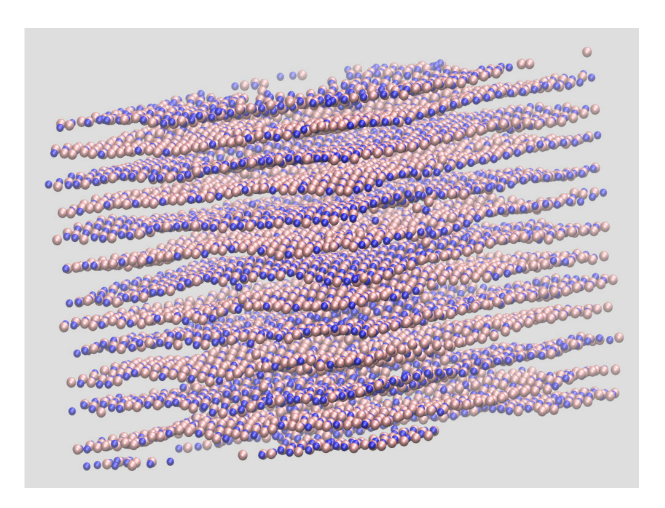


'Deep learning' casts wide net for novel 2-D materials

April 10 2019



Rice University researchers used a microstructure model of radiation-damaged hexagonal boron nitride to help them study the benefits of deep learning techniques in simulating two-dimensional materials to understand their characteristics. Credit: Prabhas Hundi/Rice University



Scientists are discovering new two-dimensional materials at a rapid pace, but they don't always immediately know what those materials can do.

Researchers at Rice University's Brown School of Engineering say they can find out fast by feeding basic details of their structures to "deep learning" agents that have the power to map the <u>materials</u>' properties. Better yet, the agents can quickly model materials scientists are thinking about making to facilitate the "bottom-up" design of 2-D materials.

Rouzbeh Shahsavari, an assistant professor of civil and environmental engineering, and Rice graduate student Prabhas Hundi explored the capabilities of neural networks and multilayer perceptrons that take minimal data from the simulated structures of 2-D materials and make "reasonably accurate" predictions of their physical characteristics, like strength, even after they're damaged by radiation and high temperatures.

Once trained, Shahsavari said, these agents could be adapted to analyze new 2-D materials with as little as 10 percent of their structural data. That would return an analysis of the material's strengths with about 95 percent accuracy, he said.

"This suggests that transfer learning (in which a deep-learning algorithm trained on one material can be applied to another) is a potential gamechanger in material discovery and characterization approaches," the researchers suggested.

The results of their extensive tests on graphene and hexagonal boron nitride appear in the journal *Small*.

Since the discovery of graphene in 2004, atom-thick materials have been touted for their strength and range of electronic properties for composites and electronics. Because their atomic arrangements have a significant impact on their properties, researchers often use molecular



dynamics simulations to analyze the structures of new 2-D materials even before trying to make them.



Rice University graduate student Prabhas Hundi, left, and Rouzbeh Shahsavari, an assistant professor of civil and environmental engineering, are using deep learning techniques to speed up simulations of novel two-dimensional materials to understand their characteristics and how they're affected by high temperature and radiation. Credit: Jeff Fitlow/Rice University

Shahsavari said deep learning offers a significant speed boost over such traditional simulations of 2-D materials and their characteristics, allowing calculations that now take days of supercomputer time to run in hours.



"Because we can build our structure-property maps with only a fraction of the data from graphene or boron nitride molecular dynamics simulations, we see an order of magnitude less computational time to get a full behavior of the material," he said.

Shahsavari said the lab decided to study graphene and hexagonal <u>boron</u> <u>nitride</u> for their high tolerance to deterioration under <u>high temperatures</u> and in radiation-rich environments, important properties for materials in spacecraft and nuclear power plants. Because the Shahsavari group had already carried out more than 11,000 radiation cascade damage <u>molecular dynamics simulations</u> for another paper on 2-D materials, they had incentive to see if they could reproduce their results with a much faster method.

They ran thousands of "deep learning" simulations on 80 combinations of radiation and temperature for <u>hexagonal boron nitride</u> and 48 combinations for graphene, hitting each combination with 31 random doses of simulated radiation. For some, the researchers trained the deep learning agent with a maximum of 45 percent of data from their molecular dynamics study, achieving up to 97 percent accuracy in predicting defects and their effects on the material's characteristics.

Adapting trained agents to different materials, they found, required only about 10 percent of the simulated data, greatly speeding up the process while retaining good accuracy.

"We tried to figure out the corresponding residual strengths of the materials after exposure to extreme conditions, along with all the defects," he said. "As expected, when the mean temperature or the radiation were too high, the residual strength became pretty low. But that trend wasn't always obvious."

In some cases, he said, the combined higher radiation and higher



temperatures made a material more robust instead of less, and it would help researchers to know that before making a physical product.

"Our <u>deep learning</u> method on the development of structure-property maps could open up a new framework to understand the behavior of 2-D materials, discover their non-intuitive commonalities and anomalies, and eventually better design them for tailored applications," Shahsavari said.

More information: Prabhas Hundi et al, Deep Learning to Speed up the Development of Structure–Property Relations For Hexagonal Boron Nitride and Graphene, *Small* (2019). DOI: 10.1002/smll.201900656

Provided by Rice University

Citation: 'Deep learning' casts wide net for novel 2-D materials (2019, April 10) retrieved 2 May 2024 from <u>https://phys.org/news/2019-04-deep-wide-net-d-materials.html</u>

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