Isolated and hybrid bilayer graphene quantum rings

M. Mirzakhani,1,2,∗ D. R. da Costa,3,† and F. M. Peeters4,‡

1School of Physics, University of the Witwatersrand, Johannesburg, Wits 2050, South Africa
2Center for Theoretical Physics of Complex Systems, Institute for Basic Science, Daejeon, 34126, South Korea
3Departamento de Fisica, Universidade Federal do Ceará, Campus do Pici, 60455-900 Fortaleza, Ceará, Brazil
4Department of Physics, University of Antwerp, Groenenborgerlaan 171, B-2020 Antwerp, Belgium

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Using the continuum model, we investigate the electronic properties of two types of bilayer graphene (BLG) quantum ring (QR) geometries: (i) an isolated BLG QR and (ii) a monolayer graphene (MLG) with a QR put on top of an infinite graphene sheet (hybrid BLG QR). Solving the Dirac-Weyl equation in the presence of a perpendicular magnetic field and applying the infinite-mass boundary condition at the ring boundaries, we obtain analytical results for the energy levels and corresponding wave spinors for both structures. In the case of isolated BLG QR, we observe a sizeable and magnetically tunable band gap which agrees with the tight-binding transport simulations. Our analytical results also show the intervalley symmetry $E_{\pm}^m(m) = -E_{\mp}^m(m)$ between the electron (e) and hole (h) states ($m$ being the angular momentum quantum number) for the energy spectrum of the isolated BLG QR. The presence of interface boundary in a hybrid BLG QR modifies drastically the energy levels as compared to that of an isolated BLG QR. Its energy levels are tunable from MLG dot, to isolated BLG QR, and to MLG Landau energy levels as magnetic field is varied. Our predictions can be verified experimentally using different techniques such as by magnetotransport measurements.

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I. INTRODUCTION

Over the past years, a family of two-dimensional (2D) graphene nanostructures, including graphene nanoribbons [1–6], quantum rings (QRs) [7–12], quantum dots (QDs) [13–27], and antidots [28–32] with different type of geometries, edge types, and stackings of graphene layers have received increasing interest. These studies showed that the electronic and optical properties of graphene QDs can be modified by size, shape, edge type, and electrostatic gating; see, e.g., also Refs. [13, 16, 17, 20, 26]. The effect of twisting on the electronic and transport properties of bilayer graphene (BLG) nanostructures has also been recently addressed in Refs. [33–39]. Within today’s technology, such as nanolithography, it is possible to realize such 2D nanostructures on a scale of a few tens [40, 41] or even only a few [42, 43] nanometers as well as, in different types of graphene layer stackings [44, 45].

Furthermore, bandstructure engineering can be performed by creating periodic arrays of holes in both MLG [29, 30, 32, 40] and BLG [46] sheets, known as graphene antidot lattices. Depending on the size and the period of the holes, such graphene nanostructures render graphene semiconducting with a sizeable band gap which displays a wide range of electronic and optical properties [46–48]. Very recently, a related system has been investigated where a MLG sheet consists of highly regular triangularly arranged holes [32]. The results showed that the structure displays both insulating behaviour and ballistic transport in excellent agreement with analytical calculations which describes the structure as a quantum system consisting of connected “Dirac rings” [32, 49]. In this model, Dirac fermions are strongly confined in a MLG QR geometry using an infinite-mass (IM) potential. While in the case of MLG such holes can act as scattering centers, several experimental works [50, 51] demonstrated that in the case of BLG, the adjacent layers can connect with each other thus resulting in the formation of periodic arrays of connected edgeless Dirac rings in BLG structure.

Here, we aim to investigate the electronic properties of such rings in BLG as an individual QR defined by an IM potential (isolated ring) as shown in Fig. 1(a). Of course in graphene nanostructures, the type of edge plays an important role and their effects are well known, in particular the existence of a zero-mode state at zigzag edges [13, 52, 53]. Using the IM boundary condition removes the edge effects and has the advantage that analytical results can be obtained while still representing a real system [27]. Theoretically, the IM boundary condition for confining neutrinos in a hard-wall billiard was derived by Berry and Modragon [54]. This boundary condition was previously employed to investigate the electronic properties of MLG nanostructures (dot, antidot, ring) [27, 31, 49, 55], BLG QDs [61], and trilayer graphene QDs [25]. Previous studies of BLG QRs based on Dirac equation includes electrostatically defined BLG QRs [62, 63] which was solved numerically in Ref. [62] and was mod-

∗ mirzakhani@ibs.re.kr
† diegorabelo@fisica.ufc.br
‡ francois.peeters@uantwerpen.be
eled as zero-width-ring geometry in Ref. [63]. However, at present there is, to our knowledge, no theoretical study on the energy spectrum of BLG QRs with IM boundary condition that models a realistic ring.

In the present work, we solve the Dirac-Weyl equation in the presence of a perpendicular magnetic field and apply IM boundary condition at the ring boundaries to obtain analytical results for the energy levels and corresponding wave functions. In the case of isolated BLG QR, we find an excellent agreement between the analytical predictions for the size of the band gap as a function of the magnetic field with that found in the conductance of two-terminal QR device simulated using the KWANT package [64] based on Landauer-Büttiker formalism and the tight-binding model (TBM). This is in contrast with previous study of zero-width BLG QR [63], showing a fixed energy band gap as a function of magnetic field. Our analytical results also show that the energy spectrum of the isolated ring exhibits intervalley symmetry $E_e^K(m) = -E_h^{K'}(m)$ for the electron (e) and hole (h) states where $m$ is the angular momentum quantum number.

