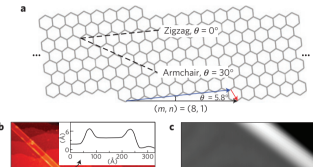
# Spatially resolving edge states of chiral graphene nanoribbons

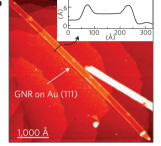
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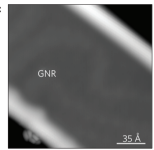
A central question in the field of graphene-related research is how graphene behaves when it is patterned at the nanometre scale with different edge geometries. A fundamental shape relevant to this question is the graphene nanoribbon (GNR), a narrow strip of graphene that can have different chirality depending on the angle at which it is cut. Such GNRs have been predicted to exhibit a wide range of behaviour, including tunable energy gaps<sup>1,2</sup> and the presence of one-dimensional (1D) edge states<sup>3-5</sup> with unusual magnetic structure<sup>6,7</sup>. Most GNRs measured up to now have been characterized by means of their electrical conductivity, leaving the relationship between electronic structure and local atomic geometry unclear<sup>8-10</sup>. Here we present a sub-nanometre-resolved scanning tunnelling microscopy (STM) and spectroscopy (STS) study of GNRs that allows us to examine how GNR electronic structure depends on the chirality of atomically well-defined GNR edges. The GNRs used here were chemically synthesized using carbon nanotube (CNT) unzipping methods that allow flexible variation of GNR width, length, chirality, and substrate 11,12. Our STS measurements reveal the presence of 1D GNR edge states, the behaviour of which matches theoretical expectations for GNRs of similar width and chirality, including width-dependent energy splitting of the GNR edge state.

The chirality of a GNR is characterized by a chiral vector (n,m) or, equivalently, by chiral angle  $\theta$ , as shown in Fig. 1a. GNRs having different widths and chiralities were deposited on a clean Au(111) surface and measured using STM. Figure 1b shows a room temperature image of a single monolayer GNR (GNR height is determined from linescans, such as that shown in Fig. 1b inset; some multilayer GNRs were observed, but we focus here on monolayer GNRs). The GNR of Fig. 1b has a width of 23.1 nm, a length greater than 600 nm, and exhibits straight, atomically smooth edges (the highest quality GNR edges, such as those shown in Figs 1 and 2, were observed in GNRs synthesized as in ref. 11). Such GNRs are seen to have a 'bright stripe' running along each edge.

This stripe marks a region of curvature near the terminal edge of the GNR that has a maximum extension of  $\sim$ 3 Å above the mid-plane terrace of the GNR and a width of  $\sim$ 30 Å (see line scan in Fig. 1b inset). Such edge-curvature was observed for all high-quality GNRs examined in this study (more than 150, including GNRs deposited onto a Ru(0001) surface). This







**Figure 1 | Topography of graphene nanoribbons (GNRs) on Au(111). a**, A schematic drawing of an (8, 1) GNR. The chiral vector (n,m) connecting crystallographically equivalent sites along the edge defines the edge orientation of the GNR (black arrow). The blue and red arrows are the projections of the (8, 1) vector onto the basis vectors of the graphene lattice. Zigzag and armchair edges have corresponding chiral angles of  $\theta=0^\circ$  and  $\theta=30^\circ$ , respectively, whereas the (8, 1) edge has an chiral angle of  $\theta=5.8^\circ$ . **b**, Constant-current STM image of a monolayer GNR on Au(111) at room temperature ( $V_{\rm S}=1.5~{\rm V}, I=100~{\rm pA}$ ). Inset shows the indicated line profile. **c**, Higher resolution STM image of a GNR at  $T=7~{\rm K}$  ( $V_{\rm S}=0.2~{\rm V}, I=30~{\rm pA}$ , greyscale height map).

