Graphene kirigami as a platform for stretchable and tunable quantum dot arrays

D. A. Bahamon,1,* Zenan Qi,2 Harold S. Park,2 Vitor M. Pereira,3,4 and David K. Campbell5

1MackGraphe—Graphene and Nano-Materials Research Center, Mackenzie Presbyterian University, Rua da Consolação 896, 01302-907, São Paulo, SP, Brazil
2Department of Mechanical Engineering, Boston University, Boston, Massachusetts 02215, USA
3Centre for Advanced 2D Materials & Graphene Research Centre, National University of Singapore, 6 Science Drive 2, Singapore 117546
4Department of Physics, National University of Singapore, 2 Science Drive 3, Singapore 117542
5Department of Physics, Boston University, 590 Commonwealth Avenue, Boston, Massachusetts 02215, USA

(Received 17 March 2016; revised manuscript received 16 May 2016; published 6 June 2016)

The quantum transport properties of a graphene kirigami similar to those studied in recent experiments are calculated in the regime of elastic, reversible deformations. Our results show that, at low electronic densities, the conductance profile of such structures replicates that of a system of coupled quantum dots, characterized by a sequence of minibands and stopgaps. The conductance and $I$-$V$ curves have different characteristics in the distinct stages of deformation that characterize the elongation of these structures. Notably, the effective coupling between localized states is strongly reduced in the small elongation stage but revived at large elongations that allow the reestablishment of resonant tunneling across the kirigami. This provides an interesting example of interplay between geometry, strain, spatial confinement, and electronic transport. The alternating miniband and stopgap structure in the transmission leads to $I$-$V$ characteristics with negative differential conductance in well defined energy/doping ranges. These effects should be stable in a realistic scenario that includes edge roughness and Coulomb interactions, as these are expected to further promote localization of states at low energies in narrow segments of graphene nanostructures.

DOI: 10.1103/PhysRevB.93.235408

The development of advanced microfabrication techniques in the final two decades of the last century allowed the creation of two-dimensional electron gases (2DEGs) at the gate oxide-semiconductor interface of heterostructures [1]. In the final two decades of the last century allowed the development of advanced microfabrication techniques in the gate oxide-semiconductor interface of heterostructures [1]. In these structures, point contacts [2,3], quantum wires [4], constrictions [5], and quantum dots [6] have been used to confine electrons in quantum dots [10–13], nanoribbons [14,15], andjunctions [16].

An interesting and different direction has recently been pursued by employing lithography to extend the range of elastic deformation that can be sustained by a graphene by patterning nanomeshes out of large graphene sheets [17]. Such structures, inspired by the Japanese art of cutting paper called kirigami, have been designed and tested in recent experiments [18] that establish their mechanical robustness and extremely high elongation limits (~240%) compared to uncut graphene. In a previous study of the elastic and mechanical characteristics of graphene kirigami, some of us showed that their stretchability limit and effective Young’s modulus can be characterized (and customized) in terms of two geometric parameters [19]: $\alpha$, the ratio of the overlapping cut length to the kirigami length, and $\beta$, the ratio of the overlapping width to the kirigami length. With reference to Fig. 1(a), these parameters are $\alpha = (w - 0.5b)/L$ and $\beta = (0.5d - c)/L$.

The ability currently demonstrated to experimentally design graphene kirigami capable of ultrahigh elastic deformations raises the question of how the electronic states and the flow of current within the kirigami are, or can be, modified under deformation. In particular, since any kirigami always involves a number of bends, indentations, and narrow regions (henceforth, “constrictions”), it can naturally harbor a number of localized states at low energies (below the threshold for electronic transmission) as a result of either the geometry alone [20] or geometry combined with disorder [21] and/or Coulomb interactions [22,23]. The experimental observation of Coulomb blockade in graphene nanoribbons and constrictions [22–26] indicates that such states are expected to be prevalent at low energies in “papercut” graphene devices and that disorder can promote or further stabilize them.

