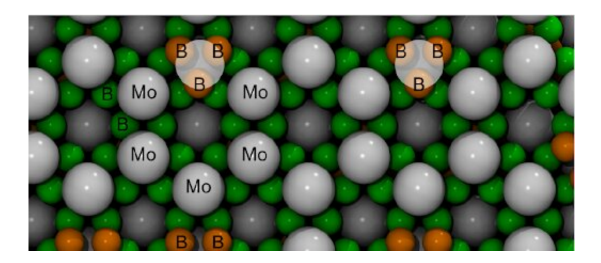


Scientists calculate crystal structure of superhard molybdenum borides

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Atom arrangement in the highest molybdenum boride (MoB5-x) crystal Credit: Dmitry V. Rybkovskiy, J. Phys. Chem. Lett.

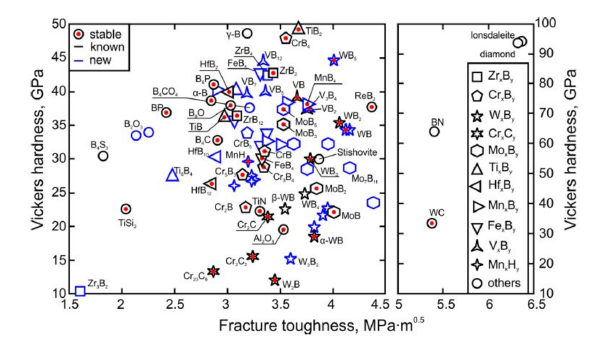
In their search for new superhard compounds, researchers have carried out a prediction of stable molybdenum borides and their crystal structures. They revealed that the highest borides contain four to five boron atoms per each molybdenum atom. The estimated Vickers hardness of MoB5 is 37 to 39 GPa, which makes it a potential superhard material. The study was published in *The Journal of Physical Chemistry Letters*.

Previously, a group of researchers led by Artem Oganov, professor at Skoltech and MIPT, published a study in the *Journal of Applied Physics*



in which they compiled a list of hard and superhard materials that may have useful industrial applications. This list, made using the evolutionary algorithm USPEX and new model of Vickers hardness (pressure required to leave a pyramid-shaped indentation in the material) and fracture toughness (ability of a material to resist propagation of a fracture), was dubbed a "treasure map" for experimenters by its authors.

In the new paper, scientists from Skoltech, Moscow Institute of Physics and Technology, A. M. Prokhorov General Physics Institute of RAS, Pirogov Russian National Research Medical University, and Northwestern Polytechnical University (Xi'an, PRC) studied the molybdenum borides region of the map. Transition metal borides are a potential replacements for traditional hard alloys and superhard materials in technological applications. Unlike widely used diamond and cubic boron nitride, transition metal borides do not require <u>high pressure</u> for their synthesis, making their production cheaper.





Hard and superhard materials on the "treasure map". The horizontal position of a symbol reflects its fracture resistance and its vertical position reflects its Vickers hardness. Black symbols represent known materials, blue ones represent new materials; symbols marked by a red spot indicate materials that are stable at ambient conditions Credit: Alexander G. Kvashnin, *Journal of Applied Physics*

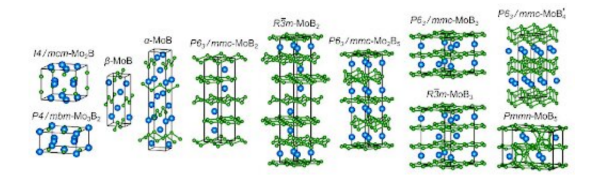
The high-valence electron density of the metal <u>atoms</u> resists compression due to electrons repelling each other, and covalent boron-boron and boron-metal bonds resist elastic and plastic deformations.

"The simulated X-ray diffraction (XRD) patterns are usually compared to those suggested in the experiment in order to determine whether the predicted structure is compatible with the experimental one. However, considering transition metal borides, such as molybdenum borides, the XRD pattern will only show signals from the heavier atoms while positions of the light boron atoms will be essentially invisible. This is why <u>crystal structure</u> models based only on experimental data are often unrealistic and unstable. Therefore, a comprehensive approach to determining crystal structures requires state-of-the-art theoretical calculations," says Alexander Kvashnin, one of the authors and a senior research fellow at Skoltech and MIPT.

Molybdenum pentaboride MoB_5 was found to be the stable highest boride, however, the simulated XRD patterns were close but not identical to the experimental data. The predicted pentaboride had a few weak peaks that were not observed in the experiments. This hinted at a higher symmetry in the experimental sample. Key structural elements of the new compound are boron atoms arranged in graphene-like layers, molybdenum layers, and triangles of boron atoms. Boron and molybdenum atoms are arranged in alternating layers with some Mo atoms replaced with B₃ triangles spread evenly throughout the crystal's



volume.



Various molybdenum boride crystal structures obtained using the USPEX algorithm Credit: Dmitry V. Rybkovskiy, J. Phys. Chem. Lett.

"We made a hypothesis that the structure of the highest boride is disordered and boron triangles statistically replace molybdenum atoms. In order to prove that, we developed a lattice model that enabled us to define the rules that governed how the boron units should position themselves to achieve the lowest energy," says Dmitry Rybkovskiy, research fellow at Skoltech and A. M. Prokhorov General Physics Institute and the first author of the work.

The brute-force search for the positions of molybdenum atoms and boron triangles, sampling different variants, revealed the patterns related to stability. Stable phases contain four to five <u>boron atoms</u> per one metal atom, and $MoB_{4.7}$ is the most stable of such compounds and gives the best match to the experimental XRD pattern.

"This study is an interesting example of interaction between theory and experiment. Theory predicted a compound that demonstrates peculiar properties and new structure, but the experiment suggested that the



actual material is more complex and its structure is partially disordered. The theory we formulated based on these findings allowed us to reproduce all experimental observations and understand the exact composition and structure of this material, as well as its detailed properties. In particular, the calculated hardness is close to the range of superhard materials," said Artem Oganov, professor at Skoltech and MIPT, and the leader of the team of authors.

Superhard materials have a wide range of industrial applications, such as machine tool building, jewelry, or mining. They are used in cutting, polishing, grinding, drilling, so the search for new superhard compounds is an important task.

More information: Dmitry V. Rybkovskiy et al, Structure, Stability, and Mechanical Properties of Boron-Rich Mo–B Phases: A Computational Study, *The Journal of Physical Chemistry Letters* (2020). DOI: 10.1021/acs.jpclett.0c00242

Alexander G. Kvashnin et al. Computational discovery of hard and superhard materials, *Journal of Applied Physics* (2019). DOI: 10.1063/1.5109782

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