Negative quantum capacitance in graphene nanoribbons with lateral gates

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We present numerical simulations of the capacitive coupling between graphene nanoribbons of various widths and gate electrodes in different configurations. We compare the influence of lateral metallic or graphene side gate structures on the overall back gate capacitive coupling. Most interestingly, we find a complex interplay between quantum capacitance effects in the graphene nanoribbon and the lateral graphene side gates, giving rise to an unconventional negative quantum capacitance. The emerging non-linear capacitive couplings are investigated in detail. The experimentally relevant relative lever arm, the ratio between the coupling of the different gate structures, is discussed.

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I. INTRODUCTION

Graphene nanoribbons1–7 are of increasing interest due to their promise of a band gap, overcoming the gapless band structure of truly two-dimensional (2D) graphene.8,9 In particular, their overall semiconducting behavior allows the fabrication of graphene field-effect transistors,10 tunneling barriers,11 and quantum devices.12 First experimental demonstrations of graphene nanoribbon based quantum dots,13,14 double quantum dots15 and charge sensors16,17 have been reported in recent years. In most of these quantum devices the local electrostatic tunability of the electrochemical potential along graphene nanoribbons is key for the device functionality. For this purpose, local top-gates and lateral gates, based either on metals or in-plane graphene have been fabricated (see, e.g., Refs. 18–21). In particular, the 2D nature of graphene makes it straightforward to pattern a number of lateral graphene gates and in-plane charge detectors from the very same graphene sheet as the adjacent top-down fabricated nanoribbon.12,22 Consequently, a better understanding of the capacitive coupling between nanoribbons (with different widths) and (graphene) gate electrodes is important for device optimization and future graphene-based nanoelectronics. This is particularly true for graphene nanodevices since their Fermi energy is tuned using capacitive coupling. The strength of these couplings, in turn, depends on the density of states (DOS) via quantum capacitance effects.23,24 The low DOS close to the Dirac point thus makes graphene a rather unusual gate material with reduced and energy dependent screening properties.25,26 Graphene sheets have been subject to a large number of theoretical and experimental studies of quantum capacitance effects.27–37 Indeed, recent advances allowed experiments based on capacitance measurements to observe such phenomena as Fermi velocity renormalization,38 fractional quantum Hall phase transitions,39 and Hofstadter’s butterfly.40 Furthermore, quantum Hall transport has been used to probe the capacitance profile at graphene edges.41 These demonstrations give vested hope that quantum capacitance effects in graphene nanodevices could be exploited in future applications.

Here we present numerical simulations of the electrostatic capacitive coupling between graphene nanoribbons of different widths and gates in different configurations. In particular we elucidate the influence of metallic and graphene lateral side gate structures (see Fig. 1) on the back gate capacitive coupling. We show that the non-linear capacitive couplings give rise to complex interplay between quantum capacitance effects in the nanoribbon and the gates. Surprisingly, we find the unconventional phenomenon of an effective negative quantum capacitance in nanoribbons with lateral graphene gates. Inclusion of quantum capacitance effects will therefore be essential in reliably interpreting experimental results.

The paper is organized as follows: in Sec. II we briefly introduce the description of quantum capacitance effects on the Thomas–Fermi level and the model system under
investigation. Results for graphene nanoribbons with different width, and with metallic and graphene side gates are discussed in Sec. III. Finally, we compare the capacitive coupling between either nanoribbon and back gate or nanoribbon and side gates. We conclude with a short summary.

II. QUANTUM CAPACITANCE AND MODEL SYSTEM

Accurate simulations of the influence of side gates require a detailed treatment of the electrostatic problem beyond analytical models available for simple geometries such as the infinitely extended graphene sheet or an isolated nanoribbon.\textsuperscript{34,42} We therefore present a self-contained treatment of the quantum capacitance effect, and present a numerical approach to calculate quantum capacitance effects by coupling the Poisson equation with a Thomas–Fermi approach for the electronic structure. As a first test, we apply our formalism to analytically solvable models.\textsuperscript{34}

