

Twisting and stacking double carbon layers

The name 'graphene' is given to a very thin form of carbon: specifically, a single layer of carbon atoms arranged in an hexagonal honeycomb-type lattice. Graphene research took off seriously in 2004 when single wafers were isolated for the first time. At about that time, scientists began to realise that graphene composed of two carbon layers (bilayer graphene) could be at least as interesting as single-layer graphene. Indeed, a growing number of researchers, both theorists and experimentalists, are now actively studying this form.

Since the early 2000s, several thousand papers have been published in this vibrant area of condensed-matter research. It is not easy to navigate this sea of information. An international team of researchers recently published a review in *Physics Reports* that should help newcomers make sense of the field.

Not all bilayer graphene systems are born equal: placing one layer on top of the other in different ways gives rise to non-equivalent forms. The most common varieties are AB-stacking (Bernal stacking), which is the most often studied;

AA-stacking; and the twisted bilayer. Initially, the interest in Bernal stacking was driven by the theoretical possibility of opening a controllable electron band gap with a transverse electric field. This field-sensitive gap was first proposed theoretically and then observed using an assortment of experimental tools. Our ability to control this non-negligible semiconducting gap suggests that the AB-bilayer might be useful in electronic applications and for photo-detection. This review discusses collective "many-body" effects in an AB-bilayer; the low-temperature, ordered phase of the system is of particular interest.



