Quantized conductance of one-dimensional strongly correlated electrons in an oxide heterostructure

H. Hou,1 Y. Kozuka,2,3,4 Jun-Wei Liao,1 L. W. Smith,1 D. Kos,1 J. P. Griffiths,1 J. Falson,5 A. Tsukazaki,6 M. Kawasaki,2 and C. J. B. Ford1,*

1Cavendish Laboratory, University of Cambridge, Cambridge CB3 0HE, United Kingdom
2Department of Applied Physics and Quantum-Phase Electronics Center (QPEC), The University of Tokyo, Tokyo 113-8656, Japan
3Research Center for Magnetic and Spintronic Materials, National Institute for Materials Science (NIMS), 1-2-1 Sengen, Tsukuba 305-0047, Japan
4JST, PRESTO, Kawasaki, Saitama 332-0012, Japan
5Max Planck Institute for Solid State Research, D-70569 Stuttgart, Germany
6Institute for Materials Research, Tohoku University, Sendai 980-8577, Japan

(Received 5 July 2018; revised manuscript received 3 March 2019; published 25 March 2019)

Oxide heterostructures are versatile platforms with which to research and create novel functional nanostructures. We successfully develop one-dimensional (1D) quantum-wire devices using quantum point contacts on MgZnO/ZnO heterostructures and observe ballistic electron transport with conductance quantized in units of $2e^2/h$. Using dc-bias and in-plane field measurements, we find that the $g$ factor is enhanced to around 6.8, more than three times the value in bulk ZnO. We show that the effective mass $m^*$ increases as the electron density decreases, resulting from the strong electron-electron interactions. In this strongly interacting 1D system we study features matching the “0.7” conductance anomalies up to the fifth subband. This Rapid Communication demonstrates that high-mobility oxide heterostructures such as this can provide good alternatives to conventional III-V semiconductors in spintronics and quantum computing as they do not have their unavoidable dephasing from nuclear spins. This paves a way for the development of qubits benefiting from the low defects of an undoped heterostructure together with the long spin lifetimes achievable in silicon.

DOI: 10.1103/PhysRevB.99.121302

I. INTRODUCTION

Physical phenomena in transition-metal oxides and their complex compounds have stimulated intense interest in research covering metallic, semiconducting, and insulating properties. At the heterointerface of two oxide layers, the symmetry breaking leads to novel properties including quantum confinement of electrons, strong correlations, superconductivity, and ferromagnetism [1]. In a MgZnO/ZnO heterostructure, the polarization mismatch between MgZnO and ZnO originating from spontaneous and piezoelectric contributions induces a two-dimensional electron gas (2DEG) at the interface [2]. By engineering the strain via Mg composition and the MgZnO thickness, the 2DEG density can reach down to $10^{11}$ cm$^{-2}$ with mobility over $10^6$ cm$^2$ V$^{-1}$ s$^{-1}$ [3]. Both integer and fractional quantum-Hall effects have been investigated [2,4,5]. Furthermore, the weak spin-orbit interaction and the low concentration of nuclear spins in ZnO lead to a long spin-coherence time [6]. These unique properties in the MgZnO/ZnO heterostructure create an excellent platform for investigating quantum physics beyond the more conventional III-V alternatives [7,8]. More recently, Andreev reflection has been demonstrated at the interface between a MgZnO/ZnO heterostructure and MoGe superconductor, and this could be a good system for investigating non-Abelian quasiparticles, such as Majorana fermions [9].

Most low-dimensional devices use gates to define nanostructures such as one-dimensional (1D) quantum point contacts (QPCs)/quantum wires or quantum dots. However, gating oxide heterostructures is a challenge. In a MgZnO/ZnO heterostructure, we have found that using standard insulators such as Al$_2$O$_3$ causes a reduction in mobility and strong parallel conduction. The latter may be because the hard Al$_2$O$_3$ deposited on the thin stressed MgZnO layer contributes to a mismatch of the spontaneous and piezoelectric polarizations, and induces another 2DEG at the Al$_2$O$_3$/MgZnO interface. So far it has been necessary to use one-off techniques to create nanostructures, such as conducting atomic force microscopy lithography on LaAlO$_3$/SrTiO$_3$, which showed quantized conductance in units of $e^2/h$ in a strong magnetic field [10–12]. In our work, we have successfully prevented parallel conduction by replacing hard Al$_2$O$_3$ insulators with soft parylene C.

