Realizing Universal Edge Properties in Graphene Fractional Quantum Hall Liquids

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Universal chiral Luttinger liquid behavior has been predicted for fractional quantum Hall edge states, but so far has not been observed experimentally in semiconductor-based two-dimensional electron gases. One likely cause of this absence of universality is the generic occurrence of edge reconstruction in such systems, which is the result of a competition between confinement potential and Coulomb repulsion. We show that due to a completely different mechanism of confinement, edge reconstruction can be avoided in graphene, which allows for the observation of the predicted universality.

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Introduction. – Topological states of matter often support protected gapless edge or surface excitations, which in turn provide a window to probe the topology of the bulk phases. This is the case for integer and fractional quantum Hall (FQH) liquids, and topological insulators and superconductors. The FQH edge states are described by chiral Luttinger liquid (CLL) theory\textsuperscript{[1]}, and predicted to exhibit certain universal low-energy properties like electron tunneling exponents, for many bulk filling factors including the celebrated Laughlin sequence\textsuperscript{[2]}. Such universality, however, has not been observed in FQH liquids formed in semiconductor-based two-dimensional electrons gases (2DEGs)\textsuperscript{[3,4]}. This is clearly one of the most significant long-standing puzzles in the field of quantum Hall effect. One likely cause of this discrepancy is edge reconstruction\textsuperscript{[3,4]}, which induces additional non-chiral edge modes that are not tied to the bulk topology; these additional modes ruin the predicted universality\textsuperscript{[3,4]}. Edge reconstruction is a consequence of competition between confinement potential that holds the electrons in the interior of the sample, and Coulomb repulsion that tends to spread out the electron density. Detailed numerical studies\textsuperscript{[1,3,4]} show that due to the electrostatic configuration of semiconductor 2DEG, edge reconstruction occurs generically in the FQH regime. This suggests it is unlikely that one can observe the predicted universality in these systems, without carefully designing their electrostatic structure.

Recently, graphene has emerged as a brand new 2DEG system, in which FQH effects have been observed\textsuperscript{[5–12]}. Its many fascinating properties due to the Dirac nature of the electrons notwithstanding, we point out that the mechanism for electron confinement is very different between graphene and semiconductor 2DEGs. In the latter electrons are provided by dopants, which are positively charged (so that the system is overall charge neutral), and provide the dominant source of confinement potential. Since these dopants are placed hundreds or thousands of angstroms away from the 2DEG, the confining potential is not strong enough to balance the Coulomb repulsion at the edge, and prevent edge reconstruction\textsuperscript{[4,8,9]}. In graphene, on the other hand, electrons come from (properly biased) metallic gates. The charge neutrality is reflected by the fact that every electron carries an opposite image charge due to the metallic gate; as a result the electron-electron interaction is of dipole-dipole type at long distance, and no additional neutralizing back ground charge is present (or needed). Thus the one-body confinement potential is provided instead by the presence of graphene boundaries.

In this work we study the combined effects of the Dirac nature of electrons and the dipole nature of interaction, and show that there is an experimentally accessible parameter window in which edge reconstruction can be avoided. This points to the possibility of realizing universal CLL physics at graphene quantum Hall edges, and resolving a long-standing puzzle in the field.

Model. – In this Letter we study $\nu = 1/3$, at which the first FQH state was observed in both semiconductor 2DEGs\textsuperscript{[13]} and in single-layer graphene\textsuperscript{[10,11]}. In a semiconductor 2DEG, the $1/3$-filled lowest Landau level (LL) originates from a single band of nonrelativistic electrons. In graphene, however, the $0$th LL has a 4-fold spin and valley degeneracy with equal number of states coming from the conduction and valence bands that form two Dirac cones in a Brillouin zone. Hence the $\nu = 0$ state corresponds to a half-filled $0$th LL and is already nontrivial as there are multiple ways to occupy the LL. At zeroth order, Coulomb interaction is spin- and valley-independent, giving rise to an internal SU(4) symmetry, which is spontaneously broken due to the exchange effect that gives rise to quantum Hall ferromagnetism\textsuperscript{[14]}. Various symmetry-breaking perturbations lift the SU(4) degeneracy and select the occupied states [or a specific direction in the SU(4)-order-parameter space]. The two obvious possibilities for the...
As we will see later, the dimensionless ratio $d/l_B$ (with $l_B = \sqrt{\hbar c/eB}$ being the magnetic length) controls the stability of the Laughlin state in both cases. However, here the image charge moves with the electrons (and thus enters the two-body interaction) while the background charge is inert and provides the dominant part of one-body confining potential. It also bears some resemblance to that in Ref. [19], in which the image charges from the dielectric media modify the form of Coulomb interaction. Here the effective interaction between electrons contains their direct interaction in the graphene layer and their interaction with image charge on the other side of the metallic gate at a distance $2d$ away:

$$ V(r) = \frac{\epsilon^2}{r \sqrt{r^2 + (2d)^2}}. $$

(1)

$V(r)$ crosses over from $1/r$ at short distance to $1/r^3$ at long distance around $r \sim d$, and the significant pseudopotentials $(V_m)$ are those with $m \lesssim d/l_B$.