is reminiscent of curved edge structures observed previously near graphite step-edges  $^{13}$ . We rule out that these GNRs are collapsed nanotubes by virtue of the measured ratio (observed to be  $\pi$ ) of GNR width to nanotube height for partially unzipped CNTs. We further rule out that the curved GNR edges observed here are folded graphene boundaries by means of a detailed comparison of terminal curved edges and actual folded edges (see Supplementary Information). Low-temperature STM images (Figs 1c and 2a) show finer structure in both the interior GNR terrace and the edge region. Figure 2a, for example, shows the atomically-resolved edge

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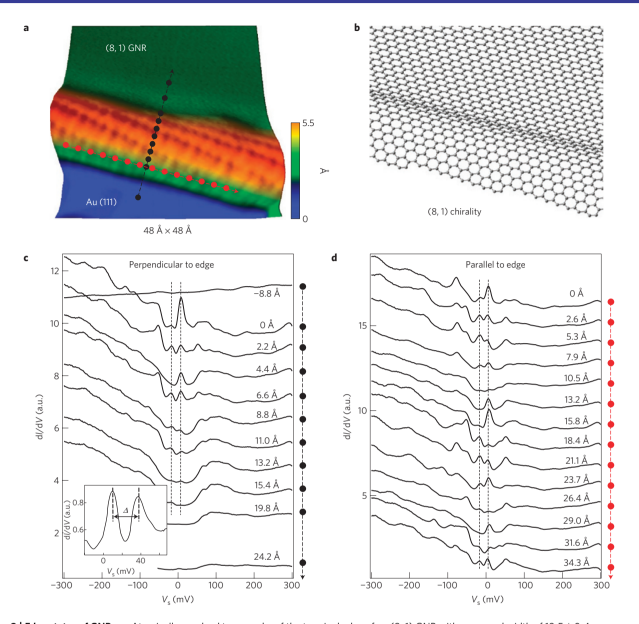


Figure 2 | Edge states of GNRs. a, Atomically-resolved topography of the terminal edge of an (8, 1) GNR with measured width of  $19.5 \pm 0.4$  nm ( $V_s = 0.3 \, \text{V}$ ,  $I = 60 \, \text{pA}$ ,  $T = 7 \, \text{K}$ ). b, Structural model of the (8, 1) GNR edge shown in a. c, dI/dV spectra of the GNR edge shown in a, measured at different points (black dots, as shown) along a line perpendicular to the GNR edge at  $T = 7 \, \text{K}$ . Inset shows a higher resolution dI/dV spectrum for the edge of a (5, 2) GNR with width of  $15.6 \pm 0.1 \, \text{nm}$  (initial tunnelling parameters  $V_s = 0.15 \, \text{V}$ ,  $I = 50 \, \text{pA}$ ; wiggle voltage  $V_{r.m.s.} = 2 \, \text{mV}$ ). The dashed lines are guides to the eye. d, dI/dV spectra measured at different points (red dots, as shown) along a line parallel to the GNR edge shown in a at  $T = 7 \, \text{K}$  (initial tunnelling parameters for c and d are  $V_s = 0.3 \, \text{V}$ ,  $I = 50 \, \text{pA}$ ; wiggle voltage  $V_{r.m.s.} = 5 \, \text{mV}$ ).

region of a monolayer GNR and clearly exhibits how the periodic graphene sheet of the GNR terminates cleanly and with atomic order at the gold surface.

Such high-resolution images allow us to experimentally determine the chirality of GNRs, and to create structural models of observed edge regions. In Fig. 2a, for example, we see rows of protrusions (with a spacing,  $\sim\!\!2.5$  Å, equivalent to the graphene lattice) near the edge of a GNR having a width of  $19.5\pm0.4$  nm. By comparing this row orientation with the orientation of the GNR terminal edge we are able to extract the GNR chirality (details in Supplementary Information). The GNR shown in Fig. 2a has an (8, 1) chirality (equivalent to  $\theta=5.8^\circ$ ), and the resulting structural model for this GNR is shown in Fig. 2b. We find the distribution of GNR chiralities to be essentially random. This is consistent with our structural data, which indicates that the CNT unzipping direction is very close to the axial direction of the precursor CNTs (see

Supplementary Information), as well as the fact that the precursor CNTs have a broad chirality distribution<sup>14</sup>.