* dario.bahamon@mackenzie.br
A segment of suitably patterned graphene hosting such states defines a quantum dot, and its periodic repetition would define an array of coupled quantum dots, analogously to what has been achieved in semiconductor structures through comb-shaped arrays of side gates [7,28,29]. The fact that graphene kirigami of possibly any desired shape can sustain repeated stretching cycles [18] suggests that one might be able to design kirigami suitable for the study of coupled quantum dots, with the advantage that the interdot coupling responds directly to the deformed state of the structure. The vast range of stretchability limits and effective Young moduli that these structures can be designed with, combined with the proven mechanical resilience of graphene at the nanoscale, hint at the possibility of designing versatile electromechanical devices based on this concept of stretchable graphene quantum dot arrays whose current response can be strongly sensitive to the interdot coupling.

That is precisely the problem we address in this work, concentrating on the local atomic aspects that contribute to localization of electronic states at certain portions of the kirigami. We confirm that such states define effective quantum dots, which result in a characteristic profile of the conductance at low energies defined by a sequence of resonant minibands and stopgaps; see Fig. 3(a). This is analogous to the conductance of periodic split-gate semiconductor heterostructures [7,28,30], where each miniband consists of a set of resonant tunneling peaks. At low energies, the conductance profile is seen to be very sensitive to the deformation, and we analyze this response in terms of the interplay between variations in interdot coupling and strain barriers that develop under stretching.

Our study combines information on the local atomic displacements obtained from molecular dynamics (MD) simulations of deformed kirigami with quantum transport calculations to assess the conductance characteristics at different stages of deformation [31]. At the structural level, the initial stage (stage 1) of the end-to-end longitudinal deformation is characterized by a very small effective Young’s modulus and an essentially negligible average stretching of the interatomic bonds because, in this early stage, elongation occurs mostly through bending and twisting of the structure in three-dimensional space [19]. Despite the low in-plane stretching that occurs during this stage, the conductance and the current are significantly reduced, and the resonant features disappear. However, further elongation into a second stage of deformation where there is substantial bond stretching leads to the revival of the resonant features, and the restoration of the magnitudes of conductance to the same levels observed in the undistorted kirigami. These changes are consistently understood as a consequence of transport at low energies being dominated by resonant tunneling between states localized at specific portions of each periodic unit of the kirigami; elongation during stage 1 of deformation perturbs the coupling (overlap) between these states and considerably degrades the conductance as a result of hopping disorder that weakens the superperiodicity of the kirigami structure; further stretching into stage 2 results in strong and localized strain barriers that reestablish the superperiodicity, favor electron localization, and hence revive resonant transmission. The terminology “strain barrier” is employed here deliberately to emphasize the direct impact that deformation hot spots have on the electronic behavior of the system at high elongations: as discussed below, the development of localized and periodic regions with extreme bond stretching reinforces the geometry-induced confinement of electronic states within each repeating unit of the kirigami.

\section{I. METHODOLOGY}

Our representative kirigami is obtained by cutting out a graphene nanoribbon of width \( w \) according to the pattern shown in Fig. 1(a). The rectangular interior holes are defined by the height \( w \) and width \( c \), while the connecting necks have length \( d \) and a cut starting from the outer edge of length 0.5\( w \). The longitudinal period is \( d \) and \( L \) defines the total length of the system. We shall concentrate our discussion in the kirigami shown schematically in Fig. 1(b); our actual system contains 11,408 atoms with the geometrical parameters \( b \approx 10 \text{ nm}, w \approx 6.8 \text{ nm}, c \approx 0.7 \text{ nm}, d = 4.8 \text{ nm}, \text{ and } L \approx 34 \text{ nm} \). For definiteness, we base our discussion on this specific structure where the graphene lattice is oriented so that all horizontal edges are zigzag edges. We also do not include any disorder or edge roughness at this stage. The results, however, are general and should hold when these two restrictions are relaxed because the only key physical ingredient is the existence of localized states defining a local quantum dot at specific constrictions or bends, and these can be stabilized by different geometries, with or without disorder [21]. For illustration, we show that explicitly by analyzing a different lattice orientation in the discussion session.