The one-body confining potential in graphene comes exclusively from the existence of a boundary where the electron wave function must vanish. In a semi-infinite graphene sheet with an edge at $x = 0$ in a magnetic field, the Dirac electron spectrum satisfies $D_\mu(-\sqrt{2}x_c) = 0$, where $D_\mu(x)$ is the parabolic cylinder function, $x_c = k l_B$, $k$ is wave vector parallel to the edge and $\mu = \epsilon^2/2$; $\epsilon$ is energy in units of $\hbar v_F/l_B$. The solutions give the energies for the LLs in the presence of such a boundary [20]. In the bulk, the solutions are $\mu = 0, 1, 2, \cdots$, hence the energy for the $n$th LL is $\epsilon_n = \sqrt{2n} \epsilon$. Close to the boundary, the LLs are no longer flat. For a sufficiently large quantum Hall droplet occupying $N_{orb}$ orbitals on a disk with the edge at a radius $r_c = \sqrt{2N_{orb}/l_B}$, we can recast the Dirac solution as

$$ D_\mu[-\sqrt{2}(x_c - r_c/l_B)] = 0. $$

(2)

We note that the LL energies solved from the Dirac equation are in units of $\hbar v_F/l_B$, while the electron Coulomb interaction is in units of $\epsilon^2/l_B$ (the choice of energy unit from now on). In analogy to the definition of the fine-structure constant $\alpha = e^2/(\hbar c) \sim 1/137$, we define the fine-structure constant for graphene $\alpha_g = e^2/hv_F = \epsilon^2/l_B^{1/2} \sim 2.2$. The largeness of the constant is due to the speed of light being much larger than the graphene Fermi velocity $v_F \sim 10^6 m/s$. We feed the solution of the lowest LL (LLL) energy at $x_c = r_c/l_B - \sqrt{2}m$, scaled by $1/\alpha_g$, as the edge potential on the $m$th orbital $U_m$ in the disk geometry. This results in the Hamiltonian

$$ H = \frac{1}{2} \sum_{mnl} V_{mn}^l c_{m+l}^+ c_{n+l} c_{m} + \sum_m U_m c_m^+ c_m, $$

(3)

where $c_m^+$ is the creation operator for an electron with angular momentum $m$ in the LLL. $V_{mn}^l$ is the interaction matrix element calculated from Eq. (1).

**Stability of Laughlin state for spin-polarized electrons.**

- The Laughlin state is the exact zero-energy ground state with a total angular momentum $M = M_0 = 3N(N-1)/2$ at $\nu = 1/3$ for the hard-core potential with $V_1 > 0$ and $V_{m>1} = 0$, in the absence of one-body (or confinement) potential. Positive $V_{m>1}$, or the longer-range components of the interaction, tend to spread out...
the electron density (against confining potential) and increase $M$ for the ground state; this is the driving force of edge reconstruction instability. Compared to that for a realistic GaAs/AlGaAs 2DEG system with long-range Coulomb interaction [4, 8] (hereafter referred to as the GaAs case), graphene has a shorter-range interaction and a different source of edge confinement potential. However, they share some common ingredients, e.g., a control parameter $d$ that dictates the electrostatic configuration. Therefore, we expect that only for a finite range of $d \sim l_B$ can the Laughlin phase be stabilized, as in the case of GaAs [4]. This window can be determined by examining the total angular momentum $M$ of the ground state and its overlap with the Laughlin state.

In Fig. 1(b) we map out in the $d$-$N_{orb}$ plane the regions in which the Hamiltonian has a global ground state at $M = M_0 = 3N(N-1)/2$ (i.e., the same as Laughlin state), for 5-10 electrons. The Laughlin regions first rapidly shift to smaller $d$ as $N_{orb}$ increases, and then level off for $d \sim l_B$. Correspondingly, for large $d$ the Laughlin-like state exists for a very small range of $N_{orb}$, while this range gets significantly larger for smaller $d$. The high sensitivity on $N_{orb}$ for larger $d$ indicates that the ground state with $M = M_0 = 3N(N-1)/2$ suffers from strong finite-size effects, related to the fact that in such finite size system the edge confining potential can also impact the bulk region (near the center of the disc) strongly. This point is supported by Fig. 1(c), where we replot the regions by shifting $N_{orb}$ to $N_{edge} = N_{orb} - (3N - 2)$, where $3N - 2$ is the least $N_{orb}$ needed to support the Laughlin state, in an attempt to separate out the sensitivity to the system size. We find in such a plot the Laughlin regions are dependent on system size for larger $d$, while they fall on top of each other and show very weak size dependence for (shaded region)

$$0.5l_B \lesssim d \lesssim 1.5l_B,$$  \hspace{1cm} (4)

indicating genuine stability of the Laughlin state in this region [21]. Our conclusion is further supported by calculating the overlap between the numerical ground states with the Laughlin state itself. As is clearly visible in Fig. 1(d), the overlap is very small for large $d$, but rapidly increases toward 1 as one approaches the stability window of Eq. (4).