We explored the local electronic structure of GNR edges using STS, in which dI/dV measurement reflects the energy-resolved local density of states (LDOS) of a GNR. Figure 2c,d shows dI/dV spectra obtained at different positions (as marked) near the edge of the (8, 1) GNR pictured in Fig. 2a. dI/dV spectra measured within 24 Å of the GNR edge typically show a broad gap-like feature having an energy width of  $\sim$ 130 meV. This is very similar to the behaviour observed in the middle of large-scale graphene sheets, and is attributed to the onset of phonon-assisted inelastic electron tunnelling<sup>15</sup> for  $|E| \geq 65$  meV. This feature disappears further into the interior of the GNR, as expected, because of increased tunnelling to the Au substrate<sup>16</sup>. Very close to the GNR edge, however, we observe further features in the spectra. The most dominant of these features are two peaks that rise up within the elastic tunnelling

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region (that is at energies below the phonon-assisted inelastic onset) and which straddle zero bias. For the GNR shown in Fig. 2a (which has a width of  $19.5 \pm 0.4$  nm) the two peaks are separated in energy by a splitting of  $\Delta = 23.8 \pm 3.2$  meV. Similar energy-split edge-state peaks have been observed in all the clean chiral GNRs that we investigated spectroscopically at low temperature. For example, the inset to Fig. 2c shows a higher resolution spectrum exhibiting energy-split edge-state peaks for a (5, 2) GNR having a width of 15.6 nm and an energy splitting of  $\Delta = 27.6 \pm 1.0$  meV. The two edge-state peaks are often asymmetric in intensity (depending on specific location in the GNR edge region), and their mid-point is often slightly offset from  $V_s = 0$  (within a range of  $\pm 20$  meV). As seen in the spectra of Fig. 2c, the amplitude of the peaks grows as one moves closer to the terminal edge of the GNR, before falling abruptly to zero as the carbon/gold terminus is crossed. The spatial dependence of the edge-state peak amplitude as one moves perpendicular from the GNR edge is plotted in Fig. 3a and shows exponential behaviour. The edge-state spectra also vary as one moves parallel to the GNR edge, as shown in Fig. 2d. The parallel dependence of the edge-state peak amplitude is plotted in Fig. 3b, and oscillates with an approximate 20 Å period, corresponding closely to the 21 Å periodicity of an (8, 1) edge.

We have also characterized monolayer GNRs having different chiralities and widths (the lengths of the GNRs used in these measurements are greater than 500 nm). In Fig. 3c, we plot the width dependence of the measured energy gap of GNR edge states for a broad range of chirality (3.7° <  $\theta$  < 16.1°). The measured edge-state energy splitting shows a clear inverse correlation with GNR width. Our gap values tend to be smaller than those observed previously for lithographically patterned GNRs (probably because of uncertainty in the edge structure of lithographically obtained GNRs; ref. 9).

The high quality of the atomically well-defined edge structures observed here allows us to quantitatively compare our experimental data to theoretical calculations of the electronic structure of chiral GNRs. We find that the spectroscopic features we observe correspond closely to the spatial- and energy-dependence predicted for 1D spin-polarized edge states coupled across the width of a chiral GNR. This behaviour is quite different from the properties observed previously for graphite step edges, armchair nanoribbons, and comparatively less ordered graphene platelet edges, where no magnetism-induced energy splitting has been seen 17-20.

