

Review

Machine learning interatomic potential: Bridge the gap between small-scale models and realistic device-scale simulations

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SUMMARY

Machine learning interatomic potential (MLIP) overcomes the challenges of high computational costs in density-functional theory and the relatively low accuracy in classical large-scale molecular dynamics, facilitating more efficient and precise simulations in materials research and design. In this review, the current state of the four essential stages of MLIP is discussed, including data generation methods, material structure descriptors, six unique machine learning algorithms, and available software. Furthermore, the applications of MLIP in various fields are investigated, notably in phase-change memory materials, structure searching, material properties predicting, and the pre-trained universal models. Eventually, the future perspectives, consisting of standard datasets, transferability, generalization, and trade-off between accuracy and complexity in MLIPs, are reported.

INTRODUCTION

Materials science stands at the forefront of interdisciplinary research, bridging various scientific domains and exploring phenomena across multiple length scales, from atomic to macroscopic.^{1–7} The field focuses on how the composition, structure, and processing of materials influence their properties and performance, leading to significant advancements in developing and refining materials.^{8,9} The evolution of scientific paradigms from traditional empirical and model-based theoretical approaches to contemporary computational and data-driven methodologies has significantly elevated the role of full-scale simulation in the design and development of materials.^{10–16} The primary component of material simulations at any scale is the interatomic potentials, which define the potential energy surfaces (PES) of atomic interactions within a material system.^{17–19} These potentials are essential for accurately predicting and understanding the physical and chemical properties of materials, which play a pivotal role not just in materials science, but also in chemistry and condensed matter physics.^{20–29} The precise determination of these potentials is crucial for reliable molecular dynamics simulations and studying material behaviors under diverse conditions.^{30–47}

Traditionally, interatomic potentials have been derived from empirical methods, while computationally efficient methods often lack the necessary accuracy and transferability for complex systems, such as the embedded-atom method (EAM),⁴⁸ empirical N-body potential,⁴⁹ Tersoff,^{50–52} and so on. First-principles methods, such as Density Functional Theory (DFT), offer higher accuracy but at a significant computational cost, limiting their applicability to smaller systems or shorter timescales.^{53,54}

The advent of machine learning (ML) has opened new horizons in the field of interatomic potentials.⁵⁵ Machine learning, with its ability to learn from and make predictions or decisions based on data, presents a promising alternative to traditional methods.^{56–60} Therefore, Behler and Parrinello proposed an artificial neural network⁶¹ for MLIP with symmetry function descriptors,⁶² effectively applied to silicon,^{63–65} vinyl bromide,⁶⁶ virtual reality,⁶⁷ Ge-Te binary compounds,⁶⁸ surface diffusion of CO/Ni,⁶⁹ Ge-Sb-Te ternary alloys,⁷⁰ synthetic chemists,⁷¹ multi-scale-shock dynamics simulations,⁷² and many transition-metal oxide compositions.^{73,74} Meanwhile, Csanyi et al. also developed the Gaussian approximation potential⁷⁵ (GAP) using SOAP descriptors,⁷⁶ applied to amorphous carbon⁷⁷ and silicon.⁷⁸ These two methods now also are the main technologies of MLIPs.^{79–94} Previously, there are some review articles on MLIPs, for example, Behler et al.^{95–97} outlined the timeline for the evolution of neural network potentials by classifying schemes for MLIPs into four generations; Deringer et al.⁹⁸ introduced the basic principles of MLIPs and highlighted the applications to some select problems in materials science, consisting of phase-change materials for memory devices, nanoparticle catalysts, and carbon-based electrodes for chemical sensing, supercapacitors, and batteries; Frieserich et al.⁹⁹ summarized the underlying machine learning methods, the data acquisition process and active learning procedures for MLIPs; Unke et al.¹⁰⁰ gave chemical insights into MLIPs and describe a step-by-step guide for constructing and testing them from scratch. In this review article, we highlight the process for constructing MLIPs from the perspective of data, consisting of the high-throughput generation of materials data, conversion of structural data into descriptors, machine learning models for data training, and the application of the

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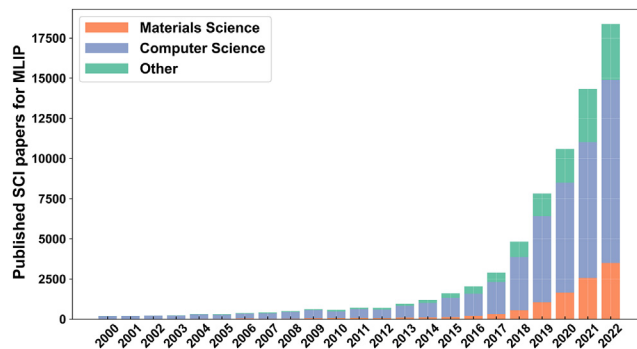


Figure 1. The number of published scientific articles on the topic of MLIP

The original data is queried by the keyword of machine learning potential, machine learning force field, and MLIP from Web of Science from 2000 to 2022, as indexed in the Science Citation Index (SCI).

model to new data. In particular, in the model section, we divided the models into six categories based on different ML algorithms and provided executable code for each method.

In detail, this review focuses on the fundamentals, flowcharts, software, applications, and challenges of MLIPs. In section [Fundamental and flowchart of machine learning interatomic potential](#), we introduce the development tendency and the entire key flowcharts related to MLIPs. The data collection methods, material structure descriptor, ML algorithm, and software are presented in sections three to five. The sixth section outlines the application of MLIPs in the selection of phase-change memory materials, structure searching, material properties prediction, and the development of pre-trained universal models. Finally, it looks forward to the challenges and opportunities of MLIPs in the future.

FUNDAMENTAL AND FLOWCHART OF MACHINE LEARNING INTERATOMIC POTENTIAL

From the year 2000–2022, the number of articles published in MLIP has increased across various fields, as shown in [Figure 1](#). Initially, MLIP research was predominantly rooted in the field of computer science, in which the earliest articles were published.⁶⁹ The trend in Computer Science has seen a consistent rise over the years, reflecting the increasing relevance and application of MLIP algorithms in this field.⁹⁷ For materials science, there were hardly any MLP publications before 2007. However, there has been a noticeable increment in recent years with the development of supercomputing power and the wave of the AI technology revolution.^{101–108}

The training process of machine learning models involves several key steps, as shown in [Figure 2](#). In this part, we focus on introducing the issues that need to be identified at each step.

- Data Collection: the first step is collecting the abundant and necessary data. This involves focusing on the problem to be solved and then gathering all relevant data that can be used to train the model. The data should be as representative as possible of the problem domain and diverse enough to capture all possible conditions.
- Materials Descriptors: once the data is collected, the most important step, which has the greatest impact on the accuracy of the MLIP, is how to convert spatial configurations into a machine learning dataset, also known as the material descriptors. The material descriptor determines the quality of the initial dataset for machine learning and also defines the highest accuracy that the MLIP can achieve. Different machine learning algorithms are merely used to infinitely approximate this optimal goal value. Afterward, several steps such as data cleaning, preprocessing, and normalization are also indispensable. Data cleaning can remove any irrelevant or duplicate data, while preprocessing involves techniques such as scaling or encoding to make the data suitable for training. Normalization is a process of adjusting the data to a common scale for better convergence during training.
- Model Selection: it is essential to choose a model that suits the nature of the problem and has the capacity to learn from the given data. Common models used for MLIP include Gaussian approximation, neural networks, and active learning. Then, it is trained on the prepared data, which involves optimizing the model's parameters using a suitable optimization algorithm such as gradient descent or Adam and so on. The model iteratively updates its parameters based on the feedback provided by the loss function, which measures how well the model is performing on the training data. A crucial aspect of model training is tuning hyperparameter, which are used to control the learning process of the model, such as learning rate, batch size, or the number of hidden layers in a neural network. Common techniques for hyperparameter tuning include grid search, random search, or more advanced methods such as Bayesian optimization or gradient-based optimization. After training, it is essential to evaluate the model's performance on a separate test dataset that was not used during training. This helps in gauging how well the model generalizes to unseen data and provides an accurate estimate of its future performance in real-world scenarios. Evaluation metrics such as accuracy, precision, recall, or F1-score are commonly used to assess the model's performance.
- Model improvement and deployment: based on the evaluation results and feedback from domain experts, further improvements can be made to the model. This might involve modifying the architecture of the model, trying different hyperparameter settings, or incorporating more advanced techniques such as ensemble methods or transfer learning to improve performance. While the model

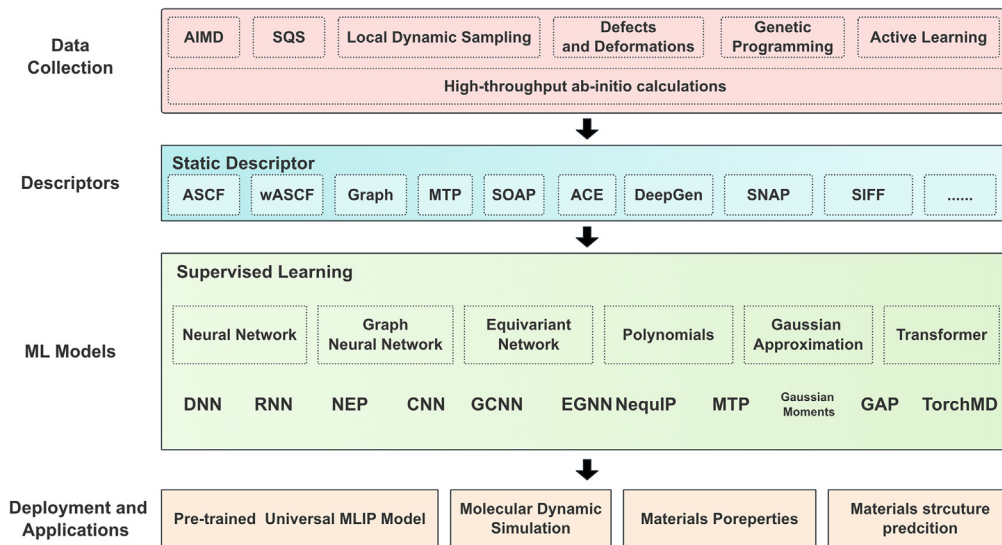


Figure 2. The flowchart and four essential stages of MLIPs

achieves satisfactory performance, it can be deployed in a production environment for real-time predictions or decision-making. It is essential to have a well-designed deployment strategy that ensures scalability, fault tolerance, and monitoring of the deployed model's performance over time.