The agreement between the obtained analytical results and TB simulations prompted us to consider also a hybrid BLG QR where a MLG ring is put on top of an infinite MLG sheet [Fig. 1(b)]. Experimental realization of such 2D heterostructures can be challenging but is doable within today’s technology. Defining the ring layer by a staggered site-dependent IM potential (e.g., using an anti-ring shaped hexagonal boron nitride as a substrate) is a way which can be used to realize the hybrid BLG QR. The hybrid structure can also be realized by (accidental) nanostructuring one of the graphene layers in BLG. For instance, topographic images have revealed that multilayer samples exfoliated from graphite often contain atomic steps and islands of one or few layers of graphene [65–69]. They have been previously investigated both theoretically and experimentally, in different configurations such as single MLG-BLG junction [70–76], double MLG-BLG junctions (MLG-BLG-MLG) [77, 78], and hybrid QD structures [79]. In all these studies, the interface between MLG and BLG regions was considered as zigzag or armchair junctions which modifies considerably the electronic properties of such structures.

A striking feature of the hybrid BLG QR is that the energy levels of the ring seem to respond to the IM potential as well as that of the layers in the LDOS can be different. This feature can be used in valleytronics applications of such graphene-based nanostructures if valley mixing is precluded.

### II. THEORY AND MODEL

We consider two different BLG nanostructures in the presence of a perpendicular magnetic field: (i) isolated BLG QR defined by a site-dependent staggered media [Fig. 1(a)] and (ii) hybrid BLG QR sandwiched between a MLG QD and an infinite MLG region, as shown in Fig. 1(b). The latter one can also be regarded as an infinite MLG sheet on which a second MLG ring is sitting on top of the first, thus realizing a BLG QR in the AB-stacking (Bernal) configuration. Dirac equation is solved for both MLG and BLG regions, with appropriate boundary conditions. By employing the IM boundary condition, we obtain analytical results for the energy levels and corresponding wave functions in each structure.

Experimentally, such a mass potential can be induced by sandwiching the BLG sheet between substrates such that the $A$ and $B$ sublattices in each graphene sheet feel different potentials [80, 81]. Equivalently, graphene nanostructures that are etched out of graphene sheets exhibit a strong confinement that can be modelled with IM-boundary conditions.

In the presence of a perpendicular magnetic field $B = B_\zeta$, the dynamics of carriers in the honeycomb lattice of carbon atoms of MLG can be described by the following Hamiltonian [7],

$$H = \nu_F \Pi \cdot \sigma + \Delta(\mathbf{r}) \sigma_z,$$  

where $\nu_F \approx 10^6 \text{ m/s}$ is the Fermi velocity, $\Pi = \mathbf{p} + e\mathbf{A}$ is the 2D kinetic momentum operator with $\mathbf{p} = -i\hbar(\partial_x, \partial_y)$, $-e$ being the electron charge, and $\mathbf{A} = (B/2)(-y\mathbf{e}_x + x\mathbf{e}_y)$ is the vector potential taken in the symmetric gauge. $\sigma = (\sigma_x, \sigma_y, \sigma_z)$ denotes the Pauli matrices and $\Delta(\mathbf{r})$ is a position-dependent mass term. In polar coordinates $(r, \varphi)$, the Hamiltonian (1) reduces to the form

$$H = E_0 \left( \begin{array}{cc} \delta & \Pi_z \\ \Pi_+ & -\delta \end{array} \right),$$  

FIG. 1. Schematic pictures of the proposed circular BLG QRs with inner and outer radii of $R_1$ and $R_2$, respectively. (a) Isolated BLG QR and (b) hybrid BLG QR sandwiched between MLG QD and MLG sheet.
where \( E_0 = \sqrt{2}h v_F / l_B \) is the cyclotron energy with \( l_B = \sqrt{\hbar / e B} \) the magnetic length, \( \delta = \Delta(p) / E_0 \), and the momentum operator

\[
\Pi_{\pm} = \Pi_x \pm i \Pi_y = -ie^{\pm irx} \frac{1}{2} \left[ \frac{\partial}{\partial \rho} \pm \frac{i \tau}{\rho} \frac{\partial}{\partial \phi} \mp \tau \rho \right]. \tag{3}
\]

Here, \( \rho = r / \sqrt{2} l_B \) is a dimensionless radial coordinate and \( \tau = \pm 1 \) distinguish the \( K \) and \( K' \) valleys. Because of circular symmetry, the two-component spinor wave function becomes \( \Psi^r(\rho, \phi) = e^{im\phi}[\phi_A^r(\rho), i\phi_B^r(\rho)]^T \), where the radial dependence of the spinor components is described by

\[
\frac{1}{2} \left[ \frac{\partial}{\partial \rho} + \frac{(\tau m + 1)}{\rho} + \tau \rho \right] \phi_B^r(\rho) = (\epsilon - \delta) \phi_A^r(\rho), \tag{4a}
\]

\[
\frac{1}{2} \left[ \frac{\partial}{\partial \rho} - \frac{\tau m}{\rho} - \tau \rho \right] \phi_A^r(\rho) = -(\epsilon + \delta) \phi_B^r(\rho), \tag{4b}
\]

where \( m = 0, \pm 1, \pm 2, \ldots \) denotes the angular momentum label and \( \epsilon = E / E_0 \) being the dimensionless carrier energy. Decoupling the above equations and using the ansatz for \( \phi_A^r(\rho) = \rho^{-m} e^{-\rho^2 / 2} f(\rho^2) \), one arrives at the associated Laguerre differential equation \( (\tilde{\rho} = \rho^2) \)

\[
\tilde{\rho} f''(\tilde{\rho}) + (m + 1 - \tilde{\rho}) f'(\tilde{\rho}) + \lambda f(\tilde{\rho}) = 0, \tag{5}
\]

where

\[
\lambda = \frac{1}{2} [-(\tau + 1) + 2(\epsilon^2 - \delta^2)]. \tag{6}
\]