This study is divided in three stages that aim to determine the electronic transport properties of the kirigami under
longitudinal tension. The deformed structures are first obtained with molecular dynamics (MD) simulations of the finite kirigami. We used the Sandia open-source code LAMMPS [32,33] with the AIREBO [34–36] potential to describe the C-C interactions: the cutoff radii are 2 Å for the REBO term and 6.8 Å for the Lennard-Jones term in the AIREBO potential. The explicit position of each carbon atom in the deformed structure is then used to build a π-band tight-binding Hamiltonian, \( H = \sum_{ij} t_{ij} (c_i^\dagger c_j + c_j^\dagger c_i) \), for the distorted kirigami. In this Hamiltonian, \( c_i \) denotes the annihilation operator on site \( i \) and \( t_{ij} \) represents the hopping amplitude between nearest-neighbor sites \( i \) and \( j \). The stretching, the compression, and the rotation of the C-C bonds created by the tensile load are taken into account in this effective Hamiltonian between nearest-neighbor sites \( i \) and \( j \) of Green’s functions, the conductance in the Landauer-Büttiker formalism can be written as \([39–41]\)

\[
G = \frac{2e^2}{h} T(E) = \frac{2e^2}{h} \text{Tr} [\Gamma_L G^\dagger \Gamma_L G^a],
\]

where \( G^a \) is the total current as a function of the applied bias voltage \( V \), are calculated from the transmission function \( T(E,V) \) as \([40]\)

\[
I(V) = \frac{2e}{h} \int_{-\infty}^{+\infty} T(E,V)(f_L(E) - f_R(E))dE,
\]

where \( f_L(R)(E) \) is the Fermi distribution of the left (right) contact.

II. CONDUCTANCE UNDER DEFORMATION

For a representative illustration showing the behavior in the two stages of deformation, we chose the kirigami structure shown in Fig. 1(b) that has a small number of cuts per repeating unit and a very high stretchability, up to 65%. In order to facilitate the presence of localized states within each segment of the kirigami even in the absence of any deformation, we chose to orient the underlying graphene lattice so that the longitudinal cuts are along a zigzag direction, as the internal mini-zigzag edges so defined are expected to naturally support localized states \([42]\) (as pointed out above, this is not a limitation).

A kirigami such as this one has four deformation stages \([19]\): (i) elongation with bending and twisting, but very little in-plane stress, (ii) elongation with stress, (iii) yielding, and (iv) fracture. We restrict our analysis to the first two, where deformations are elastic and reversible, and whose stress-strain characteristic is shown in Fig. 2. During the first stage (in this particular structure that corresponds to total deformations below \( \approx 20\% \)) horizontal and vertical segments twist and rotate and, as a result of this excursion of the graphene sheet into the third dimension, the kirigami elongates without significant modification of the average C-C bond length, except for very localized strain hot spots at the corners of the connecting elements \([19]\). A representative sample of a kirigami in this stage is shown in Fig. 1(c) for a total deformation of 15.5%. With further increase in the tensile load, the kirigami is not capable of accommodating higher elongations only by twisting, and this triggers the onset of stage 2 (here in the range 20%–40%), where further deformation occurs through

![FIG. 2. Stress-strain curve for the kirigami, showing the first two stages corresponding to the elastic and reversible deformations.](image-url)
strectching of the carbon bonds. One important consequence of the existence of these two regimes is that the effective Young’s modulus is much lower in stage 1, \(E \approx 0.69\) N/m (where it is essentially determined by the very small bending stiffness of monolayer graphene), than in stage 2, \(E \approx 15.08\) N/m. This makes mechanical manipulation of kirigami structures experimentally possible and easy in stage 1 [18].