DATA COLLECTION OF MACHINE LEARNING INTERATOMIC POTENTIALS

Datasets are fundamental to machine learning, providing the raw material from which algorithms learn and derive insights.^{109–111} Diverse datasets help in building robust models that can generalize well across different scenarios, reducing biases and improving the accuracy of predictions. In materials science, collecting comprehensive datasets through various methods is vital for exploring the entire spectrum of material properties and behaviors.^{112–116} This extensive data collection enables the identification of novel materials with desired characteristics, supports the understanding of complex material interactions, and facilitates the prediction of material performance under different conditions. In this part, we introduce some possible data collection methods for training MLIPs.

- Open-source material databases: the concepts of materials genomics have led to the creation of numerous large-scale materials databases globally, such as ICSD,¹¹⁷ Materials Project,¹¹⁸ Aflow,¹¹⁹ Materials Cloud,¹²⁰ NOMAD,¹²¹ ALKEMIE MatterDB,^{122–124} OQMAD,¹²⁵ COD,¹²⁶ OMD,¹²⁷ C2DB,¹²⁸ MatNavi,¹²⁹ among others. These databases provide a wealth of data obtained from DFT calculations, which typically require extensive computational resources. Therefore, efficiently filtering and selecting target materials from these open-source databases presents a highly effective method for gathering relevant data for MLIPs without the need for time-intensive DFT calculations.^{130–133}
- Ab Initio Molecular Dynamics (AIMD) Sampling: AIMD simulations to explore the configurational space. The temperature of the simulation determines the regions of the potential energy surface (PES) and energy ranges explored. This technique is suitable for equilibrium or near-equilibrium properties, such as studying vibrational spectra or thermodynamic properties.
- Adaptive Sampling or On-the-Fly ML: this technique starts with a small initial set of reference data to train a preliminary ML potential, which is then used in MD simulations.¹³⁴ Additional conformations are collected when model predictions become unreliable, based on an uncertainty criterion, and reference calculations are performed.^{135,136}
- Meta dynamics Sampling: similar to adaptive sampling, this method uses preliminary ML potentials in MD simulations but biases the dynamics to visit unexplored regions on the PES.^{137–139} It combines metadynamics with uncertainty estimates to select relevant structures.
- Normal Mode Sampling: this approach does not require MD simulations. It starts from a minimum on the PES and generates distorted structures by displacing atoms along normal modes. It is efficient for exploring PES but is limited to regions close to minima and is best combined with other sampling methods.

After collecting data, it is crucial to clean data by feature engineering, including removing inconsistencies, addressing missing values, and filtering out irrelevant information.^{140–144} Subsequently, data splitting segregates the refined dataset into training, validation, and test sets, which are used to assess the model's generalization capability while mitigating the risk of overfitting. Concurrently, data standardization and normalization emerge as pivotal steps in preprocessing. Specifically, standardization is the best method to rescale the data to yield a mean of zero and a standard deviation of one, effectively resolving issues arising from differing scales among features. Normalization, often achieved

Table 1. The various descriptors list for materials structure

Description	Year	Author
Atom-centered symmetry functions (ACSF)	2011	Behler ⁶²
Coulomb Matrix	2012	Rupp ¹⁵²
Smooth Overlap of Atomic Positions (SOAP)	2013	Csányi ⁷⁶
Ewald sum and Sine Matrix	2015	Faber ¹⁵³
Spectral neighbor analysis method (SNAP)	2015	Thompson ¹⁵¹
Moment Tensor Potentials (MTP)	2016	Shapeev ¹⁷⁷
Weighted atom-centered symmetry functions (wACSF)	2018	Gastegger ¹⁵⁰
Automatic selection of fingerprints	2018	Giulio ²²⁰
Atomic-position independent material descriptor (U-api)	2018	Thompson ¹⁵⁷
Deep Potential Dynamic Descriptor (DP)	2018	Zhang ¹⁵⁸
Graph Descriptor	2018	Jeffrey ¹⁵⁹
Optimizing SOAP	2019	Caro ²²¹
Atomic Cluster Expansion (ACE)	2019	Drautz ¹⁵⁶
Many-body Tensor Representation (MBTR)	2022	Huo ¹⁵⁴

by scaling data within a specific range such as 0 to 1, guarantees that all input features contribute uniformly to model training. In a word, data preprocessing, consisting of feature engineering, data standardization, and normalization, enhances the model's overall performance and stability.^{145–149}

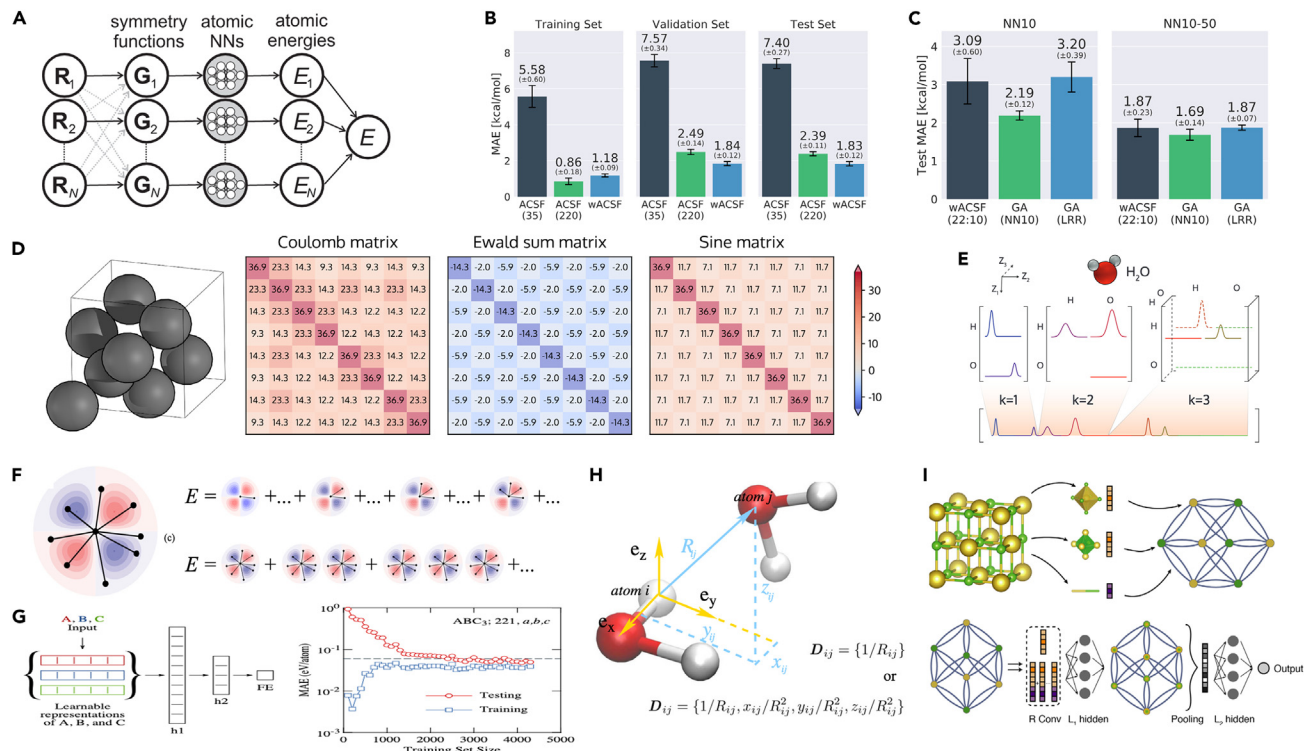
MATERIALS DESCRIPTORS OF MACHINE LEARNING INTERATOMIC POTENTIALS

To construct an effective descriptor for material structure in MLIP, the following criteria are essential.

