

## Accelerating the discovery of new materials via the ion-exchange method

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(Upper) Schematic illustration of the high-temperature synthesis method normally used for material synthesis and (bottom) the ion-exchange method suitable for synthesizing metastable phases. Credit: Issei Suzuki



Tohoku University researchers have unveiled a new means of predicting how to synthesize new materials via the ion-exchange. Based on computer simulations, the method significantly reduces the time and energy required to explore for inorganic materials.

Details of their research were <u>published</u> in the journal *Chemistry of Materials* on April 17, 2024.

In the quest to form new materials that facilitate environmentally friendly and efficient energy technologies, scientists regularly rely on the high temperature reaction method to synthesize <u>inorganic materials</u>. When the raw substances are mixed and heated to very high temperatures, they are split into atoms and then reassemble into new substances. But this approach has some drawbacks. Only materials with the most energetically stable crystal structure can be formed, and it is not possible to synthesize materials that would decompose at high temperatures.

On the contrary, the <u>ion-exchange</u> method forms new materials at relatively low temperatures. Ions from existing materials are exchanged with ions of similar charge from other materials, thereby forming new inorganic substances. The low synthesis temperature makes it possible to obtain compounds that would not be available by the usual high <u>temperature</u> reaction method.





A picture of the prediction of the availability of ion exchange. Credit: Issei Suzuki

Despite its potential, however, the lack of a systematic approach to predicting appropriate material combinations for ion exchange has hindered its widespread adoption, necessitating laborious trial-and-error experiments.

"In our study, we predicted the feasibility of materials suited for ion



exchange using <u>computer simulations</u>," says Issei Suzuki, a senior assistant professor at Tohoku University's Institute of Multidisciplinary Research for Advanced Materials, and co-author of the paper.

The simulations involved investigating the potential for ion exchange reactions between ternary wurtzite-type oxides and halides/nitrates. Specifically, Suzuki and his colleagues performed simulations on 42 combinations of  $\beta$ -M<sup>I</sup>GaO<sub>2</sub>, M<sup>I</sup> = Na, Li, Cu, Ag as precursors, and halides and nitrates as ion sources.

The <u>simulation results</u> were divided into three categories: "ion exchange occurs," "no ion exchange occurs," and "partial ion exchange occurs (solid solution is formed). To confirm their results, the researchers verified the simulation through actual experiments, confirming an agreement between simulation and experiments in all 42 combinations.

Suzuki believes that their advancement will accelerate the development of new materials suitable for improved energy technologies. "Our findings have shown that it is possible to predict whether ion exchange is feasible and to design reactions in advance without experimental trial and error. In the future, we plan to use this method to search for materials with new and attractive properties that will tackle energy problems."

**More information:** Issei Suzuki et al, Designing Topotactic Ion-Exchange Reactions in Solid-State Oxides Through First-Principles Calculations, *Chemistry of Materials* (2024). <u>DOI:</u> <u>10.1021/acs.chemmater.3c03016</u>



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