The general solution to the associated Eq. (5) is

\[
f(\rho^2) = C_1 L_\chi^{-m}(\rho^2) + C_2 U(-\lambda - 1, m, \rho^2), \tag{7}
\]

where the constants \( C_1 \) and \( C_2 \) are determined by the boundary conditions, \( U(a, b, x) \) is the confluent hyperegeometric function of the second kind and \( L_\chi^b(x) \) is the generalized Laguerre polynomial which can be defined in terms of the confluent hyperegeometric function of the first kind \( M(a, b, x) \) [an alternating notation is \( F_1(a, b, x) \)] as

\[
L_\chi^b(x) = \left( \begin{array}{c} a + b \\ a \end{array} \right) M(-a, b + 1, x), \tag{8}
\]

where \( \left( \begin{array}{c} a + b \\ a \end{array} \right) \) is the generalized binomial coefficient. Notice that \( \rho^{-m} e^{-\rho^2 / 2} L^m_\chi(\rho^2) \) and \( \rho^{-m} e^{-\rho^2 / 2} U(-\lambda - 1, -m, \rho) \) converge to finite values in the limits \( \rho \to 0 \) and \( \rho \to \infty \), respectively. So, depending on the geometry of the graphene nanostructures (dot or antidot), one can choose the appropriate wave functions to satisfy the corresponding boundary conditions.

The other spinor component of the wave function, \( \phi_B^r(\rho) \), can be obtained using Eq. (4b) and by employing the properties of \( U(a, b, x) \) and \( L_\chi^b(x) \). Thus the spinor components become

\[
\phi_A^r(\rho) = \rho^{-m} e^{-\rho^2 / 2} \times \left[ C_1 L^m_\chi(\rho^2) + C_2 U(-\lambda - 1, -m, \rho^2) \right], \tag{9a}
\]

and

\[
\phi_B^r(\rho) = \frac{-\tau}{\epsilon + \delta} \rho^{-m-m-\tau} e^{-\rho^2 / 2} \left[ C_1 (\lambda + \tau_+) L^{-m-\tau}_\chi(\rho^2) - C_2 (\lambda + \tau_+) U(-\lambda - \tau, -m + 1 - \tau, \rho^2) \right], \tag{9b}
\]

where we have defined \( \tau_\pm = (1 \pm \tau) / 2 \).

The BLG region can be described in terms of four sub-lattices, labeled \( A1, B1 \), for the lower layer and \( A2, B2 \), for the upper layer. We only include the coupling between two atoms stacked on top of each other, e.g., \( B1 \) and \( A2 \), and ignore the small contributions of the other interlayer couplings. Additional terms only cause small effects such as trigonal wrapping and electron-hole symmetry on the energy levels \[82\]. The effective Hamiltonian is \[82, 83\]

\[
\mathcal{H}_B = E_0 \begin{pmatrix} \delta & -\delta & 0 & 0 \\
-\delta & \gamma_1 & 0 & 0 \\
0 & \gamma_1 & \delta & 0 \\
0 & 0 & 0 & \delta \end{pmatrix}, \tag{10}
\]

where \( \gamma_1 = \gamma_1 / E_0 \), with \( \gamma_1 \approx 0.4 \text{ eV} \) being the nearest-neighbor interlayer coupling term. Solving the Dirac equation \( \mathcal{H}_B \Phi = E \Phi \) for the four-component wave function

\[
\Phi^r(\rho, \varphi) = e^{im\varphi} \times [e^{-i\tau\varphi} \phi_{A1}(\rho), i\phi_{B1}(\rho), i\phi_{A2}(\rho), e^{i\tau\varphi} \phi_{B2}(\rho)]^T, \tag{11}
\]

the radial dependence of the spinor components in BLG are described by

\[
\frac{1}{2} \left[ \frac{d}{d \rho} + \frac{\tau m}{\rho} + \tau \rho \right] \phi_{B1}^r(\rho) = (\epsilon - \delta) \phi_{A1}^r(\rho), \tag{12a}
\]

\[
\frac{1}{2} \left[ \frac{d}{d \rho} - \frac{\tau m - 1}{\rho} - \tau \rho \right] \phi_{A1}^r(\rho) - \gamma_1 \phi_{A2}^r(\rho) = -(\epsilon + \delta) \phi_{B1}^r(\rho), \tag{12b}
\]

\[
\frac{1}{2} \left[ \frac{d}{d \rho} + \frac{\tau m + 1}{\rho} + \tau \rho \right] \phi_{B2}^r(\rho) - \gamma_1 \phi_{B1}^r(\rho) = -(\epsilon - \delta) \phi_{A2}^r(\rho), \tag{12c}
\]

and

\[
\frac{1}{2} \left[ \frac{d}{d \rho} - \frac{\tau m}{\rho} - \tau \rho \right] \phi_{A2}^r(\rho) = (\epsilon + \delta) \phi_{B2}^r(\rho). \tag{12d}
\]