- Spatial Translation Invariance: the descriptor must remain consistent regardless of any shifts in the coordinate system, ensuring it respects the isometry of space.
- Rotational Invariance: it should be unaltered by any rotations of the coordinate system, adhering to the principle of isotropy of space.
- Permutation Invariance: the order of atomic indices should not impact the descriptor, meaning that reordering the atoms does not change the system's physical characteristics.
- Uniqueness: each descriptor should be distinctly tied to a specific atomic environment and correspond to a unique property, avoiding multiple representations for the same structure.
- Continuity: the descriptor should sensitively reflect minor variations in atomic arrangements, ensuring a proportional relationship between small structural changes and descriptor adjustments.
- Compactness: while providing comprehensive information necessary for accurate predictions, the descriptor should minimize the number of features to avoid complexity.
- Computational Efficiency: calculating the descriptor should be significantly less resource-intensive compared to direct computations of the physical properties it represents.

Here, we have listed the several commonly used and efficient descriptors, as shown in [Table 1](#). In detail, for constructing high-dimensional neural network potentials, Behler⁶² first reported atom-centered symmetry functions (ACSF), which can transform cartesian coordinates to a series of symmetry functions to accurately represent atomic environments ([Figure 3A](#)), and also can be extended to various systems such as molecules, solids, and liquids. Furthermore, Gastegger et al.¹⁵⁰ introduce weighted atom-centered symmetry functions (wACSFs) as an efficient and accurate alternative to conventional atom-centered symmetry functions (ACSFs) for MLIP, which offer better scalability with diverse chemical elements and require fewer functions for equal spatial resolution, significantly improving generalization in machine learning potentials. Utilizing the QM9 database's molecular structures and enthalpies, as shown in [Figures 3B](#) and [3C](#), they demonstrate that wACSFs achieve lower prediction errors compared to ACSFs. Moreover, it shows that simple empirical parametrization schemes are sufficient for high accuracy, and the use of genetic algorithms for optimizing wACSFs can further enhance small neural network potentials.

Nevertheless, Csanyi introduces the Smooth Overlap of Atomic Positions (SOAP) descriptor,⁷⁶ which maintains crucial properties such as differentiability with respect to atomic movement and invariance to physical symmetries such as rotation, reflection, translation, and permutation of atoms of the same species, and offers a more faithful representation that remains consistent regardless of the number of neighbors, avoiding the tradeoff between descriptor size and accuracy. This makes SOAP, integrated into GAP software, particularly effective for systems with larger clusters of atoms, where traditional descriptors tend to lose accuracy. Moreover, Thompson et al.¹⁵¹ introduce the Spectral Neighbor Analysis Potential (SNAP), whose uniqueness lies in its characterization of each atom's local environment through bispectrum components of local neighbor density, projected onto a hyperspherical harmonic basis in four dimensions. Unlike the GAP potential, SNAP

**Figure 3.** Depict of various descriptors

- (A) Atom-centered symmetry functions.⁶² (Reproduced with permission from Ref.,⁶² © J. Chem. Phys. 2011).
- (B) Weighted atom-centered symmetry functions.¹⁵⁰ (Reproduced with permission from Ref.,¹⁵⁰ © J. Chem. Phys. 2018).
- (C) Weighted atom-centered symmetry functions optimized with the GA using fitness functions based on linear ridge regression and neural network potentials.¹⁵⁰ (Reproduced with permission from Ref.,¹⁵⁰ © J. Chem. Phys. 2018).
- (D) Coulomb, Eward sum and sine matrices.¹⁵⁵ (Reproduced with permission from Ref.,¹⁵⁵ © Comput. Phys. Commun. 2020).
- (E) Many-body tensor representation.¹⁵⁵ (Reproduced with permission from Ref.,¹⁵⁵ © Comput. Phys. Commun. 2020).
- (F) Atomic cluster expansion.¹⁵⁶ (Reproduced with permission from Ref.,¹⁵⁶ © Phys. Rev. B 2019).
- (G) Atomic-position independent material descriptor.¹⁵⁷ (Reproduced with permission from Ref.,¹⁵⁷ © Phys. Rev. B 2018).
- (H) Deep potential dynamic descriptor.¹⁵⁸ (Reproduced with permission from Ref.,¹⁵⁸ © Phys. Rev. Lett. 2018).
- (I) Graph descriptor.¹⁵⁹ (Reproduced with permission from Ref.,¹⁵⁹ © Phys. Rev. Lett. 2018).

assumes a linear relationship between atom energy and bispectrum components, with linear coefficients determined via weighted least-squares linear regression against a comprehensive quantum mechanics training set.

Subsequently, as shown in Figure 3D, Rupp et al.¹⁵² report molecular representation methods based on Coulomb matrices, inspired by the nuclear repulsion term in the molecular Hamiltonian. By transforming the challenge of solving the molecular Schrödinger equation into a more manageable nonlinear statistical regression problem, the model is trained and validated on a dataset of over seven thousand organic molecules, showing a mean absolute error of approximately 10 kcal/mol. This approach demonstrates high accuracy and transferability in predicting atomization energies for a wide range of organic molecules and distorted equilibrium geometries, highlighting the broader utility of the Coulomb matrix as a descriptor in various chemical contexts. Similarly, Faber et al.¹⁵³ also generalize the Coulomb matrix representation, successful in modeling atomization energies of organic molecules, to periodic systems using three approaches: (i) a matrix based on the Ewald sum of electrostatic interactions in the unit cell; (ii) an extended Coulomb-like matrix considering neighboring unit cells; and (iii) a sine matrix representation mimicking the periodicity and elemental features of the Ewald sum matrix. These representations were compared using a Laplacian kernel with the Manhattan norm on a dataset of 3938 crystal structures. The generalization error in predicting structure formation energies was 0.49, 0.64, and 0.37 eV/atom for the respective representations. Next, Haoyan et al.¹⁵⁴ highlight the development of a many-body tensor representation (MBTR) for MLIP, which is invariant to translations, rotations, and nuclear permutations, and can represent both molecules and crystals, and is showcased in phase diagrams of Pt-group/transition-metal binary systems, as shown in Figure 3E. The Coulomb, Ewald, sine matrices, and MTBR can be performed in DScript software packages.¹⁵⁵

Additionally, Drautz et al.¹⁵⁶ report the atomic cluster expansion (ACE) as a novel and comprehensive descriptor method for MLIP, efficiently scaling linearly with the number of neighbors, a significant improvement over traditional many-atom expansion. Applied to small Cu clusters, it demonstrates smooth convergence to accuracies within the meV range, whose innovation lies in its ability to integrate nonlinear functions into the expansion, producing interatomic potentials that rival the accuracy of advanced MLIPs, as shown in Figure 3F. In addition,

Thompson et al.¹⁵⁷ also report an atomic-position independent material descriptor with the feature of using crystallographic space group and Wyckoff-sites, rather than specific atomic positions, to describe structure details of materials, as shown in Figure 3G. Employing this atomic-position independent descriptor, they also used an attention-weights sharing convolution neural network, which learns across diverse structure types without explicitly using atomic positions, to achieve a mean absolute error of only 0.07 eV/atom in predicting the formation energies of over 85,000 diverse materials.

In addition to the aforementioned static descriptors related to numerical analysis, Zhang et al.¹⁵⁸ have proposed dynamic descriptors in *DeepPotential* software, which involves training a neural network to automatically determine the coefficients of different many-body interaction terms, as shown in Figure 3H. Although this method has improved accuracy, it requires a significant number of computational resources. Particularly, with the development of mathematical graph theory, unlike traditional methods that rely on manually constructed feature vectors or complex transformations of atom coordinates, Jeffrey et al.¹⁵⁹ introduce an original machine learning framework, Crystal Graph Convolutional Neural Networks (CGCNN), which directly learns material properties from the atomic connections within a crystal structure, as shown in Figure 3I. This approach offers a universal and interpretable representation for various types of crystalline materials and has demonstrated high accuracy in predicting properties calculated by DFT for a diverse range of crystal structures and compositions, using a dataset of approximately 10^4 data points.

MACHINE LEARNING MODELS OF MACHINE LEARNING INTERATOMIC POTENTIALS

In this section, we categorize MLIPs into six types of methods and summarize the software codes that can be executed for each type of MLIP, as listed in Table 2. The first type is artificial neural networks (ANN) and deep neural networks (DNN), originally proposed and used in MLIP by Behler et al.,¹⁶⁰ collectively referred to as NN in Table 2. Initially, this model converts spatial configurations into numerical matrices based on the structural descriptors mentioned in Section [Materials descriptors of machine learning interatomic potentials](#), and then iterates the model through forward and backward propagation algorithms, obtaining the best predictive results of the model performance through the loss function, as shown in Figure 3A. Detailed descriptions of this type of model can be found in the referenced articles.¹⁶¹ This method has now extended to include more single-layer neural network NEP methods, high-dimensional approaches 4G-HDNNP,⁹⁷ Accurate Neural Network Engine (ANI-1),¹⁶² Hierarchical Interacting Particle NN,¹⁶³ and GPU-accelerated algorithms such as aenet-Pytorch.¹⁶⁴

The second method is the graph neural network (GNN) model.^{159,165–168} The process of constructing a potential function for a graph neural network model in the article can be described as follows, as shown in Figure 4A. The executable code can be found in Table 2.