To compare our experimental data with theoretical predictions for GNRs, we used a Hubbard model Hamiltonian, solved self-consistently in the mean-field approximation<sup>6</sup>, for an (8, 1) GNR having the same width as the actual (8, 1) GNR shown in Fig. 2a. The Hamiltonian:

$$H = -t \sum_{\langle ij \rangle, \sigma} [c_{i\sigma}^{\dagger} c_{j\sigma} + \text{h.c.}] + U \sum_{i} n_{i\uparrow} n_{i\downarrow}$$
 (1)

consists of a one-orbital nearest-neighbour tight-binding Hamiltonian with an on-site Coulomb repulsion term. In this expression  $c_{i\sigma}^{\dagger}$  and  $c_{j\sigma}$  are operators that create and annihilate an electron with spin  $\sigma$  at the nearest-neighbour sites i and j respectively, t=2.7 eV is a hopping integral<sup>21</sup>,  $n_{i\sigma}=c_{i\sigma}^{\dagger}c_{i\sigma}$  is the spin-resolved electron density at site i, and U is an on-site Coulomb repulsion. This GNR model is defined only by the  $\pi$ -bonding network. The terminal  $\sigma$ -bonds at the GNR edges are considered to be passivated and do not alter the  $\pi$ -system (this should, in general, correctly model a range of different possible edge-adsorbate bonding configurations<sup>7,22</sup>, including the likely oxygen-related functional group termination of our own GNRs; ref. 11). The out-of-plane curvature seen experimentally near GNR edges is not included in this model because the measured radii of curvature are sufficiently large (>20 Å) that they are not expected to significantly affect GNR electronic structure<sup>23</sup> (we

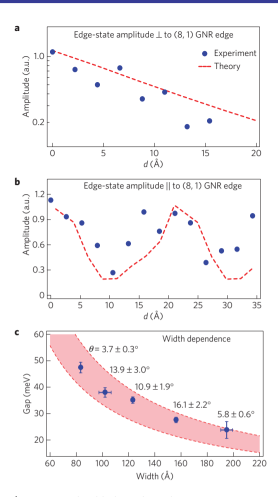
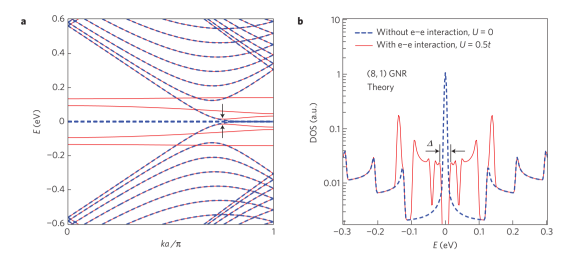


Figure 3 | Position- and width-dependent edge-state properties. a, Solid blue dots show experimental edge-state peak amplitude at points along a line perpendicular to the carbon/gold edge terminus (same positions as shown in Fig. 2c). Peak amplitude and energies were determined by fitting Lorentzian curves to the two peaks observed in the measured spectra at each location in Fig. 2c over the range  $-30 \,\mathrm{mV} < V_{\mathrm{s}} < 30 \,\mathrm{mV}$ . The energy positions of these peaks were found to be  $6.7 \pm 1.6 \, \text{mV}$  and  $-17.2\pm2.2$  mV. The positional dependence of the average peak amplitude of these two peaks is plotted. Error bars (shown when larger than plotted points) reflect the range of Lorentzian parameters that result in a good fit to the data. Dashed red line shows the calculated LDOS at locations spaced perpendicular to the edge terminus for an (8, 1) GNR (see text) at the energy of the DOS peak nearest the band-edge. Theoretical LDOS values include a single globally constant offset to model the added contribution from Au surface LDOS, and a single globally constant multiplicative factor to model the unknown total area of the STM tunnel junction. b, Solid blue dots show experimental average edge-state peak amplitude (determined as in a) at locations spaced along a line parallel to the carbon/gold edge terminus (same positions as shown in Fig. 2d). Dashed red line shows the theoretical edge-state LDOS for an (8, 1) GNR at points parallel to the edge terminus (calculated as in a). The edge-state LDOS amplitude oscillates parallel to the edge with a 21 Å period. c, Width dependence of the edge-state energy gap of chiral GNRs. From left to right, the chiralities of experimentally measured GNRs are (13, 1), (3, 1), (4, 1), (5, 2), and (8, 1) respectively, corresponding to a range of chiral angle  $3.7^{\circ} < \theta < 16.1^{\circ}$ . Energy gaps determined by Lorentzian fits to dI/dV peaks (centre-to-centre width), error bars reflect standard deviation due to spatial variation in spectra. GNR width measured as distance between the GNR edge mid-heights on opposite sides, error bars reflect standard deviation due to spatial variation along GNR axis. The pink shaded area shows the predicted range of edge-state bandgaps as a function of width, evaluated for chiral angles in the range  $0^{\circ} < \theta < 15^{\circ}$  (U = 0.5 t, t = 2.7 eV; ref. 27).