Decoupling the system of equations (12) and using \( \phi_{A2}^r(\rho) = \rho^{-m} e^{-\rho^2 / 2} g^r(\rho^2) \), we arrive at the following associated Laguerre differential equation \( (\tilde{\rho} = \rho^2) \)

\[
\tilde{\rho} g''(\tilde{\rho}) + (-m + 1 - \tilde{\rho}) g'(\tilde{\rho}) + \alpha_{\pm}(\epsilon) g(\tilde{\rho}) = 0, \tag{13}
\]

where \( \alpha_{\pm}(\epsilon) \) is given by

\[
\alpha_{\pm}(\epsilon) = \frac{1}{2} \left[ (\epsilon^2 - \delta^2) - 1 \pm \sqrt{1 + 4\gamma_1^2(\epsilon^2 - \delta^2)} \right]. \tag{14}
\]

Accordingly, similarly to the MLG region, \( \phi_{A2}^r(\rho) \) can be expressed in terms of \( L_{a}^b(x) \) and \( U(a, b, x) \) as follows:

\[
\phi_{A2}^r(\rho) = \rho^{-m} e^{-\rho^2 / 2} \times \sum_{\mu=\pm} \left[ C_{\mu}^a L_{-a}^{-m}(\rho^2) + C_{\mu}^b U(-\alpha_\mu, -1 - m, \rho^2) \right], \tag{15}
\]
where the constants $C_1^a$ and $C_2^a$ are determined by the boundary conditions. The other spinor components of the wave function can be obtained using Eqs. (12) by inserting $\phi_{A2}^r(\rho)$ and employing the properties of $U(a, b, x)$ and $L_\alpha^\rho(x)$ functions. It is possible to express the other components in a compact form as follow:

\begin{align}
\phi^r_{A1}(\rho) &= \frac{-\tau \rho^{-m+\tau} e^{-\rho^2/2}}{\gamma_1(\epsilon - \delta^2)} \sum_{\mu = \pm} \eta_\mu \left[ C_1^\mu(\alpha_\mu + \tau_-)^+ L_{\alpha_\mu+\tau_-}(\rho^2) - C_2^\mu(\alpha_\mu + \tau_-)^+ U(-\alpha_\mu + \tau, -m + 1 + \tau, \rho^2) \right], \\
\phi^r_{B1}(\rho) &= \frac{\rho^{-m-\tau} e^{-\rho^2/2}}{\gamma_1(\epsilon + \delta)} \sum_{\mu = \pm} \eta_\mu \left[ C_1^\mu L_{\alpha_\mu}(-\rho^2) + C_2^\mu U(-\alpha_\mu, -m + 1, \rho^2) \right], \\
\phi^r_{B2}(\rho) &= \frac{\tau \rho^{-m-\tau} e^{-\rho^2/2}}{\epsilon + \delta} \sum_{\mu = \pm} \left[ C_1^\mu(\alpha_\mu + \tau_+)^+ L_{\alpha_\mu+\tau_+}(\rho^2) - C_2^\mu(\alpha_\mu + \tau_+)^+ U(-\alpha_\mu - \tau, -m + 1 - \tau, \rho^2) \right],
\end{align}

where $\eta_\mu = [\epsilon^2 - \delta^2 - (\alpha_\mu + \tau_+)]$.

Berry and Modragon derived the IM boundary condition for the confinement of neutrinos in a hard-wall billiard described by the Dirac-Weyl equation [54]. Let’s consider a particle restricted in the plane $r = (x, y)$ subjected to a mass term potential $\Delta(r)$, which is vanishing inside a certain domain and equal to $\Delta \to \infty$ outside it. Solving the Dirac equation, Eq. (1), for a two-component spinor $|\psi_1(r), \psi_2(r)|^T$ leads to the following relation at the domain edge [54, 55]

$$
\psi_2(r)/\psi_1(r) = i e^{\text{i} r \theta},
$$

where $\theta$ is the polar angle of the normal vector pointing outward from the domain boundary. Within the next section, we will calculate the energy spectrum of both ring structures using the above-mentioned boundary condition.

It is worth mentioning that the IM boundary condition [Eq. (17)] does not necessarily imply that the total wave function, as well as the wave function components, go to zero at the boundaries of the quantum confined system. The use of similar Berry and Mondragon-like boundary condition to explain experimental measurements in graphene-based quantum confinement nanostructures has been reported in the literature [7, 32, 42, 55], showing a good agreement between the predicted theoretical results and the electronic properties measured experimentally. Furthermore, BLG based quantum systems with reconstructed edges present different types of edge, in addition to the conventional armchair and zigzag terminations, leading to different boundary conditions different from that assumed here [56–60].