We followed the evolution of conductance at low energies for different strain values within the reversible and elastic region and summarize the results in Fig. 3(a) for deformations of 0%, 5.3%, 15.5%, 25.1%, and 34.7%. The conductance profile of the reference (undeformed) structure displays resonant transmission within well defined bands (minibands) of width \(\approx 0.05\) \(t_0\). We can see in the DOS plotted in Fig. 3(b) for the undeformed structure that each miniband arises from the clustering of a finite number of states. At those energies transmission occurs through resonant tunneling, and is entirely suppressed otherwise, strikingly different from the conductance profile expected for a graphene nanoribbon [20,30]. In an infinite kirigami the stopgaps that separate the minibands arise from the folding of the 1D Brillouin zone of the ideal (pre-cut) graphene ribbon as a result of the new superlattice defined by the kirigami, with the expected opening of spectral gaps at the edge of the reduced zone. In our finite kirigami, Fig. 3(b) shows that the DOS is likewise strongly suppressed at the miniband edges but does not reach zero in the stopgaps. This contrasts with the sharp changes in the conductance [Fig. 3(a)]: finite (resonant) within a miniband, and clearly zero in the stopgaps. This is a clear sign that states contributing to the DOS in the stopgap regions are spatially localized, and unable to hybridize to form a tunneling pathway that spans the entire length of the system. Therefore, the robust stopgap structure in the transmission is not a result of spectral characteristics of the kirigami alone.

To be specific, let us analyze the close-up plot shown in Fig. 3(c) that captures the energy interval \(0.03 < E / t_0 < 0.09\). A mapping of the LDOS at any of the resonances shown in this panel confirms that the electronic density associated with a resonance is strongly localized around the internal longitudinal strips. Figure 4(a) shows a representative example of the “LDOS hot spots” (in red) appearing at those regions, and a good overlap/coupling with the external contacts. Another snapshot of the LDOS at a different transmission resonance is shown in Fig. 5. The existence of the minibands of resonant transmission is therefore due to the presence of localized states within each unit of the kirigami, which are localized both transversely and longitudinally by the combined effect of the internal zigzag edges and the finite extent of each segment of the kirigami [7,20,28,30,43–45]. At the lowest energies, transmission across the entire system is assisted by tunneling through these localized states that, hence, play a role similar to that of an array of coupled quantum dots that can enable resonant transmission across the entire system at (and only at) well defined energies. The width of each miniband is determined by the overlap, \(\gamma\), between localized states, and the number of resonant peaks in a miniband, \(N\), counts the number of isolated quantum dots [29,30]. Our geometry contains the 11 internal channels labeled 1–11 in Fig. 1(b) that define 11 effective quantum dots. Correspondingly, each transmission miniband has a fine structure consisting of 11 peaks as can be seen in Fig. 3(c) (the outermost two directly in contact with the metallic leads do not define quantum dots because backscattered electrons at these sections are completely absorbed by the contacts).

A quantitative estimate of the interdot overlap can be obtained assuming a one-dimensional tight-binding model, according to which the energy levels within one miniband should appear at positions

\[ E_n = -2\gamma \cos \left( \frac{n\pi}{N+1} \right), \quad n = 1, 2, \ldots, N, \]

relative to the center of the miniband. The width of a miniband is given by \(\Delta E = E_N - E_1\). The data in Fig. 3(c) allow us to estimate \(\Delta E \approx 0.04t_0\) for the undeformed kirigami, which corresponds to \(\gamma \approx 0.01t_0\). Recalling that \(t_0\) is the hopping between nearest-neighbor carbon atoms in undeformed graphene [Eq. (2)], this shows an interdot overlap two orders of magnitude smaller than the hopping in the underlying graphene lattice. Such a small value of \(\gamma\) is natural given the large spatial separation \(\sim d / 2 = 2.4\) nm; cf. Fig. 1(a) between each pair of hybridized localized states that we see in Fig. 5.

