A universal model for accurately predicting the formation energy of inorganic compounds

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ABSTRACT Harnessing recent advances in data science and materials engineering, it is feasible today to build reliable models for predicting materials properties. Here we employ a comprehensive dataset of 170,714 inorganic crystalline compounds obtained from high-throughput accurate quantum mechanics calculations, to train a machine learning model for the precise prediction of the formation energy of inorganic compounds. Distinct from previous studies, our model can be universally applied to a large phase space of inorganic materials as all the data is utilized for the training, and the model reaches a fairly good predictive ability (R² = 0.982 and mean absolute error = 0.072 eV atom⁻¹, DenseNet model). The improvement comes from several effective structure-dependent descriptors, which are carefully designed to take into account the information of the electronegativity difference between neighboring atoms and local atomic structure. This model provides a useful tool to predict the energy landscape of the compound systems in a fast and cost-effective manner.

Keywords: machine learning, formation energy, electronegativity

INTRODUCTION

The formation energy (Eform) of crystals, i.e., the energy to bind atoms together to form condensed phase of matters, is one of the most important physical properties of a material. The Eform represents the strength of the adhesion of atoms to each other within a material system, and is responsible for many thermo-dynamic- and kinetic-related properties, such as stability [1] and synthesizability [2] of a compound. The value of the Eform can be obtained from the first-principles calculations, which are relatively computation-intensive and expensive. Harnessing the advances of computational materials science databases, as well as the knowledge of data science, the Eform prediction can now be achieved in an easier and more effective way.

Previously, attempts have been made to quickly gauge the Eform of inorganic materials. For example, Cao et al. [3] utilized a convolution neural network (CNN) model and mixed Magpie descriptors [4] and orbital-field matrix (OFM) descriptors [5] as input for predicting the Eform using data of more than 4000 crystalline materials including transition metal binary alloys, lanthanide metal and transition metal binary alloys. The prediction of Eform achieved a mean absolute error (MAE) of 70 meV atom⁻¹. Ye et al. [6] used the Pauling electronegativity [7,8] and ionic radii [9] of the constituent species as the input descriptors with artificial neural networks (ANNs), and obtained a model with extremely low MAEs of 7–10 and 20–34 meV atom⁻¹ in predicting the Eform of garnets and perovskites, respectively. Ward et al. [10] built a machine learning (ML) decision tree model by training with a dataset of inorganic compounds taken from the Open Quantum Materials Database (OQMD) [11], and achieved an MAE of 80 meV atom⁻¹ in cross validation for predicting Eform by employing structural descriptors from the Voronoi tessellation of the structure of crystalline. Xie and Grossman [12] proposed a generalized graph convolutional neural networks (GGCNN) model for material property predictions, and their model for the Eform prediction reached an MAE of 39 meV atom⁻¹ with the dataset of 28,046 samples taken from the Materials Project [13]. Li et al. [14] proposed to use deep neural network-based transfer learning and a set of hybrid descriptors for perovskite Eform prediction [15].

If we look at the existing models closely, nearly all the published papers trim their training dataset in some way, and those treatments would for sure improve the predicting ability. For example, Ye et al. [6] limit their model to the Ca₂Al₂O₆ garnets and ABO₃ perovskites; the GGCNN paper [12] states that only 28,000 of compounds are used out of ~130,000 data points from Materials Project, making their model not generally applicable, especially for those removed structures; the Roost model [16] uses the “subset contains only the lowest energy polymorph for each stoichiometry” from OQMD database, meaning their model works for the stable compounds only.

Fig. 1 shows the actual predictive power of the artificial intelligence (AI) models by plugging in the entire dataset without data cleaning. All these models are reproduced using the codes, parameters, and data treatment as stated in the original papers, and all the models work as well as they are demonstrated in the paper for the cleaned dataset. But if we incorporate the entire dataset (139,368 structures from Materials Project) as the input to validate the model, the predicting power of the models becomes awful. For example, The MAE of the GGCNN model increases from 39 meV atom⁻¹ to 0.137 eV atom⁻¹.