III. NUMERICAL RESULTS

A. Analytical calculations

In the case of isolated BLG QR, applying the IM boundary condition (17) for each layer, the spinor components at the radial distances $R_1$ ($\theta = \pi + \varphi$) and $R_2$ ($\theta = \varphi$) satisfy the conditions

$$
\phi^r_{B1}(\rho_1) + \phi^r_{A1}(\rho_1) = 0, \quad \phi^r_{B2}(\rho_1) - \phi^r_{A2}(\rho_1) = 0, \quad \phi^r_{B1}(\rho_2) - \phi^r_{A1}(\rho_2) = 0, \quad \phi^r_{B2}(\rho_2) + \phi^r_{A2}(\rho_2) = 0,
$$

respectively, with $\rho_i = R_i/\sqrt{2} l_D$ ($i = 1, 2$). The eigenvalue condition is determined by inserting the obtained spinors for BLG, Eqs. (15) and (16), with $\delta = 0$ into the above four equations. Note that the effect of IM potential $\Delta \to \infty$ is now expressed by the boundary conditions (18) and (19).

For the hybrid BLG QR, the spinors corresponding to the sublattices in the lower layer are continuous at the ring boundaries, while the spinor components of the upper (ring) layer, $\phi^r_{A2}(\rho)$ and $\phi^r_{B2}(\rho)$, satisfy the IM boundary condition expressed by Eqs. (18) and (19). Thus, in this case, the boundary conditions at $R_{1(2)}$ read

$$
\phi^r_{A}(\rho_{1(2)}) - \phi^r_{A1}(\rho_{1(2)}) = 0, \quad \phi^r_{B}(\rho_{1(2)}) - \phi^r_{B1}(\rho_{1(2)}) = 0, \quad \phi^r_{B2}(\rho_{1(2)}) - \phi^r_{A2}(\rho_{1(2)}) = 0,
$$

where $- (+)$ is used at $R_{1(2)}$ boundary.

Here, we have to stress that, using the above boundary conditions when implementing them numerically gives two sets of energy levels. A set of levels corresponds to
Background colored plot shows a 2D conductance of the same ring as functions of the magnetic field. The results are plotted for several angular momenta $m = -13, \pm1, \pm2, \pm1, 0$. The energy spectrum of an isolated BLG QR, as a function of the magnetic field, has also been noticed in the presence of the magnetic field: (i) For a specific angular momentum $m$, the electron-hole (e-h) symmetry is broken, i.e., $E_e(m) \neq -E_h(m)$ and (ii) $E_i(m) \neq E_i(-m)$ ($i = e, h$), since time-reversal symmetry (TRS) is broken by the magnetic field [7]. Both symmetries are restored at $B = 0$ as expected. For large field strength, the magnetic-field dependence of the spectrum becomes approximately linear which was also observed for the MLG ring and antidot spectra [49]. Note that, as function of the magnetic field, the lowest electron and hole energy levels show a “Mexican-hat” shape similar as the Mexican-hat-shaped low-energy dispersion for a biased AB-stacked BLG [84], which is due to the fourth-order character of the dispersion relation [Eq. (14)]. Including the energy levels of both valleys [see Fig. 6(b)] we can see that the confinement-induced band gap closes with increasing magnetic field. This is in contrast with the previous study of the zero-width BLG QR [63], which shows a constant energy band gap as a function of the magnetic field. A similar result for the strong dependence of the energy gap on the magnetic field in the finite-width MLG QR as well as a comparative study with the ideal zero-width one was also discussed in Ref. [11].

To see the connection between the obtained analytical energy levels and observable physical quantity like conductance, we show in Fig. 2(c), a 2D color plot of two-terminal conductance $G$ of the same ring as functions of $B$ field and energy using the KWANT package [64] within the Landauer-Büttiker formalism using the TBM (for details of the the TB simulation see Sec. III B). We find excellent agreement between the analytical results and the tight-binding transport simulations. Specifically, the closing of the band gap as a function of $B$ field in both calculations is seen to agree remarkably well. In addition, one can see high conductance values for large energies ($|E| \gtrsim 0.2\text{ eV}$) indicating the presence of a large number of conducting channels at these energies. This is
consistent with the analytical results where we find a high concentration of energy states at high energies. Notice that, in Fig. 2(c), we plotted only the energy levels for a few values of \( m \) and only for one \( K \) valley [a complete spectrum is shown in Fig. 6(b)].

Results for the energy spectrum of a hybrid BLG QR, of radius \( R_1 = 30 \) nm and \( R_2 = 40 \) nm, as a function of magnetic field \( B \) at both valleys \( K \) and \( K' \) are shown in Figs. 3(a) and 3(b), respectively. The angular momenta are \( m = -1 \) (blue), \( m = 0 \) (green), and \( m = 1 \) (red). The \( B \)-field dependence of the hybrid BLG QR energy spectrum is strikingly different from that of the isolated BLG QR. Here, for \( B = 0 \), the spectrum is continuous because of the presence of the infinite MLG, and with increasing magnetic field the degeneracy of the states is lifted. The discrete levels (reflecting the confined states) depend on the angular momentum and the \( B \)-field strength. The spectra for both valleys show anticrossings, which are due to the influence of the MLG dot and BLG QR interface [see the inset of Fig. 3(b)]. As a result of this interface, the symmetry condition \( E^K_\mu(m) = -E^{K'}_\mu(m) \) (which holds for an isolated ring) is no longer preserved here, as shown explicitly in Fig. 3(c). At high magnetic fields, i.e., \( B > B_{\mu} \), the energy levels merge into the LLs of pristine MLG \( E_n = \pm \sqrt{2n} \hbar v_F / l_B \) with \( n = 0, 1, \ldots \), shown by dashed black curves in Figs. 3(a) and 3(b) indicating that magnetic confinement dominates and the carriers become localized at the center of the dot region (see Fig. 4 and the corresponding discussion). It is also worth noting that, here, the states with \( m < 0 \) contribute to the zero LL (\( n = 0 \)) only in one valley (\( K \)). This is in contrast with the other graphene nanostructures such as MLG [27] and BLG QDs [61] with IM boundary condition in which the states from both valleys form the zero LL. Notice that, unlike the isolated BLG QR, it is not straightforward to calculate the conductance of the hybrid QR in the TBM, since its bottom layer consists of an infinite graphene layer sheet. However, in Sec. III B, we will compare explicitly the analytical energy levels with those obtained within the TBM for both ring structures.