Under tensile load in stage 1, the kirigami elongates and becomes distorted, which significantly perturbs the overlap between the localized states. That is the conclusion that follows from the progressive disappearance of the resonant minibands under deformation that we can see in Fig. 3(a) for deformations of 5.3% and 15.5%. It is notable that, even though in stage 1 there is very little change in the average C-C bond length, the regions that are most affected by that, and which are more
FIG. 4. Normalized LDOS for the kirigami (a) undeformed at $E = 0.067 t_0$, (b) deformed by 15.5% at $E = 0.074 t_0$, and (c) deformed by 34.7% at $E = 0.073 t_0$.

strongly bent, are those in the vicinity of the localized states of the undeformed kirigami. It is, therefore, not surprising that the conductance assisted by resonant tunneling through these states can be significantly modified in this stage since the perturbations to the hopping can be significant in precisely the regions more critical for the overlap between neighboring localized states. For an elongation of 15.5%, Fig. 3(a) shows that the miniband structure disappears and the system behaves largely as an insulator through most of the energy range shown. The LDOS associated with the few remaining weak conductance peaks reflects the less effective coupling between the localized states [see Fig. 4(a)]. In the analogy introduced above, in this stage of deformation we have the equivalent of a device consisting of distinct and asymmetrically coupled quantum dots, for which the energy of transmission resonances is predominantly determined by the energy levels and coupling of individual dots [46,47].

A simple extrapolation of this picture would forecast a progressively more substantial degradation of the conductance with further stretching. However, inspection of Fig. 3(a) reveals otherwise: resonant transmission is revived beyond a geometry-controllable threshold when the system enters stage 2 of deformation. At 25.1% strain, resonant transmission is recovered at particular energies within the lowest miniband, and further stretching to 34.7% strain, rather than degrading, further improves this situation with resonant peaks reappearing again throughout the entire lowest miniband as in the undeformed device. As expected, the width of each miniband is now smaller than in the undeformed state, a combined effect of the increase in interdot distance, and the decrease of electronic hoppings in the carbon lattice. Figure 4(c) exhibits the LDOS for one of the revived conductance peaks of the same system that shows a strong similarity with the undeformed counterpart in Fig. 4(a).

It is clear from both the conductance and the LDOS that deforming the system well inside stage 2 restores the coupling between the localized states trapped at the internal mini-edges, that is strongly affected during stage 1. As pointed out earlier, the major difference between stages 1 and 2 is that, in the latter, further elongation proceeds through deformation of the C-C bonds, and therein lies the mechanism that promotes the enhanced coupling and the revival of resonant transmission. As one intuitively anticipates, and can be explicitly seen in Fig. 6(b), the deformations in question are borne almost
FIG. 6. Normalized C-C bond length for the kirigami deformed by 15.5% [panel (a), stage 1] and 34.7% [panel (b), stage 2], where the normalization is by the undeformed C-C bond length.

entirely by the bonds along the segments that link neighboring units of the kirigami. A comparison of the bond length distribution in stage 1 [Fig. 6(a)] versus stage 2 [Fig. 6(b)] indeed shows that the first regime is characterized by small and seemingly random variations, whereas the latter features a clear and periodic pattern of deformation hot spots where the C-C bond is stretched in excess of 20%.

Variations in bond length imply perturbations to the electronic hopping \[ t_{ij}, \text{ Eq. (1)} \], which is the quantity that directly affects the transport. However, it is important to mention that the C-C bond map excludes bending contributions that should dominate the hopping modifications in stage 1. To clarify this point, we plot in Fig. 7 the corresponding hopping distribution maps. The variations are of the order of \( \sim \pm 10\% \) and \( \sim \pm 20\% \) at 15.5% and 34.7% elongation, respectively. More than the magnitude, the key difference lies in the sign and spatial pattern of these hopping variations: in the first case, they appear randomly distributed and one sees places of both enhanced and reduced hopping compared to the undeformed structure; in the second, the pattern is clearly periodic and dominated by reduced hopping (\( \sim 20\% \)) spatially correlated with the strain hot spots of Fig. 6(b). These electronic “weak links” reinforce the superperiodicity of the structure, while simultaneously contributing to localize further the electronic states within each segment, thereby promoting the revival of the miniband and stopgap structure in the conductance.