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It is indeed a common issue for AI research that the ML model can be tuned with biased data to make it looking good, but it hurts the ability of extrapolation of the model. It suggests that, when we judge the AI models, both the predicting accuracy and the ability of extrapolation matter, and hence both should be validated to make sure the model is feasible and generally applicable. In this work, we build an AI model for predicting the $E_{\text{form}}$ of inorganic compounds, and the model has an improved capability of extrapolation compared with several existing models.

Evidence has been demonstrated that AI models can advance materials science significantly [17], e.g., accelerating the materials discovery of energy materials [18,19] and incorporating expert knowledge into the materials design [20]. There are three important things for building a good and reasonable ML model: (1) good data and more data (shown in Fig. S1); (2) good ML algorithm that can react to or pass over the information effectively between neurons; (3) improved descriptors those can extract the intrinsic connections between the properties. In this work, (1) we do not perform any data “cleaning” trick to purposely boost the model predictive power, rather the entire dataset is included for the training process to make sure the model is as universal as possible to cover a large configuration space of inorganic compounds; (2) a DenseNet (DN) algorithm is employed to make sure the neurons are directly and effectively connected; (3) to boost the model accuracy, it is found that the electron negativity difference and structural information are indeed the most important descriptors for determining the $E_{\text{form}}$ of compounds, and the prediction performance of $E_{\text{form}}$ is greatly improved after the addition of these descriptors. Hence, our ML model can reach a fairly good predictive ability up to $R^2 = 0.982$ and $\text{MAE} = 0.072$ eV atom$^{-1}$ for all the compounds.

**METHODS**

Models

Eight conventional ML models as well as a neural network model are adopted for predicting the density functional theory (DFT)-calculated formation energy per atom ($E_{\text{form}}$), which are AdaBoost Regression (ABR), Linear Support Vector Regression (LSVR), SVR, Ridge Regression (RR), Gradient Boosting Regression (GBR), K-Nearest Neighbors Regression (KNNR), Random Forest Regression (RFR), ExtraTrees Regression (ETR), and DN [21,22]. In this study, the commonly used ML models are implemented by scikit-learn [23] and the DN is implemented by pytorch [24]. DN is a neural network model, which has three hidden dimensions with node counts of [256, 128, 64], in addition to an input layer with 119 nodes and an output layer with 1 node. The DN model is trained using the MAE loss criterion, or called $L1$ loss criterion, shown in Formula (1) and an Adam optimizer with a learning rate of $10^{-3}$.

$$L1 = |f(x) - Y|,$$

where $f(x)$ represents the predicted value and $Y$ represents the actual DFT-calculated value retrieved from the Atomly.net materials-data infrastructure. The predictive performances of models are described as variance ($R^2$), MAE, root mean squared error (RMSE).

The training of the AI models in this work usually takes ~20 h on a single NVIDIA Tesla V100 card and the prediction of the formation energy for single structures takes less than 1 s.

**Descriptors**

In order to obtain a useful AI model, several descriptors are constructed for describing a crystal material. In general, there are two distinct types of descriptors: composition-dependent (CD) descriptors and structure-dependent (SD) descriptors. The latter incorporate the atomistic structure of a material, hence making the model geometry-sensitive. The CD descriptors are semantically extracted from the chemical formula of a crystal material based on the elemental properties, like the descriptors proposed by Magpie [4] and Oliynyk [25]. In this study, 44 different elemental properties (listed on Table S1) have been chosen for generating CD descriptors. To each elemental property, six kinds of statistical quantities (average, variance, range, skewness, kurtosis, and sum) are derived as CD (CD) descriptors and structure-dependent (SD) descriptors. The construction of a crystal material. In general, there are two distinct types of descriptors: composition-dependent (CD) descriptors and structure-dependent (SD) descriptors. The latter incorporate the atomistic structure of a material, hence making the model geometry-sensitive. The CD descriptors are semantically extracted from the chemical formula of a crystal material based on the elemental properties, like the descriptors proposed by Magpie [4] and Oliynyk [25]. In this study, 44 different elemental properties (listed on Table S1) have been chosen for generating CD descriptors. To each elemental property, six kinds of statistical quantities (average, variance, range, skewness, kurtosis, and sum) are derived as CD descriptors, ending up with 264 CD descriptors.