Consider the electrostatic problem of the capacitive coupling between two conductors at different voltages, separated by a medium with relative permittivity \(\varepsilon(r)\). The resulting electrostatic potential \(\Phi(r)\) follows the Poisson equation

\[
\nabla \cdot (\varepsilon(r)\nabla \Phi(r)) = -\frac{\rho_i(r)}{\varepsilon_0}, \quad (2.1)
\]

where \(\rho_i(r)\) is the free charge density and \(\varepsilon_0\) is the vacuum permittivity. A given potential difference \(d\Phi\) between the conductors will lead to a charge accumulation \(dQ\) on one conductor \(C\) (and \(-dQ\) on the other),

\[
dQ = \int_C d\rho_i(r) dr, \quad (2.2)
\]

according to Eq. (2.1). The classical capacitance \(C_{cl}\) then gives the ratio between charge and potential difference,

\[
C_{cl} = \frac{dQ}{d\Phi}. \quad (2.3)
\]

For nanoscale devices, an additional contribution to the capacitance can arise due to the electronic structure near the Fermi energy of the conductor. This quantum capacitance contribution is related to the additional energy cost for adding electrons to the conductor, which increases if the DOS near the Fermi edge decreases. Graphene with its vanishing DOS at the Dirac point is a prototypical case.

Within a Thomas–Fermi approach the local electron density \(n(r)\) related to the net free charge density through \(n(r) = \rho_i(r)/e\) (with \(e\) the absolute value of the elementary charge), is given by

\[
n(r) = \int_0^{E_F(r)} D(E; r) dE, \quad (2.4)
\]

with \(D(E; r)\) the local density of states and \(E_F(r)\) the local Fermi energy. In line with the semiclassical limit underlying the Thomas–Fermi approximation we consider variations only over length scales large compared to the de Broglie wavelength of the electrons. Accordingly, effects such as size quantization features\textsuperscript{43-45} of the graphene nanostructures are neglected in Eq. (2.4). They could be incorporated, e.g., through a fully self-consistent solution of the Poisson equation and the mean-field Schrödinger equation. However, for the relatively large size of the nanoribbons and in the presence of a small degree of disorder, the deviation from the Thomas–Fermi limit is expected to be small. The leading quantum correction to the classical capacitance should be captured by Eq. (2.4). Accordingly, the electrochemical potential \(\mu(r)\) contains in addition to the electrostatic potential \(\Phi(r)\) the contribution from the local Fermi energy \(E_F(r)\),

\[
\mu(r) = \Phi(r) + e^{-1} E_F(r). \quad (2.5)
\]

The solution of the coupled system of Eqs. (2.1), (2.4), and (2.5) allows the calculation of quantum capacitance effects.

Integrating the charge density over the spatial coordinates yields the total capacitance that relates the accumulated charge with the applied potential difference. The total inverse capacitance \(C_{tot}^{-1}\) is given by

\[
C_{tot}^{-1} = \frac{d\mu}{dQ} = \frac{d\Phi}{dQ} + \frac{1}{e} \frac{dE_F}{dQ} = C_{cl}^{-1} + C_{qm}^{-1}, \quad (2.6)
\]

where the inverse quantum capacitance reads

\[
C_{qm}^{-1} = \frac{1}{e} \frac{dE_F}{dQ} = \frac{1}{e^2 D(E_F)}, \quad (2.7)
\]

and is inversely proportional to the density of states at the Fermi level. For the latter we use in the following the bulk limit in line with the Thomas–Fermi approximation, while the variations of \(\mu(r)\) and \(E_F(r)\) over the length scale of the device are fully included. The relative importance of quantum capacitance corrections is governed by the ratio

\[
\frac{C_{qm}^{-1}}{C_{cl}^{-1}} = \frac{1}{e^2} \frac{dQ}{d\Phi} = \frac{1}{e^2 D(E_F)} \frac{dQ}{d\Phi}, \quad (2.8)
\]

i.e. the ratio of the electrostatically induced charge on the capacitor, \(dQ\), to the total charge \(e^2 D(E_F) d\Phi\) near the Fermi edge accessible by a potential difference \(d\Phi\).