- Defining relationships between nodes: the relationships between nodes, represented as edges in the graph, are defined. In the context of chemical molecules, nodes can represent atoms, and edges can represent chemical bonds. The similarity between nodes is also determined.
- Establishing graph convolutional network: the core of the graph neural network is the graph convolutional network, which models the relationships between nodes. Parameters such as the number of layers, the number of nodes per layer, and the feature dimensions of the nodes are determined. Additionally, the similarity calculation and weight update methods between nodes are specified.
- Extracting node features: each node in the graph convolutional network has a set of feature vectors representing its attributes. These feature vectors undergo nonlinear transformations to extract and transform more meaningful representations. The parameters for the nonlinear transformation and the dimensions of the feature vectors are determined.
- Training and optimizing the model: the potential function represents the state or behavior of the nodes in the graph neural network. It calculates the output value for each node. After defining the potential function, the model is trained and optimized using training data. Gradient descent or other optimization algorithms are employed to update the model's parameters, enhancing its prediction accuracy. Hyperparameters, loss functions, and evaluation metrics for the model are also determined during this process.

Furthermore, message passing network is one type of GNN, rapidly advancing in accuracy and generality for MLIPs. This research presents a TensorMol-0.1 model,¹⁶⁹ which describes an original approach to energy calculation using a combination of short-range and long-range potentials based on message-passing neural networks. A key feature is the use of TensorFlow for automatic differentiation in molecular potential calculations, allowing for the efficient optimization of network weights. The study also introduces a charge model for molecular dipole moments and a modified damped-shifted force for Coulomb energy calculation, ensuring smooth transition and differentiability, as shown in Figure 4B. The other message passing GNN include DimeNet++,¹⁷⁰ DimeNet,¹⁷¹ MACE,¹⁷² and so forth.

Next, the third method of MLIPs is the E(n)-Equivariant Graph Neural Networks (EGNNs),¹⁷³ an original model for learning graph neural networks that are equivariant to rotations, translations, reflections, and permutations, as shown in Figure 4C. Unlike existing methods, the EGNNs do not require computationally intensive higher-order representations in intermediate layers, yet they deliver competitive or superior performance. A distinctive advantage of the EGNNs is their scalability to higher-dimensional spaces beyond the typical 3D, broadening their applicability. The article demonstrates the model's effectiveness in various applications including dynamical systems modeling, graph autoencoder representation learning, and predicting molecular properties. In the same way, there are many other EGNN models, such as NequIP,¹⁷⁴ UNET,¹⁷⁵ Tensor-Field Networks,¹⁷⁶ and so forth.

As shown in Figure 4D, the fourth category of MLIPs is polynomial-type,¹⁷⁷ which are potential function fitting methods formed by combinations of polynomials, similar to the MTP descriptors and others mentioned in the descriptor in Section [Materials descriptors of machine learning interatomic potentials](#). The fifth MLIP is Gaussian Approximation Potential,⁷⁵ which addresses the gap between models that explicitly treat electrons and those that do not, aiming to accurately model the Born-Oppenheimer potential energy surface (PES) without simulating

Table 2. The list of implemented software of various MLIP models

Model	Type	Release time	Reference
NNP	NN	1995	Blank ⁶⁹
BPNN	NN	2007	Behler ¹⁶⁰
ænet-Fortran	NN	2016	Artrith ²²²
DTNN	NN	2017	Schutt ²²³
ANI-1	NN	2017	Smith ¹⁶²
HIP-NN	NN	2018	Nebgen ¹⁶³
SchNet	NN	2018	Schutt ²²⁴
DeepPotential	NN	2018	Zhang ^{158,225,226}
PhysNet	NN	2019	Unke ²²⁷
ACE	NN	2019	Drautz ¹⁵⁶
LieConv	NN	2020	Finzi ²²⁸
4G-HDNNP	NN	2021	Ko ⁹⁷
NEP	NN	2021	Fan ²²⁹
ænet-Pytorch	NN	2023	Artrith ¹⁶⁴
GNNFF	GraphNN	2021	Park ²³⁰
MDGNN	GraphNN	2021	Wang ²³¹
NoisyNodes	GraphNN	2022	Godwin ²³²
M3GNet	GraphNN	2022	Chen ²¹¹
TeaNet	GraphNN	2022	Takamoto ²³³
PotentialMind	GraphNN	2023	Wang ¹⁸⁷
CHGNet	GraphNN	2023	Deng ²¹⁴
TensorMol 0.1	GraphNN	2018	Yao ¹⁶⁹
DimeNet++	GraphNN	2020	Gasteiger ¹⁷⁰
DimeNet	GraphNN	2020	Gasteiger ¹⁷¹
NewtonNet	GraphNN	2021	Haghhighatlari ²³⁴
SphereNet	GraphNN	2021	Liu ²³⁵
PAINN	GraphNN	2021	Schutt ²³⁶
GemNet	GraphNN	2022	Gasteiger ²³⁷
HermNet	GraphNN	2022	Wang ²³⁸
MACE	GraphNN	2022	Batatia ¹⁷²
Allegro	GraphNN	2023	Musaelian ²³⁹
Tensor-Field	EquivariantNetwork	2018	Thomas ¹⁷⁶
Cormorant	EquivariantNetwork	2019	Anderson ²⁴⁰
UNET	EquivariantNetwork	2021	Qiao ¹⁷⁵
EGNN	EquivariantNetwork	2021	Satorras ¹⁷³
NequIP	EquivariantNetwork	2022	Batzner ¹⁷⁴
MTP	Polynomials	2016	Shapeev ¹⁷⁷
IPMLs	Polynomials	2018	Bereau ²⁴¹
aPIPs	Polynomials	2021	Allen ²⁴²
GAP	Gaussian	2010	Bartok ⁷⁵
FCHL19	Gaussian	2020	Christensen ¹⁷⁸
GMNN	Gaussian	2020	Zaverkin ¹⁷⁹
SE(3)-Transformers	Transfomer	2020	Fuchs ²⁴³
DeepMoleNet	Transfomer	2021	Liu ²⁴⁴
TorchMD-NET	Transfomer	2022	Thölke ¹⁸⁰

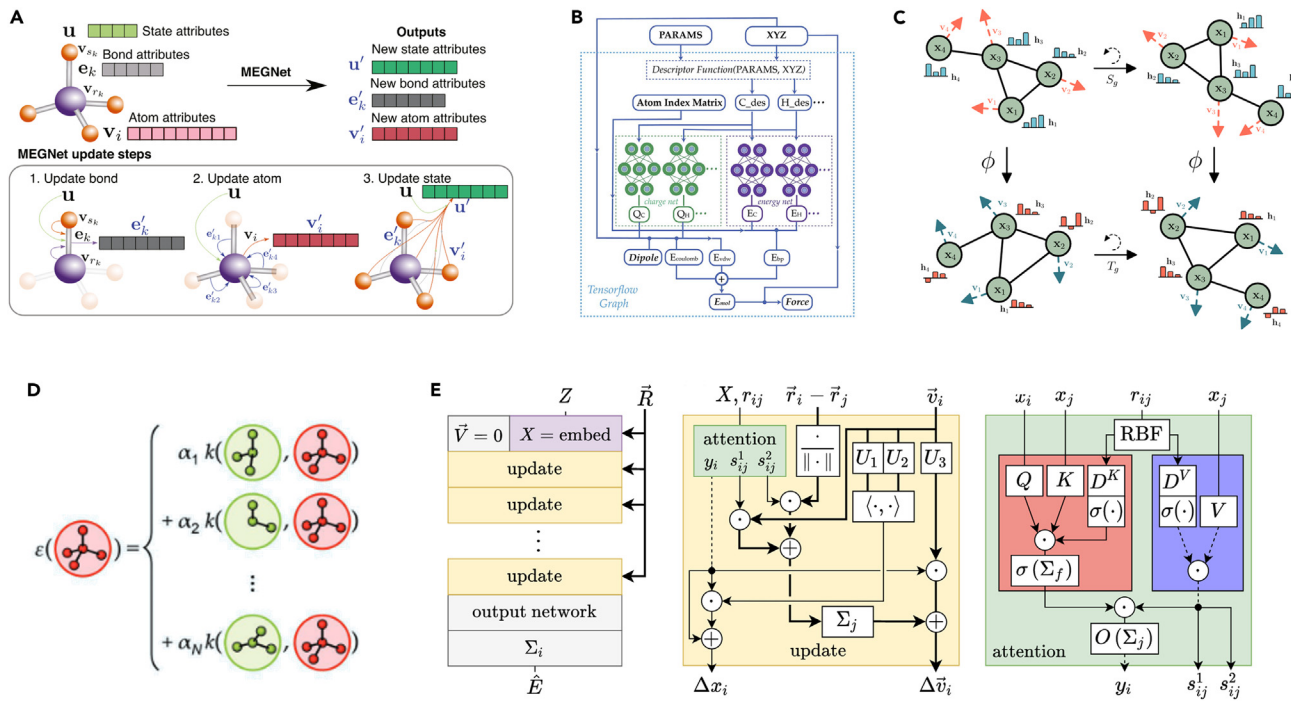


Figure 4. Schematic of the ML model in MLIPs

- (A) Many-body graph potential and the major computational blocks.¹⁶⁵ (Reproduced with permission from Ref.,¹⁶⁵ © Chem. Mater. 2019).
- (B) Charge the network and energy network for message passing network.¹⁶⁹ (Reproduced with permission from Ref.,¹⁶⁹ © Chem. Sci. 2018).
- (C) Example of rotation equivariance on a graph.¹⁷³ (Reproduced with permission from Ref.,¹⁷³ © Int. Conf. Mach. Learn. 2022).
- (D) Polynomial model.⁹⁸ (Reproduced with permission from Ref.,⁹⁸ © Adv. Mater. 2019).
- (E) Transformer model.¹⁸⁰ (Reproduced with permission from Ref.,¹⁸⁰ © The Author(s). 2022).