For the SD descriptors, beside density and band gap, we also propose some new/in-house SD descriptors related to the chemical bond (CB), coordination number (CN), and electron negativity difference ($\Delta \chi$), which are all extracted from the local environment of each atom in a structure. The density is calculated from the crystal structure via the Pymatgen [19], and the band gap was directly retrieved from the “atomly.net” database.
The local environment of an atom represents the interactions between the atom and its nearest neighbors, and directly affects the $E_{form}$ of compounds. The nearest neighbors of each atom in a structure are extracted by using the Voronoi tessellation method [10], which has also been used for generating many other descriptors related to the structure, e.g., the aforementioned OFM descriptors [5]. By using this method, the real space can be divided into a number of convex polyhedrons by constructing vertical bisectors between neighboring atoms, similar to the Wigner-Seitz cell, as shown in Fig. 2a. One can see that there is only one atom in each polyhedron, whose nearest neighbors are represented by the surface of the polyhedron.

Similar to the CD descriptors, we also use six statistical quantities to generate the new/in-house SD descriptors. That is to say, there are six CB SD descriptors ($CB^a, CB^b, CB^c, CB^d, CB^e, CB^f$) which can be calculated as follows:

$$CB^a = \frac{1}{N} \sum_{j=1}^{N} CB^a_j,$$

$$CB^b = \left[ \frac{1}{N} \sum_{j=1}^{N} \left( CB^b_j - \frac{1}{N} \sum_{j=1}^{N} CB^b \right)^2 \right],$$

$$CB^c = \max(CB^a_j) - \min(CB^a_j),$$

$$CB^d = \left( \frac{1}{N} \sum_{j=1}^{N} \left( CB^d_j - \frac{1}{N} \sum_{j=1}^{N} CB^d \right) \right)^{\frac{3}{2}},$$

$$CB^e = \left( \frac{1}{N} \sum_{j=1}^{N} \left( CB^e_j - \frac{1}{N} \sum_{j=1}^{N} CB^e \right) \right)^{\frac{4}{3}},$$

$$CB^f = \sum_{j=1}^{N} CB^f_j,$$

where $N$ is the number of atoms in the structure and $CB$ is the average of the chemical bonds of the $i^{th}$ atom with its nearest neighbors. As shown in Fig. 2b, the $CB$ between the $i^{th}$ atom and its $j^{th}$ nearest neighbor is expressed in the spherical coordinates as $(R, \theta, \phi)$, which is represented by the yellow vector $CB_{ij}$. And $CB$ is defined by

$$CB^a_j = \frac{1}{n_i} \sum_{j=1}^{n_i} CB^a_{ij} = \frac{1}{n_i} \sum_{j=1}^{n_i} (R, \theta, \phi)_{ij},$$

where $n_i$ represents the number of neighbors of the $i^{th}$ atom. Fig. 2c shows the coordination number $CN_i$ of the $i^{th}$ atom counted via the Voronoi tessellation. The six CN SD descriptors ($CN^a, CN^b, CN^c, CN^d, CN^e, CN^f$) related to statistical quantities are calculated with the formulas similar to those of the CB SD descriptors.