In the classical limit, the charge induced on a capacitor at finite voltage is small compared to the total number of electrons at the Fermi level. Conversely, a small density of states at the Fermi level implies a large \(1/C_{qm}\), and thus a reduction of the total capacitance in Eq. (2.6).

For infinitely extended two-dimensional graphene, \(E_F(r)\) is the energy difference between the highest occupied state on the Dirac cone and the charge neutrality point (i.e. the so-called Dirac point), as a function of the
We aim to describe a capacitor formed by a graphene nanoribbon, a metallic back gate and two side gates [see Fig. 1(a)]. The step from idealized, infinite graphene to a nanoribbon with side gates (of possibly different geometries) modifies the expression for the density [Eq. (2.4)] to

$$n(r; T) = \int_{0}^{\infty} D(E) f(E - E_F(r), T) \, dE.$$  \hfill (2.11)

The total charge carrier density (electron and hole charge) for the Dirac cones in graphene then becomes

$$\rho_T(T) = \frac{2\pi}{\pi} \left( \frac{k_B T}{\hbar v_F} \right)^2 \left[ \text{Li}_2(-e^\eta) - \text{Li}_2(-e^{-\eta}) \right],$$  \hfill (2.12)

where \(\text{Li}_n(x)\) is the polylogarithm and \(\eta = E_F/k_B T\). Eqs. (2.12), (2.1) and (2.5) are solved for \(\Phi(r)\). All data we present in the following are evaluated at room temperature \(T = 300 \text{ K}\).

### III. RESULTS AND DISCUSSION

We consider the role of back gate voltages on the quantum capacitance of graphene nanoribbons. At finite temperature, the quantum capacitance corrections at, e.g., room temperature are important. We therefore investigate the relevance of quantum capacitance corrections at, e.g., room temperature. At finite temperature \(T\), the occupation of electronic states is smeared out by the Fermi–Dirac distribution function \(f(E, T)\) modifying the expression for the density [Eq. (2.4)] to

$$n(r; T) = \int_{0}^{\infty} D(E) f(E - E_F(r), T) \, dE.$$  \hfill (2.11)

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Materials and at different potentials) introduces several new device-specific quantum capacitance-related effects. We disentangle them by considering configurations of increasing complexity. To test the validity of our numerical simulation, we start our discussion with the simplest case of an infinitely extended graphene sheet [see Fig. 2(a)]. The nanocapacitor is completed by an infinite back gate at a distance of $d = 300 \text{ nm}$ [see Fig. 1(a)], the gap between the two sheets filled with a dielectric substrate, SiO$_2$ with $\varepsilon = 3.9$; above the nanoribbon is air. The boundary conditions for $\mu(\mathbf{r})$ are given by the potentials applied externally to the different gates. We assume translational symmetry perpendicular to the cross-section shown in Fig. 1(b), thus reducing the calculation to 2D (For technical details, see App. A). As expected, the capacitance decreases at the Dirac point [see Fig. 2(a)] signifying the quantum capacitance effect due to the reduction of the DOS at the Dirac point. Our numerical results agree (within the numerical accuracy) with analytical models.

A. Graphene nanoribbons without side gates

For a nanoribbon of finite width $w$ but without side gates, the quantum capacitance effect is substantially increased due to the smaller DOS, leading to a more pronounced quantum capacitance dip for narrower nanoribbons [see Fig. 2(b)]. Consequently, an inclusion of quantum capacitance effects becomes more important for smaller nanostructures. Concurrently, the classical capacitance per unit area of a nanoribbon increases with decreasing nanoribbon width [see Fig. 2(c)], further amplifying the absolute change in capacitance at the Dirac point. We find good agreement between our simulation and the analytical model discussed by Lin et al.\textsuperscript{32} [see blue dashed line in Fig. 2(c)],

$$C_{cl}(w) = C_0 \left[ \frac{2}{\pi} \arctan \left( \frac{w}{4d} \right) + \frac{w}{4d} \ln \left( 1 + \frac{16d^2}{w^2} \right) \right]^{-1},$$

