When electrons spin differently

Graphene nanoribbons: it's all about the edges

As reported by the journal Nature in its latest issue, researchers from Empa, the Max Planck Institute in Mainz and the Technical University of Dresden have for the first time succeeded in producing graphene nanoribbons with perfect zigzag edges from molecules. Electrons on these zigzag edges exhibit different (and coupled) rotational directions (“spin”). This could make graphene nanoribbons the material of choice for electronics of the future, so-called spintronics.

As electronic components are becoming ever smaller, the industry is gradually approaching the limits of what is achievable using the traditional approach with silicon as a semiconductor material. Graphene, the material with a number of “miraculous” properties, is considered a possible replacement. The one atom thin carbon film is ultra-light, extremely flexible and highly conductive. However, in order to be able to use graphene for electronic components such as field effect transistors, the material has to be “transformed” into a semiconductor. This was achieved by Empa scientists some time ago using a newly developed method - in 2010, they presented, for the first time, graphene nanoribbons (GNR) only a few nanometres wide with precisely shaped edges. For this, the ribbons were grown on a metal surface from specifically designed precursor molecules. The narrower the ribbons, the larger their electronic band gap - i.e. the energy range in which no electrons can be located, which is responsible for ensuring that an electronic switch (for example, a transistor) can be turned on and off. The Empa researchers were then also able to “dope” the nanoribbons, i.e. to furnish the ribbons with impurity atoms such as nitrogen at certain points, in order to influence the electronic properties of the graphene ribbons even more.

The perfect blueprint

In the paper now published in Nature, the Empa team led by Roman Fasel reports, together with colleagues from the Max Planck Institute for Polymer Research in Mainz, headed by Klaus Müllen, and from the Technical University of Dresden led by Xinliang Feng, how it managed to synthesise GNR with perfectly zigzagged edges using suitable carbon precursor molecules and a perfected manufacturing process. The zigzags followed a very specific geometry along the longitudinal axis of the ribbons. This is an important step, because researchers can thus give graphene ribbons different properties via the geometry of the ribbons and especially via the structure of their edges.
As with floor tiling, the right tiles - or precursor molecules - for the synthesis on the surface first had to be
found for the specific pattern of the zigzag graphene ribbons. Unlike in organic chemistry, which takes the
occurrence of by-products into account on the path to achieving a pure substance, everything had to be
designed for the surface synthesis of the graphene ribbons so that only a single product was produced. The
scientists repeatedly switched back and forth between computer simulations and experiments, in order to
design the best possible synthesis. With molecules in a U-shape, which they allowed to grow together to
form a snake-like shape, and additional methyl groups, which completed the zigzag edges, the researchers
were able to finally create a “blueprint” for GNR with perfect zigzag edges. To check that the zigzag edges
were exact down to the atom, the researchers investigated the atomic structure using an atomic force
microscope (AFM). In addition, they were able to characterise the electronic states of the zigzag edges using
scanning tunnelling spectroscopy (STS).

Using the internal spin of the electrons

And these display a very promising feature. Electrons can spin either to the left or to the right, which is
referred to as the internal spin of electrons. The special feature of the zigzag GNR is that, along each edge,
the electrons all spin in the same direction; an effect which is referred to as ferro-magnetic coupling. At the
same time, the so-called antiferromagnetic coupling ensures that the electrons on the other edge all spin in
the opposite direction. So the electrons on one side all have a “spin-up” state and on the other edge they all
have a “spin-down” state.

Thus, two independent spin-channels with opposite “directions of travel” arise on the band edges, like a road
with separated lanes. Via intentionally integrated structural defects on the edges or - more elegantly - via the
provision of an electrical, magnetic or optical signal from the outside, spin barriers and spin filters can thus
be designed that require only energy in order to be switched on and off - the precursor to a nanoscale and
also extremely energy efficient transistor.

Possibilities such as this make GNR extremely interesting for spintronic devices; these use both the charge
and the spin of the electrons. This combination is prompting scientists to forecast completely new
components, e.g. addressable magnetic data storage devices which maintain the information that has been
fed in even after the power has been turned off.

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(ERC) and the US Office of Naval Research (ONR).

Reference

Graphene nanoribbons slide on gold

In collaboration with researchers at the University of Basel and other international colleagues, Empa scientists recently also studied the tribological properties of graphene nanoribbons. In an article in the journal Science, they reported on the interactions of graphene nanoribbons that, with the tip of an atomic force microscope, have been pulled in different directions over a gold surface. With these experiments and thanks to powerful computer simulations, the researchers were able to demonstrate that virtually friction-free, floating movements were possible. The reason for the smoothness (superlubricity) is that the two atomic lattices on the crystalline surfaces of gold and graphene are completely incongruent with each other; latching cannot thus take place anywhere on the atomic “rolling landscape”.


Further information

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Pattern template for graphene nanoribbons: Depending on the direction of the ribbon axis, graphene nanoribbons have an armchair edge (orange) or a zigzag edge (blue).

"Blueprint" for the fabrication of zigzag graphene nanoribbons using a specifically synthesised precursor molecule.
Illustration of a graphene nanoribbon with zigzag edges and the precursor molecules used in its manufacture. Electrons on the two zigzag edges display opposite directions of rotation (spin) - "spin-up" on the bottom edge (red) or "spin-down" on the top edge (blue).

Atomic force microscopy image of the atomic structure of a zigzag graphene nanoribbon.