electrons, such as FCHL19,¹⁷⁸ GMNN,¹⁷⁹ and so forth. Finally, this study introduces TorchMD-NET,¹⁸⁰ a novel equivariant Transformer (ET) architecture for MLIP, notably overcoming the traditional trade-off between accuracy and computational speed. The model sets a standard in accuracy and efficiency, outperforming existing methods on the MD17, ANI-1, and QM9 datasets. A key feature of TorchMD-NET is its attention-based architecture that leverages rotationally equivariant features, making it particularly effective in predicting energies and atomic forces in molecular dynamics. The study also emphasizes the importance of including off-equilibrium conformations in training datasets for a more accurate evaluation of molecular potentials. By analyzing the model's attention weights, the researchers gain insights into the molecular representation, noting differences in how the model treats hydrogen atoms in energy-minimized molecules versus those in off-equilibrium states, as shown in Figure 4E.

APPLICATION OF MACHINE LEARNING INTERATOMIC POTENTIALS IN MATERIALS SCIENCE

Accelerating the molecular dynamic calculation in multi-scale simulation

Gabriele⁶¹ firstly reports the creation and validation of a Neural Network (NN) potential for the phase change materials (PCMs) GeTe. The NN potential successfully replicates the characteristics of GeTe's crystalline, liquids, and amorphous states, closely matching previous DFT calculations, as shown in Figure 5A. The study highlights the potential's robustness in larger (4096-atom) simulations and its sensitivity to fluctuations in smaller (216-atom) models. Notably, the NN potential is also applicable to slightly altered GeTe alloys but faces limitations with off-stoichiometric compositions, such as Ge_{0.15}Te_{0.85}, due to its inability to accurately model long Te-Te chains.^{181–183} Additionally, using the GAP framework, another research¹⁸⁴ introduces an MLIP for the single ternary PCMs compounds of Ge₂Sb₂Te₅, which enabled the creation of a detailed 7200-atom model, providing insights into the material's structure, and facilitated the generation of smaller models for in-depth chemical bonding studies, as shown in Figure 5B. Furthermore, without extra datasets, they expand the above model to six Sb-Te alloy PCMs,^{185,186} with compositions ranging from 2:3 to 4:1, revealing that all exhibit similar local structural motifs of defective octahedra. Figure 5C reveals that the crystallization behavior can be influenced by the noticeable shift in medium-range order and cavity concentration with the Sb content increases. For instance, Sb₂Te₃, fulling of ABAB rings, favors nucleation-driven crystallization, while Sb-rich alloys such as Sb₄Te show more 5-fold rings, leading to growth-driven crystallization due to structural dissimilarities with the crystalline phase. Meanwhile, Wang et al.¹⁸⁷ also develop a graph convolution neural network for six binary Sb-Te alloys, with a broader composition ranging from 7:1 to 1:2, as shown in Figure 5D. For the exploration of PCM behaviors in realistic memory device geometries and conditions, Zhou et al.³⁰ introduce an ML-based potential model, trained using quantum-mechanical data, to simulate all Ge-Sb-Te compositions used in PCMs, which significantly enhances

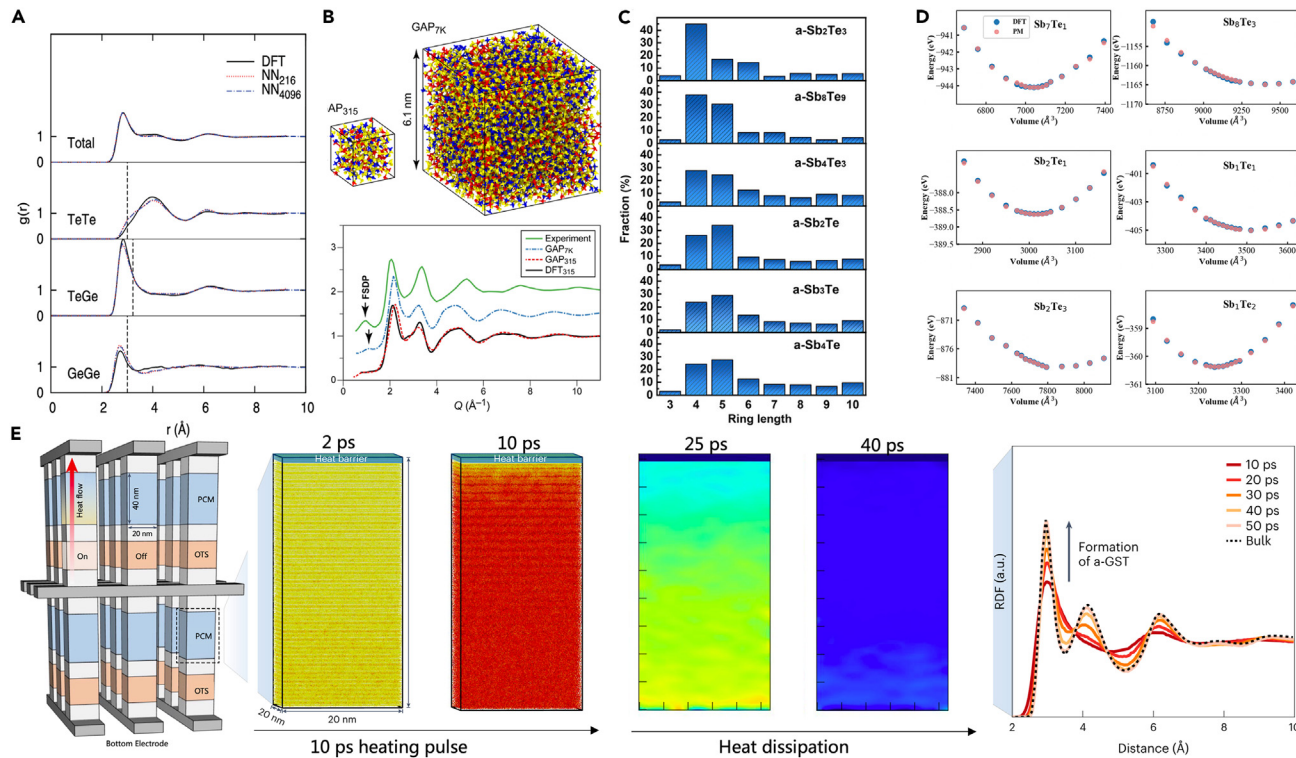


Figure 5. The molecular dynamic calculation results, which is accelerated by MLIPs in multi-scale simulation

(A) The radial distribution function compared MLIP with DFT in GeTe.⁶¹ (Reproduced with permission from Ref.,⁶¹ © Phys. Rev. B 2012).

(B) The radial distribution function compared MLIP with DFT in Ge₂Sb₂Te₅.¹⁸⁴ (Reproduced with permission from Ref.,¹⁸⁴ © J. Phys. Chem. B 2018).

(C) The fraction of primitive rings for the six amorphous Sb-Te alloys.¹⁸⁵ (Reproduced with permission from Ref.,¹⁸⁵ © Phys. Status Solidi Rrl: Rapid Res. Lett. 2021).

(D) The energy-volume curves of six crystal Sb-Te alloys.¹⁸⁷ (Reproduced with permission from Ref.,¹⁸⁷ © J. Phys. Chem. C 2023).

(E) Schematic of commercial cross-point products, the processes of melting and heat dissipation in a Ge₁Sb₂Te₄ structural model, and the evolution of the amorphous GST.²¹⁹ (Reproduced with permission from Ref.,²¹⁹ © Sci. Bull. 2023).

the speed and accuracy of atomistic simulations. Notably, it supports simulations of multiple thermal cycles and operations crucial for neuro-inspired computing, including cumulative SET and iterative RESET processes. A large-scale device model, containing over half a million atoms, demonstrates the model's capability to accurately depict critical processes in PCM-based memory devices, as shown in Figure 5E. Since the scaling of MLIPs with $O(N_{\text{electron}})$ is identical to that of a traditional force-field MD simulation, compared with the $O(N_{\text{electron}}^3)$ scaling of DFT, this cost reduction is more effective for multi-scale simulation. Based on data from experiments and *ab initio* calculations as references, Rowe et al.¹⁸⁸ developed a GAP MLIPs model to simulate the thermal expansion of graphene, detailing the spread of small molecules on graphene surfaces and addressing nuclear quantum effects through path integral molecular dynamics, which is especially useful when precise descriptions of processes are required. MLIPs have also been used in actinide molten salts in nuclear energy. For instance, Nguyen¹⁸⁹ developed an MLIP model using AIMD data with 90 atoms, which enables the acquisition of extended molecular dynamics trajectories (in nanoseconds) for larger systems (10^3 atoms) at significantly reduced computational cost, facilitating the exploration of bonding structures, thermodynamics, and dynamics across various temperatures. Deng et al.¹⁹⁰ expended MLIPs on complex ceramics with three kinds of representative structures, i.e., Ti₃SiC₂ of the MAX structure, zircon of the mineral structure, and lead zirconate titanate of the perovskite structure, facilitating the analysis and design of complex crystalline materials. Miwa et al.¹⁹¹ successfully applied MLIPs for large molecular dynamics simulations of the lithium superionic conductor Li₁₀GeP₂S₁₂. In a word, MLIPs offer an accuracy nearly equivalent to that of AIMD but at a much lower cost, which accelerates molecular dynamic calculations for large material systems (up to 100 million atoms¹⁹²), bridging the gap in multi-scale simulation.

