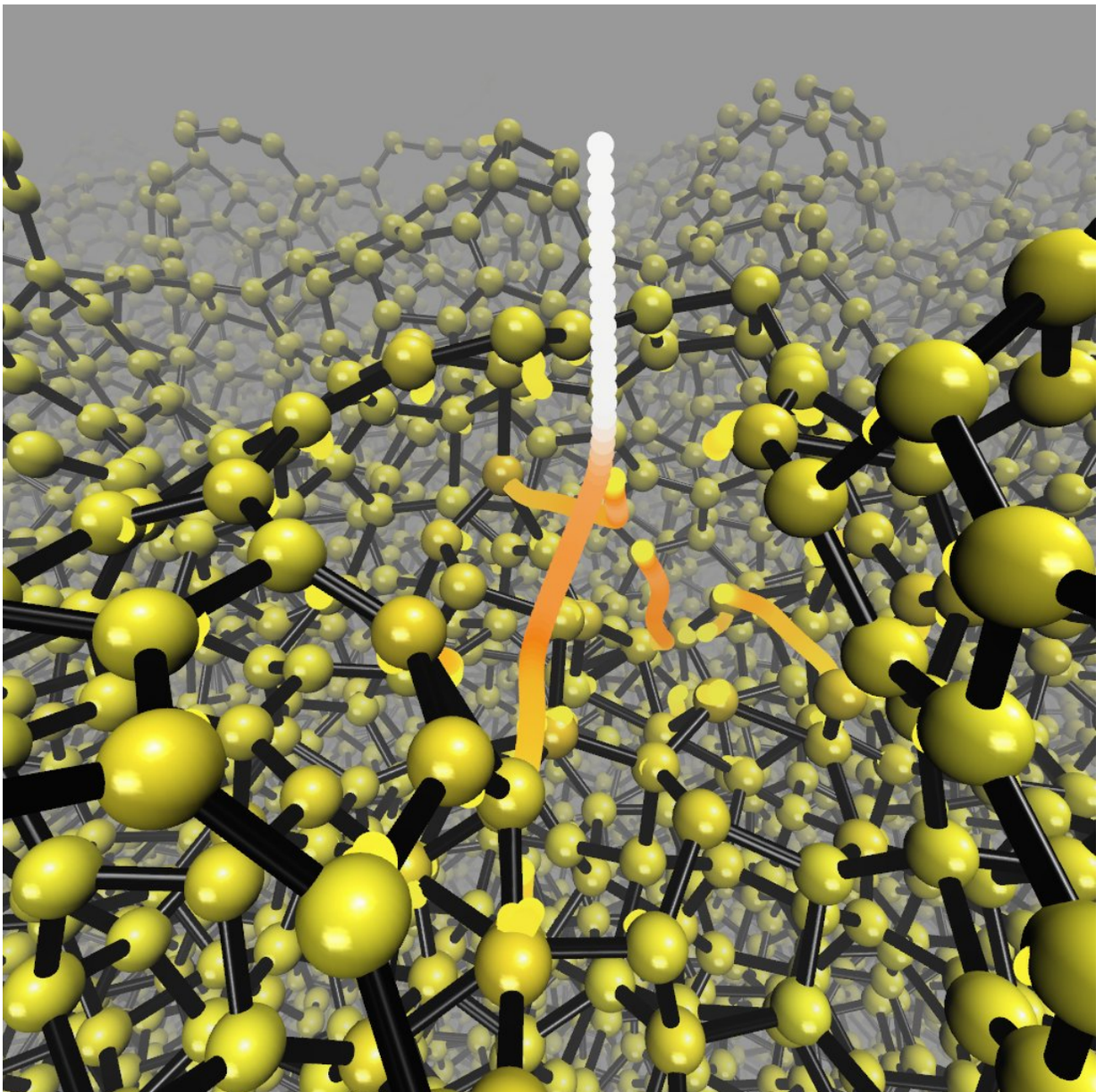


Diamond-like carbon is formed differently to what was believed – machine learning enables development of new model

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Trajectories followed by incident and knockon atoms during energetic deposition of a tetrahedral amorphous carbon thin film. Credit: Aalto University

Researchers at Aalto University and Cambridge University have made a significant breakthrough in computational science by combining atomic-level modeling and machine learning. For the first time, the method has been used to realistically model how an amorphous material is formed at the atomic level: that is, a material that does not have a regular crystalline structure. The approach is expected to have impact on the research of many other materials.

"The secret of our success is [machine learning](#), through which we can model the behaviour of thousands of [atoms](#) over long periods of time. In this way, we have obtained a more accurate model," explains Postdoctoral Researcher Miguel Caro.

The team's simulations reveal that diamond-like [carbon](#) film is formed at the atomic level in a different way than was thought. The prevailing understanding over the last 30 years of the formation mechanism for [amorphous carbon](#) film has been based on assumptions and indirect experimental results. Neither a good nor even an adequate atomic-level model has been available up to now. The new method has now overturned the earlier qualitative models and provided a precise atomic-level picture of the formation mechanism.

"Earlier, amorphous carbon films were thought to form when atoms are packed together in a small area. We have demonstrated that mechanical shock waves can cause the formation of diamond-like atoms further away from the point at which the impacting atoms hit the target, reports

Caro, who performed the simulations on CSC (IT Center for science) supercomputers, modelling the deposition of tens of thousands of atoms.

Results open up significant new avenues for research

There are countless different uses for amorphous carbon. It is used as a coating in many mechanical [applications](#), such as car motors, for example. In addition, the material can also be used for medical purposes and in various energy-related, biological and environmental applications.

"For us, the most important application is biosensors. We have used very thin amorphous carbon coatings for identifying different biomolecules. In these applications, it is especially important to know the films' electrical, chemical and electrochemical properties and to be able to customise the material for a particular application," explains Professor Tomi Laurila.

Dr. Volker Deringer, a Leverhulme Early Career Fellow, is particularly excited about using these methods for amorphous materials.

"Teaming up has been a great success," conclude Deringer and Caro, who are continuing the collaboration between their institutions through ongoing visits. The team expect that their approach will help many others in experimental materials research, because it can give information about [materials](#) with a level of precision close to that of quantum mechanical methods, but simultaneously can make use of thousands of atoms and long simulation times. Both of these are extremely important for a realistic picture of the processes in experiments.

"I'm especially excited about the kinds of opportunities this method offers for further research. This atomic-level model produces verifiably correct results that correspond exceptionally well to the experimental

results, revealing also for the first time the [atomic-level](#) phenomena behind the results. Using the [model](#), we can, for example, predict what kind of carbon surface would be best for measuring neurotransmitters dopamine and serotonin," says Laurila.

More information: Miguel A. Caro et al. Growth Mechanism and Origin of High sp^3 Content in Tetrahedral Amorphous Carbon, *Physical Review Letters* (2018). [DOI: 10.1103/PhysRevLett.120.166101](https://doi.org/10.1103/PhysRevLett.120.166101)

Provided by Aalto University

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