where $C_0$ is the capacitance of the plate capacitor per unit area, $C_0 = \varepsilon_0 \varepsilon / d$. The residual differences are most likely due to the assumption of a spatially constant effective dielectric constant used in the analytical model (the average value $\varepsilon = (3.9 + 1)/2$ of the dielectric constants of substrate and air is used). As discussed above, one can use a full quantum local density of states (LDOS) in Eq. (2.4) beyond the Thomas–Fermi approximation. Using the quantum LDOS calculated for this geometry we have verified that such a correction yields a negligible change in the capacitance for realistic nanoribbon widths. In line with Eq. (2.8) we expect the approach
of the classical limit with increasing width \(w\) [Fig. 2(b)], illustrating that \(C_{\text{qm}}^{-1}/C_{\text{cl}}^{-1} \to 0\) as \(w \to \infty\) [Fig. 3(d)].

**B. Graphene nanoribbons with side gates**

The introduction of side gates precludes an analytic solution of the problem. Side gates decrease the overall capacitance of the nanoribbon since the classical field-lines further away from the nanoribbon no longer bend towards the nanoribbon (increasing its capacitance), but towards the side gates [see Fig. 1(b)]. Therefore, charge will accumulate at the side gates which decreases the nanoribbon capacitance [compare scale on Figs. 3(c) and 2(c)]. In addition to this classical effect, the different DOS in the graphene nanoribbon and metal side gates further increase the quantum capacitance effect [see Fig. 3(a)]. Put simply, it is energetically much more costly to put electrons into the nanoribbon (due to its small size and the small DOS) than into the metal side gates. A drastically different behavior emerges for graphene side gates [see Figs. 3(b),(c) and (d)], since at the Dirac point the DOS decreases in both the side gates and the nanoribbon simultaneously.

In the experiment, the electrostatic tuning of device properties proceeds by varying the voltages of the side gates relative to that of the back gate. As two prototypical examples, we treat the symmetric (i.e. \(V_{SG1} = V_{SG2}\)) and antisymmetric (i.e. \(V_{SG1} = -V_{SG2}\)) voltage configurations. Due to the non-uniform potential distribution \(\Phi(\mathbf{r})\), different parts of the device reach the (local) Dirac point at different voltage configurations, namely when locally \(\mu(\mathbf{r}) = \Phi(\mathbf{r})\).

We first consider metal side gates and the symmetric voltage configuration: for positive side gate voltages, the position of the quantum capacitance dip shifts to negative back gate voltages where the side gate influence is compensated [see Figs. 4(a) and 4(b)]. When the energy of the Dirac point is reached in the nanoribbon, the capacitance decreases. The relative depth of the dip depends inversely on the nanoribbon width. For finite voltages \(V_{SG}\), the side gates cause a non-uniform electrostatic potential in the nanoribbon, and the Dirac point is thus reached at different positions in the nanoribbon for different back gate voltages. With increasing side gate voltage, the non-uniformity of the electrostatic potential grows. Consequently, the quantum capacitance dip becomes wider and more shallow: a capacitance landscape in the \(V_{BG}-V_{SG}\) plane reveals two lines crossing each other at a small angle [see white lines in Fig. 4(b)] denoting the voltage combinations where no local charge is induced by the gates at either the center of the nanoribbon or its edges. These combinations control the width and shape of the quantum capacitance dip. For the antisymmetric
For graphene side gates, the situation becomes more complicated and novel features emerge: since the DOS of both the nanoribbon and the side gates features minima at the respective Dirac points, quantum capacitance (QC) effects in the nanoribbon and the side gates interact, leading to features beyond the standard “QC dip”. We find an increase in capacitance for larger side gate voltages [i.e. a quantum capacitance peak, see Fig. 5(a)]. This counter-intuitive increase in capacitance due to the limited density of states rather than a reduction arises from the reduced screening response by the side gates which, in turn, restores the capacitance of the nanoribbon. At specific combinations of back gate and side gate voltages, the local chemical potential in some region of the side gates coincides with the Dirac point. In this case, the screening ability of graphene side gates suffers due to the reduced capacitance in the side gates. In turn, this increases the capacitive coupling between the back gate and the nanoribbon. The loss in screening by the side gates thus causes a positive peak in the nanoribbon capacitance [see top right in Fig. 5(a)]. These results imply the surprising phenomenon of an effectively negative quantum capacitance. Previously, negative quantum capacitance effects were only reported due to many-body effects, in superconducting qubits, or for graphene heavily doped with Ag adatoms, which lead to dispersionless resonant impurity bands near the charge neutrality point. The explanation for the splitting and broadening of the QC dips applies analogously to the QC peaks [see dashed lines in Figs. 5(c),(d)].