The strength of a chemical bond should depend on the electronegativity difference ($\Delta \chi$) between the two bonding atoms. Usually, for ionic compounds, the bigger the electronegativity difference, the stronger the bond. The $\Delta \chi$ between atoms bonded together will lead to the charge transfer from one to another, greatly affecting the charge density distribution of the local structure, and thereby the $E_{form}$ of a system will be affected because the total energy would be a functional of charge density [26]. Therefore, the $\Delta \chi$ is essential to the prediction of $E_{form}$ and thus we propose several SD descriptors of $\Delta \chi$. Fig. 2d describes the electron distribution caused by the $\Delta \chi$ bonding atoms. Here, we have chosen five different electronegativity definitions, Allred and Rochow electronegativity (AR $\chi$), Pauling electronegativity (Pa $\chi$), Martynov and Batsanov electronegativity (MB $\chi$), Gordy electronegativity (Go $\chi$) and Mullikan electronegativity (Mu $\chi$), to generate descriptors related to the $\Delta \chi$. To each electronegativity definition, the SD descriptors are calculated by the following steps: firstly, the absolute value of the $\Delta \chi$ between the $i^{th}$ atom and its $j^{th}$ neighbor ($\Delta \chi_{ij}$) is given by

$$\Delta \chi_{ij} = |\chi_i - \chi_j|,$$

where $\chi_i$ and $\chi_j$ are the electronegativity of the $i^{th}$ atom and its $j^{th}$ neighbor, respectively. Then, the $\Delta \chi^a$ of the $i^{th}$ atom with its neighbors are calculated as the calculation as follows:

$$\Delta \chi^a_i = \frac{1}{n_i} \sum_{j=1}^{n_i} \Delta \chi_{ij},$$

where $n_i$ denotes the number of neighbors of the $i^{th}$ atom. Finally, to each electronegativity definition, the six electronegativity difference SD descriptors ($\Delta \chi^a, \Delta \chi^b, \Delta \chi^c, \Delta \chi^d, \Delta \chi^e, \Delta \chi^f$) for the whole structure are calculated similarly to the aforementioned calculation method of the $CB$ SD descriptors.

In the end, we got another set of descriptors which consists of the 234 CD descriptors and the 44 SD descriptors based on 9 different structure-related properties as detailed in Table S2. The 30 CD descriptors with respect to 5 kinds of electronegativity definitions are removed and replaced by the corresponding SD descriptors related to the electronegativity definitions. Although the treatments to build the descriptors, such as the Voronoi tessellation, are well-known, those descriptors of electronegativity difference in this work are the newly invented, and the following paper will show that they are fairly important.

Data and data availability
The dataset used in this work is obtained from the “atomly.net”...
[27], which is an access DFT database. The details of the data can be found in Supplementary information. The codes that are used to generate and validate the models can be found at https://github.com/atomly-materials-research-lab/Descriptor.

RESULTS AND DISCUSSION

Nine ML models are built to predict $E_{\text{form}}$, and their performance are benchmarked by comparing their cross-validated coefficients of determination ($R^2$), RMSE, and MAE values of the test dataset, which is the entire database without data cleaning (the optimized parameters of the eight conventional ML models are listed in Table S3). The top 20 important descriptors are screened out by using the RFR method, and the correlations between each descriptor pair are studied by the Pearson correlation coefficient.

It is found that the newly proposed SD descriptor along with the entire dataset (without data cleaning) can significantly optimize the models, making it much better than the models in Fig. 1. According to the validation results illustrated in Fig. 3, the DN model (Fig. 3i) gives the best performance with $R^2 = 0.982$, while the RR method (Fig. 3a) has the lowest prediction accuracy with $R^2 = 0.858$. Other four classical ML models, GBR (Fig. 3e), ABR (Fig. 3f), ETR (Fig. 3g), and SVR (Fig. 3h), can also achieve a very high prediction accuracy of $E_{\text{form}}$ with $R^2 > 0.95$. For the two tree-based models, the ETR with $R^2 = 0.961$ is better than the RFR (Fig. 3d) with $R^2 = 0.945$. As most classical boosting algorithms, both ABR (with extremely randomized trees as the weak learners) and GBR (with decision trees as the weak learners) can implemented very high $R^2 = 0.957$ and 0.952, respectively. The $R^2$ value of the KNNR model is close to 0.9 as illustrated in Fig. 3c. Comparing Fig. 3b and h, it is found that using the SVR model with the nonlinear RBF (radial basis function) kernel function can get a better prediction performance than that with the linear kernel function.