The zero-field quantization of conductance in integer multiples of $2e^2/h$ is a signature of ballistic charge transport in 1D systems. The lateral electrostatic confinement creates a series of 1D subbands, in which spin-up ($\uparrow$) and spin-down ($\downarrow$) subbands each contribute $e^2/h$. This is already observed in many materials, including GaAs/AlGaAs [13,14], InGaAs/InAlAs [15], GaN/AlGaN heterostructures [16], strained epitaxial germanium [17], and carbon-based materials [18,19]. An anomalous feature at conductance $G = 0.7 \times 2e^2/h$ was first investigated by Thomas et al. and attributed to a possible spontaneous spin polarization [20,21]. Its origin has since been much debated [22], and other explanations proposed
including quasi-bound-state formation and the Kondo effect [23,24]. Recently, Bauer et al. used a smeared van Hove singularity to explain it and emphasized the important role that electron-electron interactions play in the 0.7 anomaly [25].

Here we report ballistic transport in a high-quality MgZnO/ZnO heterostructure and show well-defined conductance quantization. Using dc-bias spectroscopy [26] and in-plane magnetic-field measurements, we find a g factor in the 1D wire that is enhanced by a factor of ~3 compared to the bulk and is fairly constant at low 1D subband index. Additionally, we show that the effective mass \( m^* \) increases as the density decreases in our 1D system, as occurs for 2DEGs in similar heterostructures [27,28]. The strongly correlated electron system in MgZnO/ZnO heterostructures offers a novel platform to investigate interaction effects, particularly the 0.7 structure. These strong correlations arise from the high \( m^* (= 0.5m_e \text{ in bulk ZnO, where } m_e \text{ is the bare electron mass}) \) and small dielectric constant \( \varepsilon = 8.3 \) compared to GaAs. The ratio \( r_s = E_C/E_F \) can reach 10 in low-density wafers \( (E_C \text{ is the average Coulomb energy per electron and } E_F \text{ is the Fermi energy}) \). These strong interactions may be the source of “\( N.7 \)” structures in 1D subbands with higher index \( N \) apparent in our data, which behave like the 0.7 structure. Any such \( N.7 \) structures [29–32] are very weak in GaAs electron or hole systems. We confirm the importance of electron-electron interaction in the 0.7 anomaly with an independent measurement of the strength of the electron-electron interaction from the electron effective mass [25].

II. EXPERIMENTAL DETAILS

The MgZnO/ZnO heterostructures are grown by molecular-beam epitaxy [3]. Devices A and B (C) have a 2DEG 92 nm (500 nm) below the surface, density \( 3.2 \times 10^{11} \text{ cm}^{-2} \) \((1.2 \times 10^{12} \text{ cm}^{-2})\), and mobility \( 3.5 \times 10^5 \text{ cm}^2 \text{ V}^{-1} \text{ s}^{-1} \) \((6 \times 10^4 \text{ cm}^2 \text{ V}^{-1} \text{ s}^{-1})\), corresponding to an electron mean free path of \( l_e = 3.2 \mu \text{m} \) \((1.1 \mu \text{m})\). A scanning electron micrograph and schematic cross section of a device are shown in Fig. 1(a). A Hall-bar mesa is patterned by Ar ion milling and Ti/Au is thermally evaporated to create Ohmic contacts without annealing. Ti/Au split gates of length \( L = 200 \text{nm} \) and width \( W = 300 \text{nm} \) are deposited on a 30-nm-thick parylene-C insulator layer, forming a quasi-1D wire in the 2DEG. We measure conductance through the QPC using a four-terminal lock-in technique with excitation voltage \( 10 \mu \text{V} \) at 77 Hz, at a temperature of \(~50 \text{ mK})\). For perpendicular magnetic-field measurements we measure the diagonal resistance to obtain the number of transmitted Landau-level edge states [33].