A comparison of the size of the quantum capacitance dip suggests that the quantum capacitance effect is larger for grounded \(V_{SG1} = V_{SG2} = 0\) metal than grounded graphene side gates [compare Figs. 4(a) and 4(b)]. The observation that the capacitance of the nanoribbon increases when the side gates locally reach their Dirac point offers an additional explanation why the QC dip is deeper for metal side gates than for graphene side gates: since the nanoribbon and the side gates reach the Dirac point at the same potential, the large negative nanoribbon QC dip is superimposed on a positive effective side gate QC peak, and thus, the dip depth is decreased. This overlap can be seen in the capacitance distribution in the \(V_{BG}-V_{SG}\) plane [see Figs. 5(c),(d)]; regions of low (blue) and high (red) capacitance converge at the center. As soon as the graphene side gates are at a finite electrostatic potential, the QC peak and the QC dip are shifted relative to each other causing the broadening of the peak and an increase in dip depth.

Instead of considering the back gate capacitance as modified by side gates, one may alternatively consider separately the charges induced on the nanoribbon due to finite side gate and the back gate voltages: for each gate, we define a capacitance relating the voltage on the gate to the induced charge on the nanoribbon. A comparison of the relative capacitive couplings, i.e. the ratios of the capacitances, is of great interest for experiments: back gate and side gates usually feature very different coupling coefficients due to the different spatial distance to the nanoribbon. This relative coupling is usually referred to as relative lever arm \(\alpha_{rel} = C_{BG}/C_{SG}\). Accurate knowledge of \(\alpha_{rel}\) is essential for a detailed interpretation of experimental results. Due to quantum capacitance effects, \(\alpha_{rel}\) depends on \(V_{BG}\) and \(V_{SG}\), as we now investigate in detail. We determine side gate capacitance \(C_{SG}\) as \(C_{SG} = dQ_{GNR}/dV_{SG}|_{V_{BG}}\) and \(C_{BG} = dQ_{GNR}/dV_{BG}|_{V_{SG}}\). The relative lever arm as function of \(V_{BG}\) and \(V_{SG}\) features a substantial energy dependence already for metallic side gates [see Fig. 6(a)]; we find the relative lever arm decreased by roughly three per-
cent around zero carrier density in the nanoribbon. For the case of graphene side gates, we observe a decrease by up to fifteen percent of the relative lever arm along a positive diagonal in the gate voltage plane [highlighted in Fig. 6(b)]. This is of importance as experimental devices often feature side gates and nanoribbon etched from the same graphene sheet.

Interestingly, when comparing our numerical results with the experimental data reported in Ref. 49 one finds that the experimentally extracted $\alpha_{rel}$ is nearly a factor three larger than our calculation suggests. Since the back gate coupling is well controlled in experiments on large-scale graphene devices, we conjecture that this discrepancy might be related to a significantly reduced side gate coupling in the experiment. In particular, the edges of a realistic experimental nanoribbon device feature edge roughness and uncontrolled edge terminations (both not included in the present model) which may lead to a large local density of states at the edges, and thus to a significant charge accumulation at the edge of the nanoribbon. These charges may partly screen the lateral side gates and therefore increase the measured relative lever arm. Such a screening effect is also consistent with recent observations on hydrofluoric acid (HF) treated graphene nanoribbons where an increased side gate coupling strength has been found after HF dipping.\(^{50}\)