In summary, the profile of the conductance at low energies in these graphene structures is typical of a system of weakly coupled quantum dots: minibands of resonant transmission alternating with stopgaps due to the superlattice periodicity along the longitudinal direction. Direct inspection of the LDOS at any of the resonant energies provides a direct confirmation of this in real space. Analysis of the evolution of the conductance and the LDOS with deformation shows that the resonant tunneling assisted by localized states is strongly suppressed during stage 1 of deformation. The local perturbations to the electronic hopping in this regime are sufficient to disturb the overlap between the localized states. This effect is reversed in stage 2 by the appearance of strong strain barriers spanning the length of the system along the links between individual elements of the periodic structure. These have a confining nature from the electronic point of view, which consistently explains the reinforcement of overlap between localized states, and the revival of resonant transmission, at the highest deformations.
current is expressed in units of $a_f e d$.

We used a bias voltage of $V_{SD} = 100$ meV, the current is expressed in units of $I_0 = 2e/h$, and the gate voltage in units of $t_0$.

**III. I-V CHARACTERISTICS AND NEGATIVE DIFFERENTIAL RESISTANCE**

In principle, to fully examine the current-voltage characteristics of these kirigami we should calculate the transmission function for different bias and gate voltages, according to Eq. (5). However, similar to the experimental setup used in Ref. [18] where the kirigami was immersed in an electro-chemically controlled liquid gate with a fixed bias voltage $V_{SD} = 100$ meV, we restrict our analysis to the electric current as a function of gate voltage for a fixed bias. Figure 8 displays current for kirigami deformed by 0%, 15.5%, and 34.7% under a fixed $V_{SD} = 100$ meV and at $T = 300$ K.

One can see a clear oscillation of the current with increasing $V_g$, a behavior that traces back to the underlying miniband structure of the conductance discussed in the context of Fig. 3. The fact that below $V_g = 0.1t_0$ the current magnitude first decreases with deformation, but then increases again in stage 2, is a natural consequence of the revival of resonant transmission at low energies promoted by the localized strain that emerges in stage 2. The most interesting aspect of the I-V characteristics shown in Fig. 8 is that the current oscillation immediately implies that these kirigami have negative differential resistance over reasonably large intervals of $V_g$, which is a much sought-after property for new classes of nonlinear electronic devices [48,49].

**IV. INTRODUCING DEPHASING**

So far, our electronic transport model considers coherent transport, including only edges and mechanical deformations as source of scattering. To address the robustness of the strain modulation of the conductance in a more realistic scenario, we now turn to effects of random scatterers modeled within the dephasing Büttiker-probe model [50]. To achieve that, we distribute voltage probes as phase-breaking scatterers all over the kirigami (except the left and right edges), and set the current at each probe to zero. Given that we are not interested in the particular physical mechanism (impurities, phonons, or electrons) behind the destruction of quantum coherence, we fix the self-energy of each Büttiker probe to $\Sigma_\phi = -i\eta$ [51], which is related to the phase relaxation time by $\tau_\phi = h/2\Gamma_\phi = h/4\eta$ [40,52]. In Fig. 9, we plot the effective conductance that obtains for weak ($\eta = 0.01t_0, \tau_\phi \approx 6.1 \times 10^{-15}$ s) and strong ($\eta = 0.5t_0, \tau_\phi \approx 1.2 \times 10^{-16}$ s) phase-breaking scatterers for different deformations. The dashed blue line corresponds to the conductance of a pristine zigzag nanoribbon with the same width of our contacts. Be aware of the different scales in the vertical axes.

![Graph showing current vs. gate voltage for kirigami deformed by 0%, 15.5%, and 34.7%](image1)

**FIG. 8.** Current vs gate voltage for the kirigami deformed by 0%, 15.5%, and 34.7%. We used a bias voltage of $V_{SD} = 100$ meV, the current is expressed in units of $I_0 = 2e/h$, and the gate voltage in units of $t_0$.

![Graphs showing effective conductance for weak and strong dephasing](image2)

**FIG. 9.** Effective conductance in the presence of (a) weak ($\eta = 0.01t_0, \tau_\phi \approx 6.1 \times 10^{-15}$ s) and (b) strong ($\eta = 0.5t_0, \tau_\phi \approx 1.2 \times 10^{-16}$ s) phase-breaking scatterers for different deformations. The dashed blue line corresponds to the conductance of a pristine zigzag nanoribbon with the same width of our contacts. Be aware of the different scales in the vertical axes.
energy. Measurements of the phase-relaxation length at low temperature in graphene have found $\ell_\phi \approx 1 \mu m$ [53], which is much smaller than the size of the typical internal lengths used in the kirigami experimentally probed by Blees and collaborators [18]. This might partially explain why deformed kirigami used in that experiment did not show noticeable sensitivity of the conductance to the elongation.