III. RESULTS AND DISCUSSION

A. Quantized 1D conductance

Figure 1(b) shows the 1D conductance vs split-gate voltage \( V_{\text{SG}} \). At \( V_{\text{SG}} = -1.5 \text{V} \), electrons below the gate are depleted and a quasi-1D wire is defined in the gap [inset, Fig. 1(b)]. This definition voltage matches the expected value calculated using a parallel-plate capacitor model with this 2DEG density, dielectric constant and thickness. 1D conductance plateaus appear as \( V_{\text{SG}} \) is made more negative, quantized in units of \( \Delta G \approx 2e^2/h \). They are visible up to \( 14e^2/h \) (devices A and B). For \( G < 2e^2/h \), Coulomb-blockade (CB) peaks appear, probably because dots form owing to the reduced electron screening increasing disorder, so we will discuss results from the second plateau and above.

Figure 1(c) shows the transconductance \( dG/dV_{\text{SG}} \) vs source-drain bias \( V_{\text{dc}} \) (devices A and B). The dc bias is calibrated with series resistance at zero gate voltage [34]. Dark (light) regions correspond to plateaus (transitions between plateaus). The splitting of the source and drain chemical potentials causes the risers to split until quantized plateaus appear between them at odd-integer values \( G = Me^2/h \) \((M = 3, 5, 7, \ldots)\), giving diamond-shaped features in Fig. 1(c) [26].

The 1D subband energy spacing \( \Delta E \) is given by the size of the diamond \( \Delta V_{\text{dc}} \) times \( e \), and is around 0.4 meV here. Surprisingly, this remains reasonably constant as the 1D subband index decreases. Because of the high \( m^* \) and hence relatively
small subband spacing, the plateaus are strongly temperature dependent, disappearing for $T \gtrsim 1 \text{ K}$ [35]. Note the straight line at $V_{SG} = -2.9 \text{ V}$ in Fig. 1(c) (device A) or near $V_{SG} = -2.6 \text{ V}$ (device B) must come from a random impurity or dot, perhaps under a gate, as they are only very slightly dependent on $B$ or $V_{d}$ and so provide a conduction path in parallel to the 1D wire.

B. Magnetic-field dependence and enhancement of $g$ factor

In an in-plane field, the quantized plateaus at even multiples of $e^2/h$ split due to the Zeeman energy $E_Z$ [Fig. 2(a)]. At $B_1 = 0.5 \text{ T}$, plateaus occur at $G = ne^2/h$ ($n = 3, 4, 5, \ldots$), when a 1D subband of the lower-energy spin-polarization direction becomes fully transmitted before the subband with opposite spin. As $B_1$ increases further to 1 T, the spin-split 1D subbands cross, leaving plateaus only at $G = M e^2/h$ ($M = 3, 5, 7, \ldots$). From $E_Z$ and the subband spacing, we estimate the $g$ factor as [26]

$$g^* = \frac{1}{\mu_B} \frac{\delta E}{\delta V_{SG}} = \frac{e}{\mu_B} \frac{\Delta V_{dc}}{\Delta B},$$

with $e$ the electronic charge and $\mu_B$ the Bohr magneton. We estimate $g^* \approx 6.8$ (6.4) for the second (fifth) subband, enhanced above the bulk value $g^* \approx 2$ for heterostructure and bulk ZnO [6]. An enhancement of $|g^*|$ by a similar factor occurs for electrons in GaAs QPCs, with values from 0.75 to 1.5 [20,23,36], compared to 0.44 in the bulk. For electrons in GaAs QPCs, $|g^*|$ decreases fairly rapidly with subband index [29], whereas in our data $g^*$ is almost constant, as the Zeeman splittings of subbands are almost identical [Fig. 2(a)]; the gradient of subband edges and the field at which subbands cross are very similar for subbands 2 to 6.