**Figure 4** | **Theoretical band structure and density of states (DOS) of a 20-nm-wide (8, 1) GNR. a**, Dashed blue line shows the calculated GNR electronic structure in the absence of electron-electron interactions (U = 0). Solid red line shows the calculated GNR electronic structure for U = 0.5t (t = 2.7 eV). Finite U > 0 splits degenerate edge states at E = 0 into spin-polarized bands, opening a bandgap (arrows). **b**, Dashed blue line shows the (8, 1) GNR DOS for the U = 0 case in **a**. The peak at E = 0 is due to the degeneracy of edge states in the absence of electron-electron interactions. Solid red line shows the (8, 1) GNR DOS for U = 0.5t. The opening of the bandgap (arrows) reflects the predicted energy splitting due to the onset of magnetism in spin-polarized edge states for U > 0, and compares favourably with the experimental data for the (8, 1) GNR of Fig. 2.

tested this conjecture by including the observed curvature in some calculations, and found that it has no significant effect—either from  $\sigma$ - $\pi$  coupling or from pseudofield effects—on the calculated GNR electronic structure). The effect of the gold substrate here is taken only as a charge reservoir that can slightly shift the location of  $E_{\rm F}$  within the GNR band structure and reduce the magnitude of the effective U parameter by means of electrostatic screening (the experimental charge-induced energy shifts seen here are within the range of charge-induced energy shifts observed previously for CNTs on Au; ref. 24).

We first calculated the GNR electronic structure for U = 0, which effectively omits the electron-electron interactions responsible for the onset of magnetic correlations. This results in the theoretical band structure and density of states (DOS) shown in Fig. 4a, b (blue dashed lines). The finite width of the GNR leads to a family of subbands in the band structure, with no actual bandgap (Fig. 4a). A flat band at E = 0 due to localized edge states leads to a strong van Hove singularity (that is, a peak) in the DOS at E = 0 (Fig. 4b). The DOS in this case does not resemble what is seen experimentally. We next calculated the (8, 1) GNR electronic structure for U > 0. Here the electron–electron interactions lift the degeneracy of the edge states by causing ferromagnetic correlations to develop along the GNR edges and antiferromagnetic correlations to develop across the GNR. This leads to a spin-polarization of the edge states that splits the single low-energy peak seen in the U = 0 DOS into a series of van Hove singularities, thus opening up a gap at E = 0. Such behaviour is seen in the band structure and DOS of Fig. 4a, b (solid red lines). We identify the lowest-energy pair of van Hove singularities with the pair of peaks observed experimentally near zero bias for GNR edges. We focus our experiment/theory comparison on the low-energy regime  $(|E| \le 65 \text{ meV})$  because higher energy experimental features are complicated by the onset of phonon-assisted inelastic tunnelling<sup>25</sup> (the low-energy edge-state peaks, by contrast, do not have the characteristics of inelastic modes).