With the objective to estimate the effect of the new SD descriptors proposed in this study on predicting the $E_{\text{form}}$ of...
crystals, we investigated the changes in evaluation metrics ($R^2$, RMSE, and MAE) of the ML models with and without the SD descriptors. As shown in Fig. 4 and Fig. S2, the prediction performances of $E_{\text{form}}$ for all the nine ML models have been improved by increasing the SD descriptors, which indicate that the local structure information describes $E_{\text{form}}$ very well for the inorganic materials. Fig. 4a depicts that the increase rate of $R^2$ values varies from 6.3% to 17.0% in combination of the SD descriptors with the CD ones. The SD descriptors dramatically improve the $R^2$ value of the ABR model for $E_{\text{form}}$ from 0.829 to 0.957. For DN model, which has the highest predictive power, the $R^2$ value increase from 0.885 to 0.982, which is 11.0% increase. The changes of RMSE and MAE values are shown in Fig. 4b, c. One can see that the decreasing rates of RMSE and MAE values fluctuate in the range of [20.5%, 60.4%] and [10.0%, 50%], respectively. These results prove that the newly proposed SD descriptors, in particular the electronegativity difference of neighboring atoms, can significantly improve the prediction performances of $E_{\text{form}}$ in the inorganic materials.

It is also observed that the electronegativity difference, unlike the average electronegativity employed in previous studies, indeed improves the model significantly. To analyze the control factors of $E_{\text{form}}$, the RFR method is chosen for ranking the importance of different descriptors, since the RFR method is a tree-based learning algorithm and has advantages on both accuracy and interpretability [28]. Fig. 5 shows the top 20 descriptors ranked for $E_{\text{form}}$ prediction, and the correlation between descriptor pair in the way of Pearson correlation coefficient matrix. It is found that the Pauling electronegativity is the most important factor for the prediction model of $E_{\text{form}}$ without and with SD descriptors (47.1% of Pa $\chi$ in Fig. 5a and 46.4% of $\Delta$Pa $\chi$ in Fig. 5b, respectively). Although the ratio of $\Delta$Pa $\chi$ slightly drops when using the SD descriptors, the overall ratio of the top 20 important descriptors increases from 74.6% to 82.6%. In Fig. 5a, b, there are five descriptors (Pa $\chi$, Pa $\chi^2$, AR $\chi^2$, AR $\chi^4$, Mu $\chi^2$) related to electronegativity and two descriptors ($\Delta$Pa $\chi$, $\Delta$AR $\chi^2$) related to “electronegativity difference” appearing in the top 20 important descriptors, respectively. Metallic valence represents the oxidation state of the metal in alloy similarly to that of covalent compound. In the previous studies for predicting the thermodynamic stability of perovskites, the most common oxidation state is one of the most important descriptors. The three descriptors ($MV^s$, $MV^f$, $MV^r$) related to the metallic valence [29,30] appear in the 20 important descriptors in both Fig. 5a, b, respectively. Fig. 5b illustrates that the density ($\rho$), coordination number ($CN^a$), band gap ($E_g$) are also important descriptors for improving the prediction performance of $E_{\text{form}}$, which is consistent with the previous results [14,31,32]. In the prior study of Ward et al. [4], it is pointed out that the $E_{\text{form}}$ of intermetallic compounds is best described by the variances in the melting point ($MP^v$) and number of d electrons between constituent elements ($V_d^v$). The similar result is also observed in this study, because there exist linear relationships between the melting point and boiling point. As shown in Fig. 5a, b, the variance of the boiling point ($BP^v$) as well as the variance of d electrons and unfilled d electrons ($V_d^v$, $UV_d^v$) appear in the top 20 important descriptors. The variance of cohesive energy, the skewness and kurtosis of the 1st ionization

Figure 4  (a) $R^2$, (b) RMSE, and (c) MAE values of the nine different ML methods for $E_{\text{form}}$ prediction with and without the new proposed SD descriptors. The red and blue bars represent the corresponding values before and after adding these descriptors, respectively.
potential (CE, 1st IP, 1st IP), also appear in the top 20 important descriptors.