For a field applied perpendicular to the 2DEG ($B_\perp$), the electron energy contains Zeeman ($E_Z = g^* \mu_B B$) and cyclotron-energy ($E_c = \hbar \omega_c$) terms:

$$E_N(B) = (N + \frac{1}{2}) \hbar \omega_0 + \omega_c^2 \pm \frac{1}{2} g^* \mu_B B$$

($N$ is subband index, $\hbar \omega_0$ subband spacing assuming a parabolic potential, and $\omega_c = eB_\perp/m^*$ the cyclotron frequency) [37].

In the low-field regime ($\hbar \omega_0 \gg \hbar \omega_c$), spin-degenerate plateaus at even multiples of $G = e^2/h$ are split by $E_Z$, leading to additional plateaus at $G = me^2/h$ ($n = 3, 4, 5, \ldots$). These merge to odd multiples of $G = e^2/h$ with increasing $B_1$ as adjacent spin-split subbands cross. However, for high $B_1$, Landau-level formation leads to the creation of hybrid magnetoelectric subbands in the constriction [13,14,37,38], for which plateaus again occur at both even and odd integers. The onset of this regime can be quantified by the ratio $\kappa = \hbar \omega_c/E_Z = h e / (g^* m^* \mu_B)$. We estimate $m^* = (0.36 \pm 0.02)m_e$ from temperature-dependent Shubnikov–de Haas measurements and the Dingle formula [39,40]. This gives $\kappa < 1$ for $g^* = 6.8$. The smaller $m^* = 0.067 m_e$ and $g$ factor ($\approx 1$) in GaAs devices leads to $\kappa \gg 1$, so odd-integer plateaus are not observed.

Figures 2(b) and 2(c) show the evolution of subbands with $B_\perp$. Bright regions correspond to risers in conductance between plateaus, where subband edges cross the Fermi energy. Plateau heights are labeled. We model the subband energy vs $B$ using Eq. (2). Results for constant $m^* = 0.4 m_e$, $g^*$, and $\hbar \omega_0$ are plotted in Fig. 2(d), black and red lines representing spin-down (↓) and spin-up (↑) subbands, respectively. The pattern of plateaus matches the experiment well. However, the value of $B_\perp$ at which subbands cross is independent of subband index, which does not match the measurements. In Fig. 2(b), the crossing between $N = 2_1$ and $N = 3_1$ subbands occurs around $B_\perp = 0.8 \text{ T}$, and between $N = 3_1$, $4_1$ around $B_\perp = 1.1 \text{ T}$. For subbands $N > 4$, the required $B_1$ decreases. Given that $m^*$ in ZnO increases with decreasing density [27,28], we repeat the calculation with $m^*$ increasing from 0.4 to 1$m_e$ as the subband index decreases [Fig. 2(e)]. This reproduces the trend from experimental data that the value of $B_\perp$ at which spin-split subbands cross initially increases, then
devices, compared with the bulk value for ZnO. Conductances are indicated (units require strong negative voltages to define the 1D wire. The plateau decreases for higher subbands (although the model tends to underestimate due to energy blurring). We vary subband pairs decreases at lower subband index, unlike in GaAs. In addition, dc-bias spectroscopy [Fig. 3(c)] indicates a reasonably constant subband spacing over this range. Experimentally, the precise points at which bands cross at low index cannot be determined, and may be because of a strong electron-electron interaction, and nonparabolicity, disorder, and electron-phonon interactions may also contribute significantly to this increase [46].

FIG. 3. Transconductance $dG/dV_{SG}$ as a function of $V_{SG}$ and $V_{dc}$ for (a) device A at $B_{1} = 1$ T, and (b) device C at $B_{1} = 2$ T. In device C, the high electron density and thick MgZnO (500 nm) require strong negative voltages to define the 1D wire. The plateau conductances are indicated (units $e^2/h$). (c) $m^*$ measured vs $G$ in both devices, compared with the bulk value for ZnO.