\section{IV. SUMMARY AND OUTLOOK}

We have shown that side gated graphene nanoribbons exhibit significant variation in capacitance as a function of gate voltages due to classical screening and quantum capacitance effects in the nanoribbon. Both the nanoribbon geometry as well as the presence and material of side gates strongly influence quantum capacitance effects. We find both positive and, surprisingly, unconventional negative quantum capacitance corrections. The former is the usual decrease of total capacitance due to the additional energy required to fill electronic states at a low density of states. The latter occurs due to a decrease of screening by graphene side gates. Since the capacitance of a graphene nanodevice frequently enters in the tuning of the effective energy of electrons in, e.g., transport experiments, inclusion of quantum capacitance effects is critical for the correct interpretation of experimental data. The proposed increase in total capacitance for finite voltage at graphene side gates and the significant change in the relative gate lever arms should be observable experimentally for clean samples.

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\section*{Appendix A: Numerical method}

In the following, we provide further technical details on our calculations. We solve the Poisson equation (2.1) using a finite difference scheme with a grid spacing of 1 nm. Dirichlet boundary conditions are applied to the conducting parts of the system: the lower boundary (where the back gate fixes the potential), the nanoribbon and the side gates. The left, right and top border are best modeled using Neumann boundary conditions, i.e. $\frac{\partial \Phi}{\partial x} = 0$, as these emulate the infinitely extended system better than Dirichlet boundary conditions. For the case of $T = 0$ K, we solve Eq. (2.5) for $\Phi(\mathbf{r})$ using the charge density for graphene, Eq. (2.9), in combination with the Poisson equation (2.1). After the discretization, the resulting nonlinear system of equations can be solved using any nonlinear solver. On all grid points $\mathbf{r}_i$ of the graphene parts, there is a difference between the electrochemical potential $\mu(\mathbf{r}_i)$ and the electrostatic potential $\Phi(\mathbf{r}_i)$, given by a discretized version of Eq. (2.5). Inserting these constraints into the Poisson equation (2.1) yields

$$\varepsilon_0 \Delta \Phi(\mathbf{r}) = \left\{ \begin{array}{ll} \frac{q}{\pi} \left( \frac{\mu(\mathbf{r}_i) - \Phi(\mathbf{r}_i)}{\text{vol}_h} \right)^2 & , \mathbf{r} \in \mathbf{r}_i \\ 0 & , \mathbf{r} \notin \mathbf{r}_i \end{array} \right. \quad (A1)$$

where $\mathbf{r}_i$ are the grid points on the graphene parts. This system of equations (for $\mathbf{r} = \mathbf{r}_i$, $i = 1, \ldots, N$) is solved for $\Phi(\mathbf{r}_i)$. For the temperature-dependent case, the procedure remains unchanged, except for the use of the temperature-dependent expression $\rho_f(E_F, T)$, i.e., Eq. (2.12) instead of (2.9).

Quantum capacitance can also be formulated for a non-local relationship between energy and charge (as required, e.g., for ab-initio calculations of the system)

$$\mu(\mathbf{r}) = \Phi(\mathbf{r}) + \frac{1}{e} E_F [\varepsilon_0 \nabla \cdot (\varepsilon(\mathbf{r}) \nabla \Phi(\mathbf{r}))] (\mathbf{r}). \quad (A2)$$

This equation can be reformulated in terms of charge densities instead of energies by applying the inverse map $\rho_f[\mu(\mathbf{r}), \Phi(\mathbf{r})]$,

$$\rho_f[\mu(\mathbf{r}), \Phi(\mathbf{r})] (\mathbf{r}) = \varepsilon_0 \nabla \cdot [\varepsilon(\mathbf{r}) \nabla \Phi(\mathbf{r})], \quad (A3)$$

which gives the charge density, $\rho_f$, as a function of the potential $\Phi(\mathbf{r})$ and the chemical potential $\mu(\mathbf{r})$. Assuming a local dependence between $E_F[\rho_f(\mathbf{r})] (\mathbf{r})$ and $\rho_f(\mathbf{r})$ allows replacing the functional formulation (A2) by the simpler (2.5) used in the main text.