V. FINAL REMARKS

Our study shows that, when the typical feature sizes of a graphene-based kirigami are in the nanoscale, the electric conductance at low energies might be governed by resonant tunneling through states that are localized by the specific local geometry of each repeating element of the kirigami. The longitudinal periodicity of the structure results in an efficient coupling (strong overlap) between these states in the undeformed configuration. This state of affairs is, however, strongly sensitive to stage 1 deformations, and the conductance easily degrades in that regime where the overall elongation is a result of the structural twisting and bending, rather than extensive stretching of the C-C bonds. The electronic overlap is reinforced for stage 2 deformations as a result of the confining nature of the localized strain barriers that set in during this stage, and resonant transmission is hence revived at high overall elongations. The regime of resonant tunneling exists within well defined energy minibands isolated from each other by sizable transmission stopgaps. Their existence results in a strong oscillation of the $I$-$V$ characteristic as a function of gate voltage, and opens the possibility of driving the system from a conventional resistive regime to one of negative differential resistance, by simple electrostatic gating.

To explain the evolution of the conductance profile with elongation we analyzed directly the bond stretching and the perturbations to the nearest-neighbor hopping. It is customary in discussions of strained graphene to introduce the concept of pseudomagnetic field (PMF) [54–56] and map strain fields to PMFs in order to, for example, obtain a semiclassical intuition about how certain strain patterns disturb the motion of electrons [57]. In the present case, however, the characteristic dimensions of our device are small and, in addition, the strain distribution displays sharp variations within these small scales. This restricts the usefulness of the PMF concept and any semiclassical picture to interpret the effects of strain. We therefore focused directly on the changes affecting the nearest-neighbor hopping amplitudes since these are the quantities that more directly and fundamentally determine the conductance.

Finally, in our main discussion we focused on a kirigami whose horizontal segments consisted of zigzag strips of graphene. Not surprisingly, the mini-zigzag edges parallel to the transverse direction play an important role in stabilizing the localized states in the undeformed structure that are key for the conductance profile at low energy. It is important to reiterate that this is not a limitation, for the key physical ingredient is the existence of localized states defining local quantum dots, and the interplay between the strain-induced changes of the C-C hopping and effective interdot coupling upon mechanical stretching. To be specific, in Fig. 10(a) we show that rotating the underlying lattice by $90^\circ$ in the same kirigami leads to a similar profile of miniband and stopgap low-energy conductance, and that it is equally sensitive to deformation: for example, at about 16% elongation the miniband structure is entirely suppressed. In a future study we shall undertake the impact of edge roughness. We can, however, anticipate that edge roughness is expected to improve the scenario we describe here because this type of disorder promotes further localization of low-energy states in narrow graphene structures [22,23], and that explains the experimentally observed gaps and Coulomb blockade in rough (lithographically patterned) graphene nanoribbons [22,24–26]. We therefore predict that, in a realistic scenario, individual segments of a kirigami can behave as true quantum dots with a coupling (and, consequently, an overall transmission) amenable to modulation through the same type of deformation discussed here. The conditions for such behavior are expected to be very encompassing, depending only on choosing appropriate combinations of scales and geometries capable of hosting localized quantum-dot states, supported either by the geometry, disorder, or interactions.

ACKNOWLEDGMENTS

D.A.B. acknowledges support from FAPESP Grant No. 2012/50259-8. Z.Q. acknowledges the support of the Mechanical Engineering and Physics Departments at Boston University. V.M.P. acknowledges the support of the National Research Foundation (Singapore) through the CRP grant “Novel 2D materials with tailored properties: beyond graphene” (R-144-000-295-281).