C. Effective mass measurements

To investigate $m^*$ further we measure the dc-bias dependence at $B_{1} = 1$ T (device A) [Fig. 3(a)]. Adjacent spin-split subbands cross near this value of $B_{1}$, indicated by only odd plateaus being present for $V_{dc} = 0$. Since the subband spacing is roughly constant, each pair of subbands $2_{1}/3_{1}, 3_{1}/4_{1}$, etc., is degenerate since spin $\uparrow/\downarrow$ subbands are shifted by $+/-1/2g_{\parallel}B$, respectively, canceling out $E_{Z}$. The energy difference between these pairs of subbands is $\Delta E = \hbar\sqrt{\omega_{c0}^2 + \omega_{c}^2}$. In contrast to the $B = 0$ case, at $B_{1} = 1$ T the spacing between subband pairs decreases at lower subband index $N$ [Fig. 3(a)]. This could be explained by increasing $m^*$ at lower density, leading to a smaller $E_{c} = \hbar eB/m^*$ for lower $N$. Figure 3(b) shows the measurement repeated for device C with different

D. 0.7 anomaly

We now return to our discussion on the 0.7 anomaly. Figure 1(b) shows several shoulderlike features below the main plateaus. We test whether they behave similarly to the 0.7 anomaly or to CB-like resonances from impurities.

(i) Resonant peaks from CB should split with $V_{dc}$ at a rate determined by the size of the dotlike impurity and coupling to the gates, but dc-bias spectroscopy for our devices shows an orderly splitting above the second plateau typical of clean 1D devices [47,48] [Fig. 1(c)].

(ii) In Figs. 2(a)–2(c), even low $B$ lifts spin degeneracy. The edges of spin-split subbands do not meet at $B = 0$, showing clear gaps [for example, $\beta_0$ in Fig. 2(a)] at the plateau, as seen for the 0.7 anomaly in GaAs [20]. Figure 2(a) also shows gaps at higher-order crossings (labeled $\alpha_2$, $\beta_1$, $\beta_2$ matching labels in Ref. [49]), which is an important sign of the 0.7 analog [49]. The gaps can be explained as an effect of interactions [50].

(iii) The conductance sweeps in Fig. 2(a) are replotted (Fig. 4) as lines up to $B_{1} = 1$ T. The $N.7$ plateaus [just below $G = 2(N + 1)e^2/h$] appear to evolve smoothly to spin-polarized plateaus at $G = 2(N + 1/2)e^2/h$, then they split to form an extra plateau [indicated by $\ast$ in Figs. 2(a) and 4]. This split before the crossing was observed in GaAs [49], but was much weaker. How interactions contribute needs further theoretical investigation. In Fig. 4, the $N.7$ structures strengthen and occur lower on the riser for lower subbands, consistent with more significant interactions [25,51] due to the lower density.

(iv) Plateaus and $N.7$ structures stay reasonably constant as the wire is shifted laterally by asymmetric bias on the QPC gates [52]. The wire position should not significantly affect $n_{2D}$ and device dimensions ($L = 300$ nm and $W = 800$ nm), in which more plateaus are evident. The same trend of increasing spacing with subband index occurs.

Figure 3(c) shows $m^*$ vs plateau height (units $e^2/h$), estimated using $\Delta E$. At high conductance ($\sim 15e^2/h$), $m^* = (0.31 \pm 0.03)m_e$, close to the bulk effective mass found above for this wafer. When $G$ decreases to $5e^2/h$, $m^*$ increases to $(0.96 \pm 0.2)m_e$, which is comparable to that of a 2D system at a lower density of $1 \times 10^{11}$ cm$^{-2}$ ($0.8 \pm 0.2)m_e$ [28]. The large error bar is due to the blurred nature of this subband crossing. However, the trend of increasing $m^*$ as $G$ decreases is clear.

