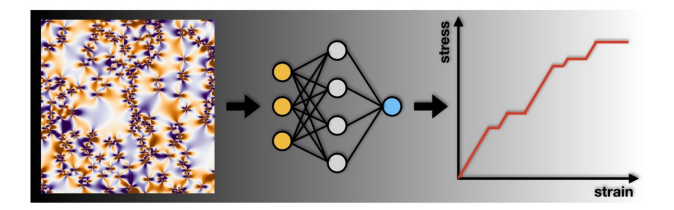


Machine learning to predict and optimise the deformation of materials

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Credit: Aalto University

Researchers at Tampere University of Technology and Aalto University taught machine learning algorithms to predict how materials stretch. This new application of machine learning opens new opportunities in physics and possible applications can be found in the design of new optimal materials. The study has been published in the prestigious journal *Nature Communications*.

Most regular objects tend to stretch 'evenly', that it: scientists can predict how much force is required to make a material stretch by a certain



distance. Recent experiments have shown that these predictions don't hold up at the micrometre scale. The stretching of microscopic crystals happens in discrete bursts with a very wide size distribution. Since the bursts occur sporadically, seemingly identical micro-scale samples can stretch in very different ways. This variability of the strength characteristics of the samples poses a challenge for the development of novel materials with desired properties. In their article "Machine learning plastic deformation of crystals" published in *Nature Communications*, the researchers use <u>machine learning</u> to predict the characteristics of individual samples.

"The machine learning algorithms succeeded in measuring how predictable the stretch process of small crystalline samples is. This would have been practically impossible with traditional means, but machine learning enables the discovery of new and interesting results," explains Associate Professor Lasse Laursonfrom the Laboratory of Physics at Tampere University of Technology.

The irreversible plastic deformation of crystalline substances occurs when crystallographic defects, called dislocations, move from one location in the crystal to another. Crystalline materials, such as metals or ice almost always contain networks of dislocations, with each crystal containing its own network.

The researchers trained machine learning algorithms to recognise the connection between an object's microscopic structure and the amount of force required to stretch a sample. The study revealed, amongst other things, that the predictability of the amount of force required changes on the stretching of the sample: At first, it becomes harder to predict the force required as the stretch grows, which depends mostly on the stretch bursts' sporadic nature. Surprisingly, however, predictability improves as the stretch continues to grow. Size also affects predictability: it is easier to predict the deformation process of larger crystals than smaller ones.



"As the stretch grows, the number of bursts reduces, consequently improving predictability. This is promising in terms of predicting the yield of individual samples, which is a key objective in material physics," says Henri Salmenjoki, <u>doctoral candidate</u> at the Department of Applied Physics at Aalto University.

"Our research indicates that machine learning can be used to predict very complex and non-linear physical processes. In addition to the development of optimal <u>materials</u>, possible applications can be found in the prediction of dynamics of many other complex systems," Laurson explains.

More information: Henri Salmenjoki et al. Machine learning plastic deformation of crystals, *Nature Communications* (2018). DOI: 10.1038/s41467-018-07737-2

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