We find that our experimental spectroscopic edge-state data for the (8, 1) GNR is in agreement with model Hamiltonian calculations for U = 0.5t. The theoretical bandgap of 29 meV is very close to the experimentally observed value of  $23.8 \pm 3.2$  meV (the value of U used here is lower than a value obtained previously from a first-principles calculation<sup>26</sup>, presumably because of screening

from the gold substrate). Our experimentally observed energy-split spectroscopic peaks thus provide evidence for the formation of spin-polarized edge states in pristine GNRs (such splitting does not arise for the non-magnetic U = 0 case described above). We are further able to compare the spatial dependence of the calculated edge states with the experimentally measured STS results. The dashed line in Fig. 3a shows the theoretical LDOS calculated at the energy of the low-energy edge-state peaks as one moves perpendicularly away from the GNR edge and into the (8, 1) GNR interior. The predicted exponential decay length of  $\sim$ 12 Å is in reasonable agreement with the experimental data. The variation seen in the calculated LDOS of the edge state in the direction parallel to the GNR edge also compares favourably with our experimental observations (Fig. 3b). The oscillation in edge-state amplitude is seen to arise from 'kinks' in the zigzag edge structure resulting from the chiral nature of the (8, 1) GNR edge (see the edge structure of Fig. 2b, each of the two dips in spectroscopic amplitude occurs at the location of a kink).

We are similarly able to compare the GNR width dependence of our experimentally measured edge-state gaps to theoretical calculations. As the measured GNRs having different widths also have different chiralities, we have calculated the theoretical edge-state gap versus width behaviour over the chirality range  $0^{\circ} < \theta < 15^{\circ}$  (it will be useful in the future to measure the local electronic properties of armchair GNRs ( $\theta = 30^{\circ}$ ), which are predicted to have no edge states). The pink shaded region in Fig. 3c shows the results of our calculations, and compares favourably with our experimentally observed width-dependent edge-state gap. This provides strong evidence that the edge-state gap we observe experimentally is not a local effect, as might occur, say, in response to some unknown molecules bound to the GNR edge, but rather depends on the full GNR electronic structure, including interaction between the edges.

### Methods

The GNRs shown in this study were produced by unzipping carbon nanotubes  $^{11}$ . GNRs were deposited onto clean Au(111) surfaces using a spin-coating method. Au(111) substrates were first cleaned by sputtering and annealing in ultra-high vacuum (UHV) before spin-coating. The samples were then transferred into the UHV chamber of our STM system (base pressure  $\sim\!2.0\times10^{-10}$  torr). After heat treatment up to 500 °C in UHV, the samples were directly transferred onto the STM stage in the same chamber for measurements.

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STM measurements were performed using a home-built STM held at low temperature ( $T=7\,\mathrm{K}$ ) for maximum spatial and spectroscopic resolution. STM topography was obtained in constant-current mode using a PtIr tip, and  $\mathrm{d}I/\mathrm{d}V$  spectra were measured using lock-in detection of the a.c. tunnelling current driven by a 451 Hz, 1–5 mV (r.m.s.) signal added to the junction bias (the sample potential referenced to the tip) under open-loop conditions. We also performed large-scale topographic surveys of GNR samples before low-temperature measurement. This was done in an Omicron variable temperature STM in UHV at room temperature. The three large-scale topographic STM images shown (Fig. 1b and Supplementary Figs S1a,b) were obtained in the Omicron STM at room temperature.

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#### **Author contributions**

C.T., L.J., Y-C.C., J.F., X.Z. and M.F.C. performed STM measurements and analysed STM data. L.J., C.T., Y-C.C., J.F., X.Z., J.M.T. and H.D. prepared samples. A.Z. assisted in structural analysis. O.V.Y., R.B.C. and S.G.L. carried out theoretical GNR simulations. All authors discussed the results and wrote the paper.

## **Additional information**

The authors declare no competing financial interests. Supplementary information accompanies this paper on www.nature.com/naturephysics. Reprints and permissions information is available online at http://www.nature.com/reprints. Correspondence and requests for materials should be addressed to M.F.C.