In previous studies of MgZnO/ZnO heterostructures, $m^*$ measured from temperature-dependent Shubnikov–de Haas oscillations increases as the 2DEG density decreases, while $m^*$ from cyclotron resonance is roughly constant [27,28]. This indicates that the increase in transport effective mass arises from electron-electron interactions, which are more significant at lower density. A similar effect is observed in other 2DEG systems [43,44], but much weaker (a factor of $\sim 1.4$ rather than 3 as in these ZnO heterostructures). In 1D GaAs wires, an increase in $m^*$ by at most 30% was reported when the 1D density decreased from $2.6 \times 10^{10}$ m$^{-1}$ to $1 \times 10^{8}$ m$^{-1}$ [45]. However, the ratio of electron-electron interaction energy to kinetic energy in GaAs is relatively low, and nonparabolicity, disorder, and electron-phonon interactions may also contribute significantly to this increase [46].
shoulder features resembling the 0.7 anomaly evolve with relative to the previous trace for clarity. Red dots illustrate how the resistance of around 110Ω is consistent with the model [25].

become better defined as the 1D subband index decreases, relative to the kinetic energy explains why estimated the strength of electron-electron interactions using electron-electron interactions. From the cyclotron energy, we effective mass and small dielectric constant leading to strong rapidly smeared by temperature, disappearing by small 1D subband spacing in our samples, any structure is additional test is the temperature dependence. Because of the complex magnetic-field behavior in GaAs are striking. An [35].

Either quantization or interaction effects inherent in the 1D system such as the 0.7 structure.

While tests (i)–(iv) described above are not fully comprehensive, they give a strong indication that these structures belong to the 0.7 family, and the great similarities with the complex magnetic-field behavior in GaAs are striking. An additional test is the temperature dependence. Because of the small 1D subband spacing in our samples, any structure is rapidly smeared by temperature, disappearing by T > 1 K [35]. N.7 structures are more visible in QPCs in MgZnO/ZnO compared to GaAs heterostructures because of the large effective mass and small dielectric constant leading to strong electron-electron interactions. From the cyclotron energy, we estimated the strength of electron-electron interactions using the electron effective mass. The increasing interaction energy relative to the kinetic energy explains why N.7 structures become better defined as the 1D subband index decreases, which is consistent with the model [25].

This MgZnO/ZnO heterostructure also shows dilute ferromagnetic properties with an anomalous Hall effect brought about by spin-dependent electron scattering off localized magnetic moments, which are likely to arise from point defects in epitaxial ZnO with localized unpaired electrons [53]. This may increase the strength of the 0.7-like structure [54] because this generally appears to be strongly related to spin [20,25]. However, the dilute ferromagnetic moments cannot be strong enough to produce full spontaneous electron spin polarization in the 1D wire, as only plateaus at even multiples of e²/h are observed at B = 0. Instead, the dilute ferromagnetism may possibly help to enhance the local spin susceptibility in the channel at B = 0 [25], making it easier for interactions to give rise to the shoulders on each plateau (the N.7 structure) seen, for example, as a pair of vertical lines in Fig. 2(c).

IV. CONCLUSION

To conclude, we have shown ballistic electron transport with conductance quantization in 1D quantum wires defined on a MgZnO/ZnO heterostructure. We also find an increasing effective mass at lower density, consistent with measurements on 2D ZnO systems. Because of the large g* and m*, a perpendicular field drives the system into a regime where the Zeeman energy is greater than the cyclotron energy, leading to only odd plateaus in the conductance. At zero field we see evidence of 0.7-like anomalies up to the fifth 1D subband. Such structures are rarely observed in GaAs, a key reason for which is probably the significantly higher interaction strength in ZnO. The ballistic transport and the importance of interactions and spin, together with a long spin-coherence time owing to the low concentration of nuclear spins in ZnO, could make high-quality MgZnO/ZnO heterostructures an interesting alternative to III-V semiconductors as platforms for quantum information and spintronics technologies.

ACKNOWLEDGMENTS

We thank S. Holmes for helpful discussions. H.H. acknowledges the Chinese Scholarship Council and Cambridge Trust for financial support. This work was partly supported by JST, PRESTO Grant No. JPMJPR1763 and JST, CREST Grant No. JPMJCR16F1, Japan.