Figure 3(c) shows the energy spectrum as a function of the angular momentum at a specific magnetic field \( (B = 10 \) T) for both valleys \( K \) (solid blue circles) and \( K' \) (open red circles). As seen, there is no symmetry between the e-h energies as well as between the valleys. For a given magnetic field, the energy levels at both valleys are affected by the ring interface for a particular range of \( m \) and converge to the MLG LLs for larger \( m \)'s (green lines), indicating carrier localization inside the dot region. Note that at the \( K \) (\( K' \)) valley, the maximum value of the angular momentum \( m_{\text{max}} \) that is converged to the \( n \)th electron LL of MLG is \( m_{\text{max}} = n - 1 \) \( (m_{\text{max}} = n, n \neq 0) \). Further, in both valleys, the electron states show a smooth oscillatory behavior as function of magnetic field and strong anticrossings are visible also for hole states. Similar behavior was also noted for the double MLG-BLG junctions studied in Ref. [78] when the spectra are plotted as a function of the center coordinate of the cyclotron orbit.

To better understand the behavior of the energy levels in the spectrum of the hybrid ring, we plot in Fig. 4 the energy levels of both types of rings and MLG dot for a specific angular momentum, \( m = -1 \) [panel 4(a)], as well as the corresponding spinor components \( [\phi_\mu(\rho), \nu = A, B, \ldots] \) and probability densities \( [\phi_A^2(\rho) + \phi_B^2(\rho), \ldots] \) for the representative energy states marked by (b), (c), and (d) in Fig. 4(a), [panels (b)-(d)]. As seen, the energy levels of the hybrid ring seem to be tunable from the MLG dot, to isolated BLG QR, and to MLG Landau energy levels as \( B \) increases. At the representative state (b), the
energy level (blue circles) resembles that of the MLG dot (dashed green curves), the corresponding quantum state is mostly confined inside the dot region with a substantial probability density in the ring region, see 4(b) panels. When the energy level approaches the one of the isolated ring (red curves), e.g., point (c), the carrier is mostly confined inside the ring region [Fig. 4(c)]. For strong magnetic field, e.g., point (d), the energy level converges to the MLG LL (magenta curve), the state is completely confined inside the dot region, as seen in Fig. 4(d).

It is also interesting to investigate the effect of ring width on the energy spectrum of both structures. For this purpose, we plot in Fig. 5, the lowest energy levels as a function of the width of the ring \( W = R_2 - R_1 \) for (a) isolated and (b) hybrid BLG QRs. In both cases, for the sake of clarity, the results are presented only for particular angular momentum \( m = -20 \) at \( B = 20 \) T. In the case of an isolated ring, the energy levels are separated by a gap, depending on the angular momentum and ring width, which closes when \( W \) increases and the energy levels for both valleys approach the LLs of bulk BLG as depicted by the horizontal gray lines. In the hybrid ring, Fig. 5(b), at small ring widths, the energy levels correspond to the LLs of MLG. When \( W \) increases, the electron (hole) energy levels, due to the influence of MLG-BLG interface, exhibit a flat plateau (strong anticrossing) features for certain ranges of \( W \), and eventually merge into the LLs of bulk BLG. The plateau-like and oscillatory features appearing in the energy spectrum can be understood as hybridization of the energy levels of the terminated systems, MLG QD and BLG antidot. Similar behavior was found for previous studies of MLG-BLG junctions [24, 72, 78, 79].

**B. Comparison with tight-binding model**

In order to check the validity of the continuum approximation, we compare explicitly our analytical results with the energy levels calculated within a nearest-neighbor TB approach. We include only the nearest-neighbor hopping parameters \( \gamma_0 = -2.7 \) eV and \( \gamma_1 = 0.4 \) eV as **intralayer** and **interlayer** couplings, respectively. The effect of an external magnetic field can be introduced into the calculations via the Peierls substitution \( t_{ij} \to t_{ij} e^{i \Phi_{ij}} \), where \( \Phi_{ij} = (1/\Phi_0) \int_{R_i}^{R_j} A(\mathbf{r}) \cdot d\mathbf{r} \) is the Peierls phase [85] with \( \Phi_0 = h/e \approx 4.14 \times 10^{-15} \) Wb the magnetic flux quan-
FIG. 6. (a) Schematic geometry of the circular BLG QRs with inner and outer radii of $R_1$ and $R_2$, respectively, defined by the IM boundary used in TBM calculation. We define the honeycomb lattice of BLG with the lattice vectors $a_1 = a_0(\sqrt{3}/2, 1/2)$ and $a_2 = a_0(\sqrt{3}/2, -1/2)$, where $a_0 = 0.246$ nm is the lattice constant. The coordinates $x$ and $y$ are defined along the armchair and zigzag directions, respectively (Fig. 1). The upper (lower) panel depicts the isolated (hybrid) BLG QR geometry. The atoms of the two layers are represented by blue (layer 1) and red (layer 2) circles. The ring regions are surrounded by a site-dependent staggered potential (yellow areas), where the atoms belonging to the sublattices $A_1$ ($A_2$) and $B_1$ ($B_2$) have mass-term potentials of $+M_0$ ($+M_0$) and $-M_0$ ($-M_0$), respectively. In the case of hybrid ring, to simulate the lower layer as an infinite graphene sheet, we consider it as a large circular flake (layer 1) on which a second MLG ring (layer 2) is sitting on top of it. Further, to eliminate the specific edge effects of layer 1, here also, we apply the staggered potential on the atoms located within the ribbon width of $\Delta r = 2a_0$ at the edge of the flake (yellow region). (b,c) Energy levels of (b) an isolated BLG QR and (c) a hybrid BLG QR as a function of the magnetic field $B$, calculated within the Dirac (red circles) and the TB (solid blue curves) models. For both cases, the results are presented for rings with inner and outer radii of $R_1 = 8a_0 = 1.97$ nm and $R_2 = 18a_0 = 4.43$ nm. The mass potential is $M_0 = 2$ eV. In the hybrid ring, we use a circular flake of radius $R_0 = 38a_0 = 9.35$ nm for the lower layer.