Prediction of material properties

MLIP has been widely applied to predict various material properties. For metal alloys, a study has developed a MTPs approach¹⁹³ for materials prediction using MLIP to approximate quantum-mechanical energies, coupled with an active learning algorithm for optimal training dataset selection. Unlike methods limited to a few lattice types, this approach can predict structures with lattice types absent in the training dataset. For Cu-Pd, Co-Nb-V, and Al-Ni-Ti metallic alloys, they discovered stable structures beyond AFLOW's listings, thanks to exploring a vast number of candidate structures (40,000 for Cu-Pd, 27,000 for Co-Nb-V, and 377,000 for Al-Ni-Ti) and using MTPs for relaxation. This method drastically cuts

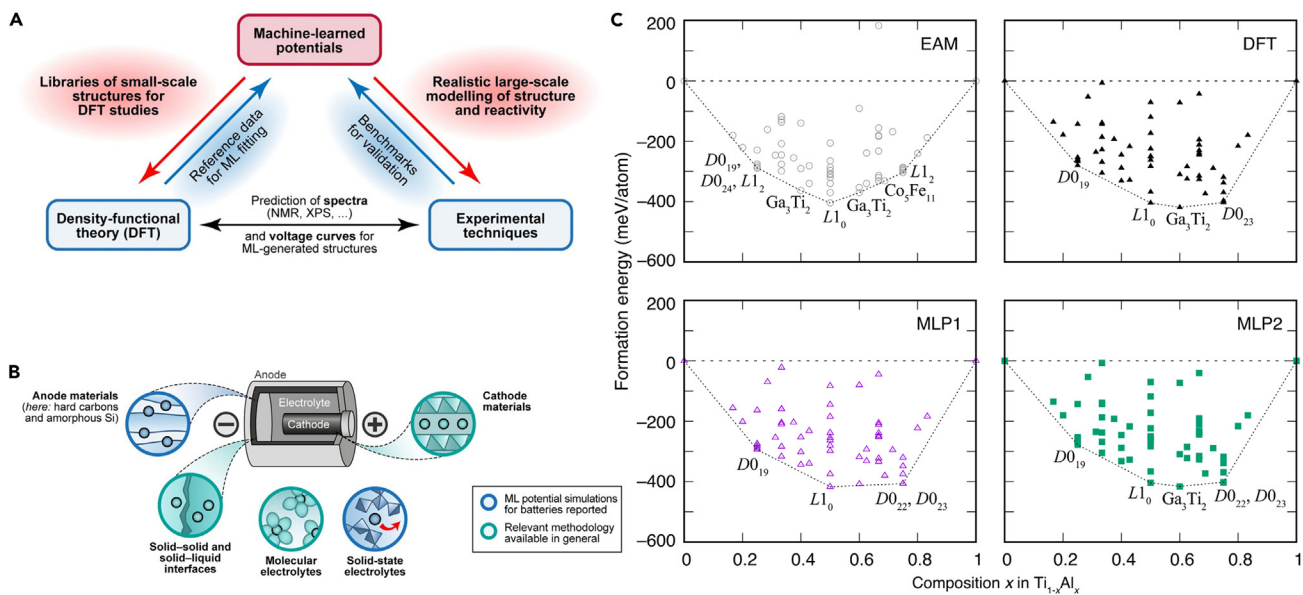


Figure 6. The flowcharts and results of the application of MLIPs in the prediction of material properties

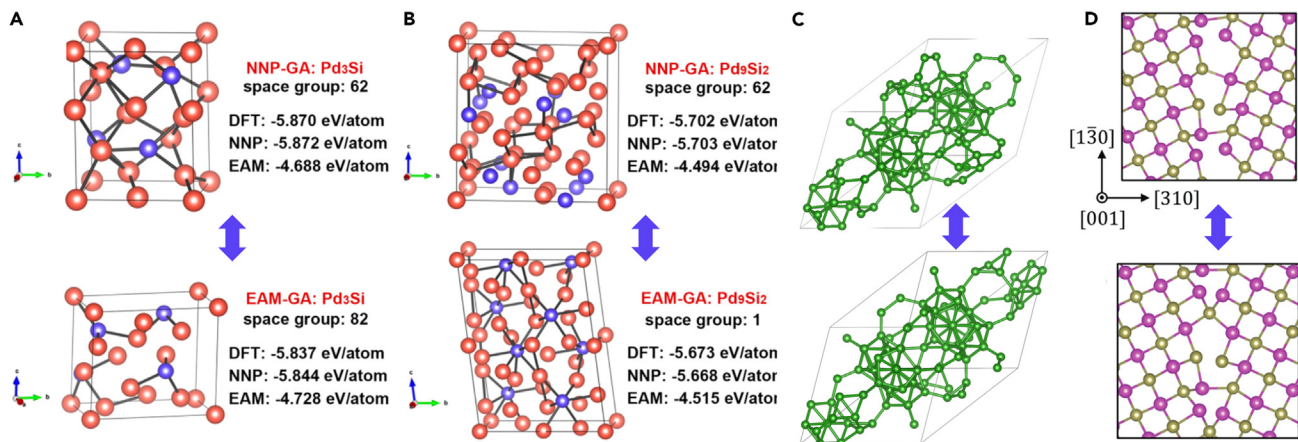
(A) A viewpoint on the synergy between machine learning-based potentials, traditional density functional theory (DFT) methods, and experimental approaches in pursuing the characterization and comprehension of battery materials at the atomic level.¹⁹⁵ (Reproduced with permission from Ref.,¹⁹⁵ © J. Phys. Energy 2020). (B) A highly simplified sketch of different components of a battery, each requiring different atomistic modeling approaches.¹⁹⁵ (Reproduced with permission from Ref.,¹⁹⁵ © J. Phys. Energy 2020).

(C) Formation energies and convex hull of $Ti_{10x}Al_x$ with different compositions.¹⁹⁸ (Reproduced with permission from Ref.,¹⁹⁸ © Phys. Rev. B 2020).

down on the number of required DFT calculations, marking a significant advancement in the discovery of alloy phases, especially for those with multiple components. For studying phononic properties of materials,¹⁹⁴ particularly focusing on low-symmetry and 2D nanomaterials. Bohayra et al. have developed an MLIP method as an efficient alternative to the traditional DFT simulations with PHONOPY code, which often face high computational demands and can produce nonphysical results for such complex structures. This study demonstrate that this approach can accurately and efficiently substitute traditional DFT methods, offering a practical solution for assessing the dynamical stability and phononic properties of complex materials. In addition, MLIP serves as a pivotal tool, complementing established DFT methods and experimental techniques, particularly in characterizing and understanding battery materials at an atomic scale,^{195,196} as shown in Figure 6A. This integration of ML-based potentials with traditional approaches enhances our capability to delve into the complex nature of battery components. The utilization of ML-based potentials is especially crucial given the diverse atomistic modeling requirements of different battery components, as illustrated in a simplified sketch of a battery's various parts in Figure 6B. This development marks a stride forward in the precision and efficiency of modeling battery materials, providing a more comprehensive understanding of their atomic-level behavior and interactions.

As shown in Figure 6C, for the prediction of mechanical properties, MLIPs are used to predict the elastic constants and mechanically induced rippling in graphene and hBN,¹⁹⁷ stacking fault energy of Ti-Al alloys,¹⁹⁸ elastic tensor of C_{11} , C_{21} , and C_{44} , and dislocation emissions of Al-Cu alloys.¹⁹⁹ For instance, Arabha et al.²⁰⁰ highlight the application of MLIPs for assessing the mechanical properties of nitrogenated holey graphene (C_2N), a two-dimensional nanomaterial. They revealed a thermal conductivity of 85.5 ± 3 W/m-K and an elastic modulus of 390 ± 3 GPa, demonstrating close agreement with DFT results. Additionally, their study explored the impact of point defects on C_2N 's mechanical properties, showing a reduction in the elastic modulus, fractural stress, and ultimate strength with the incorporation of defects. The effectiveness of MLIPs in simulating the thermal and mechanical behavior of C_2N , including defective structures, underscores their potential as a powerful tool for predicting the properties of two-dimensional nanostructures. By developing a model named UNEP-v1, Fan et al.²⁰¹ constructed unified general MLIPs for 16 elemental metals and their alloys, showing an efficient approach to represent the vast chemical space without the need to generate training data for all possible combinations. This work highlights the MLP's application in investigating the mechanical behavior of complex materials, such as the plasticity and primary radiation damage in MoTaVW refractory high-entropy alloys. The study's approach, which employs distinct neural networks for each species and multiple loss functions for parameter optimization, is scalable and adaptable, paving the way for future models that could encompass the entire periodic table.