Fig. 5c, d show the Pearson correlation matrix for the top 20 important descriptors without and with the SD descriptor set. For each pair of descriptors, A and B, the correlation coefficient $R_{AB}$ is defined as

$$R_{AB} = \frac{\sum_{i=1}^{n} (A_i - \bar{A})(B_i - \bar{B})}{\sqrt{\sum_{i=1}^{n} (A_i - \bar{A})^2} \sqrt{\sum_{i=1}^{n} (B_i - \bar{B})^2}}$$

where $\bar{A}$ and $\bar{B}$ are the sample means of A and B descriptors over the total n crystal materials, while $A_i$ and $B_i$ are the descriptors of the ith material. As shown in Fig. 5c, strong correlations are found in several pairs of descriptors: (1) two descriptors related to the $d$ valence electrons orbitals (UV_d, V_d); (2) three descriptors related to the electronegativity (Pa $\chi$, AR $\chi$, Mu $\chi$); (3) two descriptors related to the thermodynamic property (BP, HA). Comparing Fig. 5c, d, the changes in the color distributions at the lower right corner (marked by the black rectangles) show that the correlations of the top 20 descriptors decrease obviously by adding the SD descriptors. Our results show that there is a correlation between different definitions of electronegativity. As shown in Figs S3 and S4, when the most important descriptor $\Delta Pa \chi$ in Fig. 5b is removed from the collection of the input descriptors, the $\Delta AR \chi$ descriptor becomes the most important descriptor after ranking again by the RFR method. Repeatedly with the above steps, when removing $\Delta AR \chi$, the $\Delta AR \chi$ becomes the most important descriptor, which is consistent with the observed result in Fig. 5d that there exist strong correlations among the descriptors ($\Delta Pa \chi$, $\Delta AR \chi$) and the relation on prediction performance of $E_{form}$ follows the trend $\Delta Pa \chi > \Delta AR \chi > \Delta AR \chi$.

Then, the $E_{form}$ of compounds can be predicted with the constructed models. Fig. 6 shows the thermodynamic phase diagram for compounds in Ti-O, V-O, Mn-O and Li-P chemical systems obtained from the DN model and DFT. These four
chemical systems are selected to demonstrate the accuracy of the model for their importance, since TiO\textsubscript{x} materials are commonly used in photovoltaic devices to improve device performance \cite{33,34}, VO\textsubscript{x} is a promising cathode material and likely to be commercialized applicable in the future \cite{35}, MnO\textsubscript{x} is widely studied as catalysts \cite{36–38}, and LiP\textsubscript{x} is the intermedia product in certain anode reactions of Li-ion batteries \cite{39}. The thermodynamic phase diagrams are generated using the same energy correction mechanism as the Materials Project \cite{1,40}, to make sure the energy hulls in this work are comparable to either Materials Project \cite{1,40} or Atomly \cite{27} (also adopts the same energy correction for generating the phase diagram). As shown in Fig. 6a–d, The DFT data tell us that there are 6 stable crystalline compounds (Ti\textsubscript{6}O, Ti\textsubscript{3}O, TiO, TiO\textsubscript{2} and TiO\textsubscript{3}) in the Ti-O chemical system, 4 stable crystalline compounds (V\textsubscript{2}O\textsubscript{3}, V\textsubscript{3}O\textsubscript{5}, VO and VO\textsubscript{2}) and 2 unstable ones (V\textsubscript{6}O\textsubscript{13} and VO\textsubscript{3}) in the V-O chemical system, 4 stable crystalline compounds (MnO, Mn\textsubscript{3}O\textsubscript{4}, Mn\textsubscript{2}O\textsubscript{3} and MnO\textsubscript{2}) and 1 unstable crystalline compound (MnO\textsubscript{3}) in the Mn-O chemical system, and 4 stable crystalline compounds (Li\textsubscript{3}P, LiP, Li\textsubscript{3}P\textsubscript{7} and LiP\textsubscript{7}) in the Li-P chemical system, respectively. Note that the DN model can assess the $E_{\text{form}}$ of those compounds fairly well. It is seen from Fig. 6c that in the Mn-O chemical system, there is a distinct advantage of the model with the SD descriptors on predictive performance over that without the SD descriptors. The other panels in Fig. 6 show that both models, with or without the SD descriptors, can capture the shape of the energy hull to some extent. When the SD descriptors are taken into account in the model, the predicted $E_{\text{form}}$ is off only by ~135 meV atom\textsuperscript{-1} on average (ranging from 0 to 373 meV atom\textsuperscript{-1}) for the Ti-O chemical system, ~40 meV atom\textsuperscript{-1} (ranging from 0 to 102 meV atom\textsuperscript{-1}) for the V-O chemical system, ~33 meV atom\textsuperscript{-1} (ranging from 0 to 120 meV atom\textsuperscript{-1}) for the Mn-O chemical system, and ~11 meV atom\textsuperscript{-1} (ranging from 0 to 27 meV atom\textsuperscript{-1}) for the Li-P chemical system, compared with the DFT values. Hence, our model makes the prediction of $E_{\text{form}}$ of inorganic materials easy and accurate.