The vector potential corresponding to the external magnetic field $B = B\hat{z}$ perpendicular to the BLG flakes is chosen in the Landau gauge $A(r) = (0, Bx, 0)$ for which one finds that $\Phi_{ij}$ is only nonzero in the $y$ direction and is given by $\Phi_{ij} = \text{sign}(y_j - y_i) \frac{\pi + \pi_i}{2\sqrt{3}a_0} \Phi_0$, where $\Phi = B(\sqrt{3}a_0^2/2)$ is the magnetic flux threading one carbon hexagon ($a_0$ is the graphene lattice constant). Here, the QRs are defined by a staggered site-dependent potential such that the atoms belonging to the sublattices $A_1$ ($A_2$) and $B_1$ ($B_2$) have a mass-term potential of $+M_0$ ($+M_0$) and $-M_0$ ($-M_0$), respectively, as shown in Fig. 6(a). This simulates the substrate effect and can be regarded as the IM boundary condition in the TBM. More details of the geometries are provided in the caption of Fig. 6. Furthermore, our calculations show that in the Dirac model, adopting the same radius used to define the atomic geometries in TBM, results in the energy levels which are slightly larger (i.e., shifted up) than those obtained by TBM. Two possible explanations for this discrepancy can be considered. First, in the TBM to simulate the IM boundary, we have used a narrow ribbon of $\Delta r$ containing atoms with different finite on-site potentials in addition to the ring radii as shown in Fig. 6(a). The second which has also been proposed in Ref. [86], can be attributed to the $\pi$ electrons that in the TBM extend over the whole geometry. Accordingly, to compensate for this difference, we use a larger width for the rings in the Dirac model. Also note that, due to the computational limitation of the TBM, we have considered the small sizes of the rings and the applied magnetic field in our calculations is too large to be achievable in experiments. However, for the study of the electronic properties of graphene nanostructures in the presence of a perpendicular magnetic field, one can define a scaling factor and thus extend the results to lower magnetic field and larger sample sizes, e.g., see Refs. [12, 49, 87].

In Figs. 6(b) and 6(c), we compare the results obtained within the TB and Dirac approaches for the lowest-energy levels of both rings as a function of magnetic field $B$, see the figure caption for details. Notice that the Dirac results agree with the TB ones, especially for the lower-energy states. In the case of an isolated ring, there are some discrepancies between the two models for the hole energy levels. This is due to the fact that in the TBM, applying IM boundary at the edge boundary which is now not a perfect circle, breaks the e-h symmetry. This symmetry, however, is preserved in the Dirac model as a result of its perfect circular geometry. Moreover, in the
(a) Isolated BLG QR  
(b) Hybrid BLG QR

FIG. 7. Valley- and layer-resolved local density of states [LDOS (arbitrary units)] of (a) an isolated BLG QR and (b) a hybrid BLG QR with the inner and outer radii of \( R_1 = 30 \text{ nm} \) and \( R_2 = 40 \text{ nm} \), respectively. The magnetic field is \( B = 10 \text{ T} \). Layer 1 (2) is represented by the blue (red) curves. The solid (dashed) curves correspond to the \( K \) (\( K' \)) valley. For the isolated BLG QR, the LDOS is computed in the energy range of \([0,0.2] \text{ eV}\), while for the hybrid one, the states in the energy window of \([0,0.11] \text{ eV}\) [states between the lowest MLG LLs, \( n = 0 \) and \( n = 1 \) in Fig. 3(c)] are sampled.

TBM results of the hybrid ring [solid blue curves in Fig. 6(c)], we see a bunch of energy levels decreasing in energy and approaching the zero energy. They are known as the quantum Hall edge states [88] and are confined at the edges of the graphene flakes and are a result of the finite size of the flake. These Hall edge states are absent in the Dirac model for which the first layer is considered as an infinite graphene sheet. In general, we expect good agreement between the Dirac and TB models for the low-energy states. For the higher energy levels and high magnetic field, discrepancy between the Dirac and TB results becomes more significant. The reason is that the linear spectrum invoked in the Dirac equation is no longer valid.