In addition to their strong potential in simulating and assessing mechanical properties, MLIPs have also been shown to predict characteristics of thermal properties effectively. Arabha et al.²⁰⁰ reviewed the thermal conductivity of a few 2D and 3D structures, which have been calculated using MLIPs, and compared with their experimental and quantum counterparts. Ladygin et al.²⁰² introduced a groundbreaking method integrating active learning and MLIPs, specifically moment tensor potentials (MTPs), for predicting lattice dynamics properties with accuracy comparable to DFT. The research thoroughly evaluated the accuracy of these potentials by investigating four materials (Al, Mo, Ti, and U) with varying phonon and thermodynamic properties, achieving high fidelity in reproducing both harmonic and anharmonic

**Figure 7. Comparison of the structure searching results by DFT and MLIPs**

Lowest-energy structures obtained from the classical GA searches by NNP and EAM at the composition of (A) Pd₃Si and (B) Pd₉Si₂ with the corresponding DFT, NNP, and EAM energies.²⁰⁶ (Reproduced with permission from Ref.,²⁰⁶ © Phys. Rev. B 2019).

(C) Comparison of the found 106-atom structure and the existing structure.²⁰⁷ (Reproduced with permission from Ref.,²⁰⁷ © Phys. Rev. B 2019).

(D) Grain boundary structures predicted by ANN and DFT structural relaxation.²⁰⁹ (Reproduced with permission from Ref.,²⁰⁹ © Phys. Chem. Chem. Phys. 2022).

behaviors. Furthermore, the study showcases the MLIPs' capability to efficiently calculate phonon dispersion curves and vibrational density of states, providing insights into materials' thermodynamic performances. Notably, MLIPs offer substantial computational speed-ups over traditional DFT calculations without compromising accuracy, marking a significant advancement in the field of materials science and enabling detailed exploration of complex material behaviors on an unprecedented scale. In addition, MLIPs are used for predicting the phonon density of states for β -Ga₂O₃ consisting of 2500-atoms,²⁰³ unraveling the significant regulation of stress on the contribution of optical phonons to thermal conductivity in layered Li₂ZrCl₆,²⁰⁴ exploring the four-phonon scattering in WS₂,²⁰⁵ to name a few.

Structure searching

In materials science, the conventional approach of calculating material energies through DFT is widely adopted due to its high accuracy. However, a significant drawback of DFT is its time-consuming nature, particularly in the structural search of complex material systems. To overcome this limitation, MLIP methods leverage machine learning techniques to predict material properties and energies, significantly speeding up the structure search process while maintaining reasonable accuracy. By learning from existing DFT computational data, these methods rapidly evaluate a vast array of potential material structures, greatly enhancing the efficiency of material design and discovery. Recent research highlights the development of neural network MLIP for the Pd-Si system,²⁰⁶ addressing the challenge of achieving both accuracy and efficiency in structure prediction. The NNP methods demonstrate superior accuracy over the existing embedded-atom method (EAM) potential, especially in predicting ground-state structures for Pd₃Si and Pd₉Si₂ compositions, where the EAM potential fails, as shown in Figure 7A. Additionally, Evgeny et al.²⁰⁷ introduces a crystal structure prediction methodology combining the evolutionary algorithm USPEX with an MLIP that learns on-the-fly, which significantly accelerates the process, offering a computational efficiency several orders of magnitude greater than DFT methods. It automates the construction of the interatomic interaction model, eliminating the need for manual training dataset assembly and seamlessly replacing DFT in the prediction algorithm. Tested on various elements including carbon, sodium under high pressure, and boron, as shown in Figure 7B. In order to predict grain boundary, Takayuki et al.^{208,209} highlight the effectiveness of MLIP in predicting grain boundary energies in face-centered-cubic (fcc) elemental metals such as Ag, Al, Au, Cu, Pd, and Pt. Despite the absence of grain boundary structures in the training datasets, MLIPs demonstrated high accuracy in forecasting grain boundary structures and energies, aligning well with DFT. Furthermore, Tatsuya et al. present the development of an artificial neural network (ANN) MLIP for accurately predicting grain boundary (GB) atomic structures in CdTe. Using a comprehensive training dataset that includes DFT data of point defects, surfaces, and GBs, the ANN potential was trained to cover various atomic environments. The potential's effectiveness was demonstrated by its ability to predict low-energy structures and GB energetics with accuracy comparable to DFT results, outperforming conventional empirical potentials, as shown in Figure 7C. The study also outlines a method for improving the ANN potential's transferability to more complex GBs, using limited training data. Except for predicting grain boundary configurations, MLIP methods for predicting crystal defects have also been developed. As shown in Figure 7D, Alexandra et al. introduce a novel quadratic noise ML (QNML) approach for bcc Fe and W, focusing on accurately modeling radiation defects and dislocations, enhancing the accuracy of standard linear MLIP without compromising transferability. These potentials, trained on a range of defect configurations, accurately predict complex issues such as the stability and mobility of defects, which is challenging for semiempirical potentials.

Pre-trained universal machine learning interatomic potential model

Existing MLIPs are generally designed for narrow target materials, making them unsuitable for broader applications in material discovery. Therefore, Takamoto et al.²¹⁰ first report PreFered Potential (PFP), a universal Neural Network Potential (NNP) capable of handling any

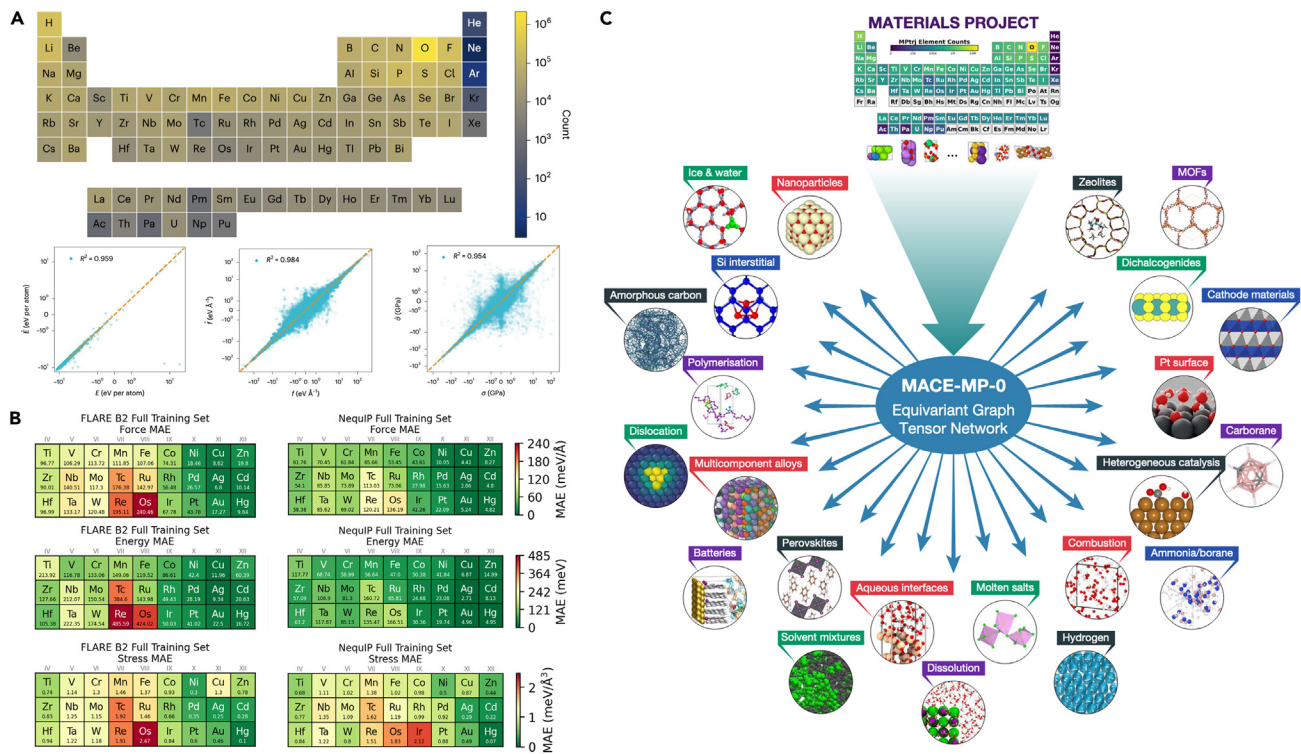


Figure 8. The three pre-trained universal MLIPs model

(A) Element counts for all atoms in the dataset, covering 89 elements across the periodic table. And the model predictions on the test dataset compared to DFT calculations.²¹¹ (Reproduced with permission from Ref.,²¹¹ © Nat. Comput. Sci. 2022).