The stability window of the Laughlin state, Eq. (4), is quite similar to that of GaAs case modeled in Refs. 4, 8, 9, where a closely related parameter $d$ characterizes the distance between the 2DEG and the background charge due to the ions. This is not an accident; the stability window is dictated by the very similar electrostatic configurations of both cases. As discussed in detail in Refs. 8 (see in particular its Fig. 1), the 2DEG and its corresponding neutralizing positive ion charge form a capacitor; the fringe field near the edge is the driving force of edge reconstruction instability. Once $d$ becomes large compared to $l_B$ which is the size of the LLL wave function, such an instability occurs. Here the situation is almost identical, with the capacitor formed by graphene 2DEG and the metallic gate. In GaAs $d$ is typically of order 1000 Å or more than $10l_B$ (essential for high mobility), thus edge reconstruction is essentially unavoidable. In graphene on boron-nitride, however, $d$ can be as small as a few nanometers or a small fraction of $l_B$ [12, 22]; as a result the stability window is within reach. We predict that in such a window edge reconstruction and other related instabilities can be avoided, resulting in a single chiral edge mode for the 1/3 FQH state. In this window single electron tunneling [25, 26] will exhibit a non-linear I-V characteristic $I \propto V^2$ with a universal exponent $\alpha = 3$, as predicted by theory [1].

Spin partially-polarized ground states. We now switch on the spin degrees of freedom. Earlier numerical studies on torus [22] and sphere [24] indicate that electrons at $\nu = 1/3$ are fully spin-polarized with Coulomb interaction, even in the absence of Zeeman splitting. Our numerical calculation confirms that a system of 7 electrons on a sphere with dipolar interaction has a ground state with total spin $S = 7/2$, as shown in Fig. 2(a). These results clearly suggest that we have a spin fully-polarized ferromagnet in the bulk. The situation, however, can be more complicated at the edge. It was shown [25, 26] that the $\nu = 1$ quantum Hall ferromagnet can support a spin texture at the edge. In the following we show this can also happen for the 1/3 edge when the Zeeman splitting is sufficiently small, and calculate the magnetic field needed to fully polarize the edge.

We choose 7 electrons in 26 orbitals at $d = 1.8l_B$ as an example, for which the ground state of spin-polarized electrons is inside the Laughlin-like region as shown in Fig. 1. Fig. 2(b) plots the low-lying energy spectrum of the system, in the absence of Zeeman splitting. The global ground state now has $S = 5/2$, indicating the ground state is not spin fully-polarized. To fully polarize the ground state, we need to apply a finite Zeeman energy, which exceeds the energy difference per electron.
FIG. 3: (Color online) (a) Zeeman energy necessary to fully-polarize the ground state of 7 (6 for inset) spinful electrons in the \( d_{N_{orb}} \) parameter space. (b) The Zeeman energy and the ground state total angular momentum for the 7 electrons along the (green) dashed line in (a) (i.e., \( N_{orb} = 26 \)). (c), (d), (e) and (f) plot the spin-resolved density profiles for 4 selected ground states in the 4 different spin regions in (b).

between the two states: \( E_z \geq \delta E/N = (E_{\text{polarized}} - E_{gs})/N \). Fig 3a plots in color the minimum Zeeman energy required to polarize the ground state to be Laughlin-like for systems of 6 and 7 electrons. In both cases we find the required \( E_z \sim 0.003 e^2/l_B \) with essentially no size dependence; this indicates both bulk and edge electron spins are fully polarized for \( B \geq 2 \) Tesla (with electron spin \( g \) factor 2). In these cases results of the previous section remain valid.

To reveal the nature of the spin partially-polarized states (which can be stable at sufficiently low magnetic fields), we fix \( N_{orb} = 26 \) and vary \( d \) inside the Laughlin-like region, indicated by the vertical (green) dashed line in Fig. 3a. As shown in Fig. 3b), the total spin \( S \) and the angular momentum \( M_{gs} \) for the ground state both vary with \( d \), indicating the interplay between edge confinement and spin structure. Fig. 3(c)-(f) show the spin-resolved density profiles of 4 ground states in the corresponding regions distinguished by the ground state spin \( S \) in Fig. 3b). The total density profile for these cases are almost identical. However, when the ground state is partially polarized, the minority-spin electron(s) (2 electrons for \( S = 3/2 \) and 1 for \( S = 5/2 \)) form a ring-like puddle along the edge; there is no electron with the minority spin near the center. This clearly indicates the reduction of polarization is due to spin texture formation at the edge, similar to that of \( \nu = 1/3 \). These edge spin textures, while eliminated by moderately high magnetic fields and unfavorable for the universal edge physics sought in this paper, are interesting in their own right and deserve further study, in light of the current experimental and theoretical interest in spin physics at quantum Hall edges.

In conclusion, we have demonstrated through detailed numerical studies that edge reconstruction can be avoided in an experimentally accessible parameter window for the \( \nu = 1/3 \) graphene fractional quantum Hall liquid, allowing for observation of universal chiral Luttinger liquid behavior. It can also support interesting edge spin textures that can be probed experimentally.

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