We also notice that the energy spectra in both cases, exhibit periodic oscillations as the magnetic flux varies. This is a direct consequence of the AB effect [89] which is well known and has been investigated for QRs (metallic, semiconductor as well as graphene) [8, 90–93]. Similar to the AB effect in semiconductor QRs, the energy oscillations manifest itself in AB oscillations in the persistent current \( j(\Phi) = -\frac{\partial}{\partial \Phi} E \).

Finally, we consider the valley- and layer-resolved LDOS for the studied structures which can be probed by quantum capacitance measurements and by scanning tunneling microscopy. We consider the LDOS in the energy window of \( \Omega \equiv [E_1, E_2] \) which can be obtained using

\[
\rho^\ell(r) = \sum_{E_i \in \Omega} \delta(E - E_i)|\psi^E_i(r)|^2, \tag{21}
\]

where \( \psi^E_i(r) = [\phi^{E_i}_{A\ell}(r), \phi^{E_i}_{B\ell}(r)]^T \) denotes the quantum state of the two layers (\( \ell = 1,2 \)) with energy \( E_i \). The components \( \phi^E_{A\ell}(r) \) and \( \phi^E_{B\ell}(r) \) correspond, respectively, to the different sublattices \( A\ell \) and \( B\ell \) in each layer at the given valley (\( \tau = K, K' \)).

In Fig. 7, we show the valley- and layer-resolved LDOS of the lowest-electron-energy states for both ring structures at \( B = 10 \text{ T} \). In the case of isolated BLG QR [Fig. 7(a)], for which the LDOS is computed in the energy range of \([0,0.2] \text{ eV}\), one can see that the LDOS of layer 1 (blue curves) is mostly centered inside the ring whereas in layer 2 the LDOS has its maximum value at the inner side of the ring. This behavior happens for both valleys, \( K \) and \( K' \) (solid and dashed curves, respectively). Besides, in each layer, the maximum contribution in the LDOS belongs to the \( K' \) valley. The lower panel of Fig. 7(a) shows the LDOS for each valley separately. We see that the LDOS contribution of the \( K' \) valley at the inner boundary of the ring is dominant which decreases with radial distance and at the outer side of the ring, contributions of both valleys in the LDOS are almost equal. Shown in panel (b) of Fig. 7 is the LDOS of the hybrid BLG QR. In this case, we compute the LDOS over the electron-energy states located between the lowest MLG LLs, \( n = 0 \) and \( n = 1 \). Here too, the contribution of the \( K' \) valley in the LDOS is dominant in both layers. While the LDOS of the lower layer (blue curves) is centered in the middle of the ring, in the upper layer (red curves) the LDOS form rings with maximaums at the inner and outer boundaries of the ring. Thus the difference between the valley contributions as well as from the different layers in the LDOS at a given magnetic field can open up novel ways for applications of such graphene-based nanostructures in valleytronics.

IV. SUMMARY AND CONCLUDING REMARKS

In summary, based on continuum (Dirac-Weyl equation) and tight-binding models, we studied the electronic properties of BLG QR, defined by IM potential, in two different configurations: an isolated BLG QR and a hybrid one where a MLG ring is put on top of an infinite MLG sheet. By applying the Dirac approximation and the IM boundary condition, we first obtained analytical results for the energy levels and corresponding wave spinors for both structures as function of a perpendicular magnetic field.

In contrast to the previously investigated zero-width BLG QR, here, the isolated BLG QR features a sizeable and magnetically tunable band gap that decreases as the magnetic field strength is increased. Our analytical findings are in excellent agreement with the tight-binding transport simulations. Further, the theoretical results show the intervalley symmetry \( E^K_c(m) = -E^K_h(m) \) between the electron and hole states for the energy levels of the isolated BLG QR, where \( m \) is the angular momen-
tum quantum number and $K, K'$ refer to the two Dirac valleys.

The results for hybrid BLG QRs showed that the presence of an interface boundary in a hybrid BLG QR modifies drastically the energy levels as compared to that of an isolated BLG QR and its energy levels interplay between the MLG dot, isolated BLG QR, and MLG Landau energy levels as magnetic field varies. No symmetry between the energy levels is found in this case.

Further, the energy spectrum of both structures exhibits Aharonov-Bohm oscillations as the magnetic field varies. We also explicitly confirmed the validity of our results by simulating the QRs by a staggered site-dependent potential in the TBM. We found good agreement between our analytical results obtained in the continuum approximation and those calculated within the TBM. Finally, we analyzed the spatial dependence of the valley- and layer-resolved LDOS for the proposed BLG TBM. Finally, we analyzed the spatial dependence of the valley- and layer-resolved LDOS for the proposed BLG TBM. Further, the energy spectrum of both structures exhibits Aharonov-Bohm oscillations as the magnetic field varies. No symmetry between the energy levels is found in this case.

Although many-body effects, such as those coming from electron-electron and electron-phonon interactions, may appear in some experiments in certain regimes and initial conditions, thus affecting the confinement properties in BLG-based nanostructures \cite{18, 94, 95}, the analytical solution proposed here allows us to have physical insights into the basic mechanisms behind the results, which is of fundamental importance for a theoretical understanding of some electronic properties in BLG nanostructures. Moreover, recent experimental measurements of quantum confined states in BLG QDs by using scanning tunneling microscope \cite{96–101} have been confirmed by single-particle tight-binding calculations, even in the presence of charge defects, impurities, dopants and adatoms \cite{98, 100, 101}.

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