**CONCLUSIONS**

In summary, we develop a universal model for predicting the formation energy of compounds as all the data points are employed for training without the “data cleaning”. The “data cleaning”, as performed by many other literature, makes their model looking good, but worsens the capability of extrapolation of the model. We have also identified several new SD descriptors related to the electronegativity difference and coordination numbers for predicting the formation energy of a crystal material, derived from the local environment of the material via the Voronoi tessellation method. We demonstrate that these new descriptors can significantly improve the prediction accuracy of formation energy not only for eight classical ML methods but also for another neural network method. The neural network
model achieves the high prediction accuracy of $R^2 = 0.982$ and MAE = 0.072 eV atom$^{-1}$. The tree-based RFR model is chosen for screening the key descriptors for best describing the formation energy. It is found that the Pauling electronegativity difference between bonding atoms is the most important factor with a ratio of 46.4% for the prediction of formation energy. Our work shows that by adopting the physics-based descriptor as well as a good dataset, the predictive power of ML models can be significantly improved. There might be still a large room out there to keep enhancing the predictive power, if we can find better descriptors with better data. We expect that the model presented here can provide a useful tool for materials science community immediately towards extremely fast, large-scale, and accurate materials modelling and prediction without costly computations.

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Author contributions Liang Y and Liu M conceived the idea, designed the experiments, and wrote the paper with support from Meng S and Wang Z; Liang Y; Chen M, Wang Y; Jia H; Lu T; and Xie F prepared the data, built the database, wrote the machine learning code, and analyzed the training as well
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Conflict of interest  The authors declare that they have no conflict of interest.

Supplementary information  Supporting data are available in the online version of the paper.

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一种预测无机晶体形成能的高精度泛化模型

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摘要  随着数据科学和材料科学的进步，人们如今可构建出较为准确的人工智能模型，用于材料性质预测。本文中，我们以170,714个无机晶体化合物的高通量第一性原理计算数据集为基础，训练得到了可精确预测无机化合物形成能的机器学习模型。相比于同类工作，本项研究以超大数据集为出发点，构建出无机晶体形成能的高精度泛化模型，可外推至广阔相空间，其中的DenseNet神经网络模型精度可以达到R² = 0.982和平均绝对误差(MAE) = 0.072 eV atom⁻¹。上述模型精度的提升源自一系列新型特征描述符，这些描述符可有效提取出原子与领域原子间的电负性和局域结构等信息，从而精确捕捉到原子间的相互作用。本文为新材料搜索提供了一种高效、低成本的结合能预测手段。