

## **Exploration of Next-Generation Solar Cell Materials**

## Using the AI analysis platform Multi-Sigma, we will construct a prediction model of formation energy and bandgap based on material descriptors. The models were linked for multi-objective optimization to identify compounds with promising properties for solar cell applications.

## 1. Formation Energy and Bandgap Prediction

Using Multi-Sigma's prediction function, we trained models to map material descriptors to properties. Model 1 used electronegativity, period, group, density, and volume to predict formation energy, while Model 2 used electronegativity, period, group, and formation energy to predict the bandgap.



Formation Energy factor analysis



## 3. Multi-Objective Optimization

Multi-Sigma's optimization function suggests the optimal parameter combinations to achieve target performance metrics.

- · Minimization of formation energy (ensuring long-term stability)
- Achieve a bandgap of around 1.5 eV (optimal light absorption characteristics relative to the sunlight spectrum)

Multi-objective optimization with Multi-Sigma yielded a promising solar cell material candidate with the following physical properties.

Formation energy	Band Gap
-3.754 eV/atom	1.528 eV

Data source: https://next-gen.materialsproject.org/api

AlZOTH inc. provides a range of Al services, including Multi-Sigma, Al consulting, experimental condition optimization support, and contract research and development. Multi-Sigma is a cloud-based Al software designed for research and development, significantly reducing experimental workload and enabling researchers to discover innovative solutions to realworld challenges with minimal experimental datasets (A) dep pund dep pund

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