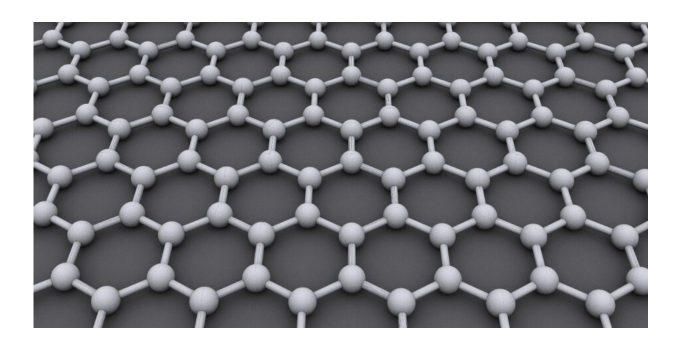


Graphene grows—physicists find a way to visualize it

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The typical two-dimensional hexagonal structure of graphene. For actual graphene, this honeycomb structure consists of carbon atoms – but now, it can also be recreated using micrometre-size polystyrene particles. Credit: Wikimedia Commons

Graphene is one of the strongest materials. On top of that, it is exceptionally good at conducting heat and electrical currents, making it one of the most special and versatile materials we know. For all these reasons, the discovery of graphene was awarded the Nobel Prize in



Physics in 2010.

Yet, many properties of the material and its cousins are still poorly understood—for the simple reason that the atoms they are made up of are very difficult to observe. A team of researchers from the University of Amsterdam and New York University have now found a surprising way to solve this issue.

Two-dimensional materials, consisting of a hyper-thin single layer of atomic crystal, have attracted a lot of attention recently. This well-deserved attention is mainly due to their unusual properties, very different from their three-dimensional 'bulk' counterparts. Graphene, the most famous representative, and many other <u>two-dimensional materials</u>, are nowadays researched intensely in the laboratory.

Perhaps surprisingly, crucial to the special properties of these materials are defects, locations where the <u>crystal structure</u> is not perfect. There, the ordered arrangement of the layer of atoms is disturbed and the coordination of atoms changes locally.

Visualizing atoms

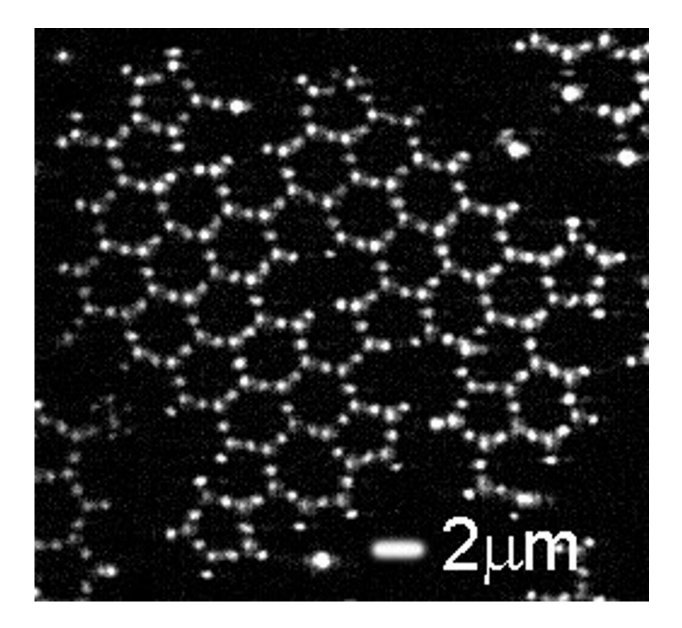
Despite the fact that defects have been shown to be crucial for a material's properties, and they are almost always either present or added on purpose, not much is known about how they form and how they evolve in time. The reason for this is simple: atoms are just too small and move too fast to directly follow them.

In an effort to make the defects in graphene-like materials observable, the team of researchers, from the UvA-Institute of Physics and New York University, found a way to build micrometer-size models of atomic graphene. To achieve this, they used so-called 'patchy particles'. These particles—large enough to be easily visible in a microscope, yet small



enough to reproduce many of the properties of actual atoms—interact with the same coordination as atoms in graphene, and form the same structure.

The researchers built a <u>model system</u> and used it to obtain insight into defects, their formation and evolution with time. Their results were published in *Nature Communications*.





Pieces of a graphene lattice made from patchy particles. Because the particles can be followed one-by-one, defects can be studied at the particle scale. Credit: University of Amsterdam

Building graphene

Graphene is made up of <u>carbon atoms</u> that each have three neighbors, arranged in the well-known 'honeycomb' structure. It is this special structure that lends graphene its unique mechanical and electronic properties. To achieve the same structure in their model, the researchers used tiny particles made of polystyrene, decorated with three even tinier patches of a material known as 3-(trimethoxysilyl)propyl—or TPM for short.

The configuration of the TPM patches mimicked the coordination of carbon atoms in the graphene <u>lattice</u>. The researchers then made the patches attractive so that the particles could form bonds with each other, again in analogy with the carbon atoms in graphene.

After being left alone for a few hours, when observed under a microscope the 'mock carbon' particles turned out to indeed arrange themselves into a honeycomb lattice. The researchers then looked in more detail at defects in the model graphene lattice. They observed that also in this respect the model worked: it showed characteristic defect motifs that are also known from atomic graphene.

Contrary to real graphene, the <u>direct observation</u> and long formation time of the model now allowed the physicists to follow these defects from the very start of their formation, up to the integration into the lattice.



Unexpected results

The new look at the growth of graphene-like materials immediately led to new knowledge about these two-dimensional structures. Unexpectedly, the researchers found that the most common type of defect already forms in the very initial stages of growth, when the lattice is not yet established. They also observed how the lattice mismatch is then 'repaired' by another defect, leading to a stable defect configuration, which either remains or only very slowly heals further to a more perfect lattice.

Thus, the model system not only allows researchers to rebuild the graphene lattice on a larger scale for all sorts of applications, but the direct observations also allow insights into atomic dynamics in this class of materials. As defects are central to the properties of all atomically thin materials, these direct observations in model systems help further engineer the atomic counterparts, for example for applications in ultra-lightweight materials and optical and electronic devices.

The findings are published in Nature Communications.

More information: Piet J. M. Swinkels et al, Visualizing defect dynamics by assembling the colloidal graphene lattice, *Nature Communications* (2023). DOI: 10.1038/s41467-023-37222-4. www.nature.com/articles/s41467-023-37222-4

Provided by University of Amsterdam

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