(B) Force, energy, and stress mean absolute errors for the 27 TM systems, using the B₂ invariant ACE descriptors and NequIP.²¹² (Reproduced with permission from Ref.,²¹² © The Author(s). 2023).

(C) A foundation model for materials modeling, which is based on Materials Project data. MACE-MP-0 is capable of molecular dynamics simulation across a wide variety of chemistries in the solid, liquid, and gaseous phases.²¹³ (Reproduced with permission from Ref.,²¹³ © The Author(s). 2023).

combination of 45 elements. Unlike existing NNPs tailored for specific materials, PFP's versatility is demonstrated in diverse applications, including lithium diffusion in LiFeSO₄F, molecular adsorption in metal-organic frameworks, Cu-Au alloy transitions, and Fischer-Tropsch catalyst discovery. PFP shows high quantitative accuracy and computational efficiency, even reproducing structures and properties not initially considered in its design.

In addition, Shyue et al.²¹¹ introduce a universal MLIP model (M3GNet) based on graph neural networks with three-body interactions, trained on the extensive Materials Project database. M3GNet can be applied to a wide range of materials for structural relaxation, dynamic simulations, and property prediction. Figure 8A reveals the distribution of the initial datasets and the accuracy on the test dataset compared to DFT calculations. Using M3GNet, around 1.8 million potentially stable materials were identified from 31 million hypothetical structures, with 1,578 out of the top 2,000 low-energy materials confirmed as stable via DFT. M3GNet's broad applicability extends to molecular dynamics simulations, identifying potential lithium superionic conductors and serving as a surrogate for DFT in other structural exploration techniques. Its current version, the best achievable with existing data, anticipates future enhancements through more accurate training data and active learning strategies.

For many-body interactions, recent research focuses on the accuracy of MLIPs for bulk solid and liquid phases of d-block elements (TM23 dataset).²¹² It compares two FLARE and NequIP models in transition metals, revealing that early transition metals pose a greater challenge in learning accuracy than later group metals, as shown in Figure 8B. The complexity of interatomic interactions varies among these metals, influenced by their electronic structures and the many-body character of interactions. Additionally, based on 1.5 million structures dataset, Ceder et al. also trained a universal graph MLIP, named Crystal Hamiltonian Graph Neural Network (CHGNet), to learn and accurately represent the orbital occupancy of electrons, enhancing its capability to describe both atomic and electronic degrees of freedom.

Another work introduces MACE-MP-0,²¹³ as shown in Figure 8C, a general-purpose machine-learned force field model, trained on a public database of 150k inorganic crystals. Capable of stable molecular dynamics across various materials, MACE-MP-0 marks a significant advancement in atomistic modeling, demonstrating both qualitative and quantitative accuracy in diverse physical science problems, including water and aqueous systems, catalysis, metal-organic frameworks and the performance on calculating phonon dispersions, bulk and shear moduli of crystals, atomic energies and lattice constants of elemental solids, the cohesive energies of molecular crystals, the reaction barrier heights, and the homonuclear diatomic binding curves. Despite its broad applicability, the model has limitations, including its reliance on the PBE

exchange-correlation functional, lack of explicit long-range interactions, and challenges in capturing intermolecular interactions and high-pressure conditions. Future improvements may involve refitting with modern functionals, incorporating electrostatic and spin interactions, and extending training data. MACE-MP-0, while a foundational model, still requires fine-tuning for specific systems, but shows promise as a step toward democratizing MLIP by lowering barriers to entry in material simulations.²¹⁴

PERSPECTIVE OF MACHINE LEARNING POTENTIAL

Standard datasets for machine learning interatomic potentials

The vast range of time and spatial scales in material computational simulations, from quantum-level calculations to macroscopic continuum models, results in a diverse array of simulation software and data formats tailored to specific applications. Therefore, standard datasets play a crucial role in the development and evaluation of MLIPs. These datasets serve as the foundational tools upon which the robustness, accuracy, and versatility of MLIPs are built and assessed. Standard datasets provide a common ground for benchmarking various MLIP models. By training and testing different MLIPs on the same dataset, researchers can objectively compare their performance, ensuring a fair and consistent evaluation of different approaches. Particularly, the diversity and comprehensiveness of high-quality standard datasets are essential for training and validating MLIPs, which determine the generalization capability of the models. In order to learn a broader spectrum of atomic interactions, the initial datasets must cover a wide range of chemical compositions, crystal structures, and physical conditions.

To overcome the lack of standardized, universal material datasets for MLIPs, firstly, the development of unified data formats and interoperability standards across different simulation software will facilitate data sharing and integration, such as Pymatgen,²¹⁵ ASE,²¹⁶ and so forth. Secondly, establishing centralized databases for material properties, accessible to researchers worldwide, could promote data standardization, consisting of material structure databases, such as Materials Project,²¹⁷ AFLOW,^{119,218} Materials Cloud,¹²⁰ and ALKEMIE-Matter DB.¹²² Finally, leveraging machine learning and artificial intelligence to harmonize and translate between different data formats could bridge the gap between diverse datasets, enhancing the field's overall efficiency. Recently, Justin et al.⁴⁷ reported the ANI-1ccx and ANI-1x datasets, which especially can be used to build generalized and accurate ML models and assess the accuracy of different models.

Trade-off between accuracy and complexity in machine learning interatomic potentials

The development of MLIPs often faces a critical trade-off between accuracy and complexity. As the complexity of an MLIP increases to achieve higher accuracy, the computational efficiency typically decreases. This can negate one of the main advantages of MLIPs over traditional ab initio methods, which is to enable faster and larger-scale simulations. In addition, a more complex MLIP might fit the training data exceptionally well, but lead to poor performance on unseen data. Finally, complex models may not scale well with system size, limiting their applicability to larger systems or longer time scales, which are often the primary motivation for using MLIPs. Therefore, it is preferable to construct models with minimal complexity while ensuring accuracy for specific material problems.

To balance accuracy and complexity in ML models, feature engineering and dimensionality reduction methods are essential to improve the quality of datasets while retrieving material information from the data ocean. In the training process, regularization techniques such as Lasso or Ridge are utilized to prevent overfitting without overly simplifying the model. Additionally, cross-validation and ensemble methods are used to combine multiple models to improve accuracy without significantly increasing individual model complexity. Lastly, iterative refinement, by gradually adjusting model complexity based on performance metrics, can ensure a balanced approach, optimizing both accuracy and computational efficiency.

Transferability and generalization

Training MLIPs involves datasets of immense complexity and size, whose cost associated with gathering data, computational processing, and model training is significant. Therefore, transferability and generalization are cornerstone features that determine the practical usefulness of MLIPs in diverse computational simulations. Transferable and generalizable MLIPs can accurately predict the potential energy surface for a wide range of chemical elements and compounds, not just those included in the training dataset. This is crucial for the exploration of materials and chemical phenomena. If MLIPs can generalize well, it reduces the need to create and train other models for every unique material system, saving computational resources and time.

To extend MLIP models that are applicable across various materials, incorporating a diverse dataset representing a broad range of materials and their properties can enhance the model's generalizability. Second, using transfer learning techniques allows models trained on one material to adapt to new materials with minimal additional training. Finally, developing models with scalable architectures (such as the Transformer model in large language model) ensures they can handle the complexity of different materials. In addition, a unified API for MLIPs supports the integration of these models into existing computational frameworks, enabling seamless workflow and interoperability between different simulation tools. Lastly, continuous validation and updating of the model with strange data ensure its applicability to emerging materials. This further democratizes access to advanced simulation capabilities, allowing researchers to readily apply MLIPs across various applications without needing to navigate disparate software ecosystems.

CONCLUSIONS

In this review, the advancements and applications of MLIPs in materials science are explored, highlighting their role in overcoming the limitations of high computational costs and specificity inherent in density-functional theory and classical molecular dynamics. The four essential

stages of MLIP development are thoroughly examined, including data generation techniques, material structure descriptors, machine learning algorithms, and executable software, providing a roadmap for researchers in the field.

A significant portion of the review is dedicated to the practical applications of MLIPs in various domains such as phase-change memory materials, structural searching, material properties prediction, and the development of pre-trained universal models. These applications showcase the versatility and robustness of MLIPs in addressing complex material science challenges. Notably, the review delves into the future perspectives of MLIPs, discussing the need for standard datasets, the balance between accuracy and complexity, and the importance of transferability and generalization in these models.

In summary, this review paints a comprehensive picture of the current state and future potential of MLIPs in materials science. It underscores the transformative impact of MLIPs in bridging the gap between small-scale models and realistic device-scale simulations, thereby revolutionizing the approach to materials research and design. The advancements in MLIPs promise not only enhanced efficiency and accuracy but also open new horizons for material discovery and innovation.

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AUTHOR CONTRIBUTIONS

Conceptualization, G. W., J. Z., and Z. S.; investigation, G. W., C. W., X. Z., and Z. L.; resources, G. W., C. W., X. Z., and Z. L.; writing – original draft, G. W., J. Z., and Z. S.; writing – review and editing, G. W., J. Z., and Z. S.; funding acquisition, G. W., J. Z., and Z. S.; supervision J. Z. and Z. S.

DECLARATION OF INTERESTS

The authors declare no financial interest.

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