numerical calculations suggest that the density of states is relatively low between the Landau levels. Therefore, it should be experimentally possible to probe individual quantum states when the broadening of the Landau levels is small.

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