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SE(3)-Equivariant Graph Neural Networks for Data-Efficient and Accurate Interatomic Potentials

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This work presents Neural Equivariant Interatomic Potentials (NequIP), a SE(3)-equivariant neural network approach for learning interatomic potentials from *ab-initio* calculations for molecular dynamics simulations. While most contemporary symmetry-aware models use invariant convolutions and only act on scalars, NequIP employs SE(3)-equivariant convolutions for interactions of geometric tensors, resulting in a more information-rich and faithful representation of atomic environments. The method achieves state-of-the-art accuracy on a challenging set of diverse molecules and materials while exhibiting remarkable data efficiency. NequIP outperforms existing models with up to three orders of magnitude fewer training data, challenging the widely held belief that deep neural networks require massive training sets. The high data efficiency of the method allows for the construction of accurate potentials using high-order quantum chemical level of theory as reference and enables high-fidelity molecular dynamics simulations over long time scales.

INTRODUCTION

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dynamics (MD)simulations 22 indispensable tool for computational discovery in fields 37 as diverse as energy storage, catalysis, and biological 38 processes [1–3]. While the atomic forces required to ³⁹ integrate Newton's equations of motion can in principle 40 be obtained with high fidelity from quantum-mechanical 41 calculations such as density functional theory (DFT), 42 in practice the unfavorable computational scaling of 43 first-principles methods limits simulations to short time 44 This prohibits 45 scales and small numbers of atoms. 31 the study of many interesting physical phenomena 46 beyond the time and length scales that are currently 47 accessible, even on the largest supercomputers. Owing to their simple functional form, classical models for the atomic potential energy can typically be evaluated orders of magnitude faster than using first-principles methods, thereby enabling the study of large numbers of atoms over long time scales. However, due to their limited mathematical form, classical interatomic potentials, or force fields, are inherently limited in their predictive accuracy which has historically led to a fundamental trade-off between obtaining high computational efficiency while also predicting faithful dynamics of the system under study. The construction of flexible models of the interatomic potential energy based on Machine Learning (ML-IP), and in particular

Neural Networks (NN-IP), has shown great promise in 82 providing a way to move past this dilemma, promising 83 to learn high-fidelity potentials from ab-initio reference 84 calculations while retaining favorable computational 85 51 efficiency [4–13]. One of the limiting factors of NN-IPs 86 is that they typically require collection of large training 87 53 sets of ab-initio calculations, often including thousands 88 or even millions of reference structures [4, 9, 10, 14–16]. 89 This computationally expensive process of training data 90 collection has severely limited the adoption of NN-IPs 91 as it quickly becomes a bottleneck in the development of 92 force-fields for new systems. Kernel-based approaches, 93 such as e.g. Gaussian Processes (GP) [5, 8] or Kernel 94 Ridge Regression (KRR) [17], are a way to remedy this 95 problem as they often generalize better from limited 96 62 sample sizes. However, such methods generally tend 97 63 to exhibit poor computational scaling with the number 98 of reference configurations, in both training (cubic in 99 65 training set size) and prediction (linear in training set₁₀₀ This limits both the amount of training data101 67 they can be trained on as well as the length and size of 102 simulations that can be simulated with them. 103 69

In this work, we present the Neural Equivariant 104
Interatomic Potential (NequIP), a highly data-efficient
deep learning approach for learning interatomic 105
potentials from reference first-principles calculations. 106
We show that the proposed method obtains high 107
accuracy compared to existing ML-IP methods across 108
a wide variety of systems, including small molecules, 109
water in different phases, an amorphous solid, a reaction 110
at a solid/gas interface, and a Lithium superionic 111
conductor. Furthermore, we find that NequIP exhibits 112
exceptional data efficiency, enabling the construction of 113

accurate interatomic potentials from limited data sets of fewer than 1,000 or even as little as 100 reference ab-initio calculations, where other methods require orders of magnitude more. It is worth noting that on small molecular data sets, NeguIP outperforms not only other neural networks, but is also competitive with kernel-based approaches, which typically obtain better predictive accuracy than NN-IPs on small data sets (although at significant additional cost scaling in training and prediction). We further demonstrate high data efficiency and accuracy with state-of-the-art results on a training set of molecular data obtained at the quantum chemical coupled-cluster level of theory. validate the method through a series of simulations and demonstrate that we can reproduce with high fidelity structural and kinetic properties computed from NeguIP simulations in comparison to ab-initio molecular dynamics simulations (AIMD). We directly verify that the performance gains are connected with the unique SE(3)-equivariant convolution architecture of the new NeguIP model.

Related Work

First applications of machine learning for the development of interatomic potentials were built on descriptor-based approaches combined with shallow neural networks or Gaussian Processes [4, 5], designed to exhibit invariance with respect to translation, permutation of atoms of the same chemical species, and rotation. Recently, rotationally invariant graph neural networks (GNN-IPs) have emerged as a powerful architecture for deep learning of interatomic potentials

that eliminates the need for hand-crafted descriptors₁₄₈ and allows to instead learn representations on graphs₁₄₉ 115 of atoms from invariant features of geometric data (e.g.150 116 radial distances or angles) [9–11, 13]. In GNN-IPs,151 117 atomic structures are represented by collections of 152 nodes and edges, where nodes in the graph correspond₁₅₃ 119 to individual atoms and edges are typically defined 154 by simply connecting every atom to all other atoms₁₅₅ 121 that are closer than some cutoff distance r_c . Every 156 122 node/atom i is associated with a feature $\mathbf{h}_i \in \mathbb{R}^h$,157 consisting of scalar values, which is iteratively refined₁₅₈ 124 via a series of convolutions over neighboring atoms j_{159} based on both the distance to neighboring atoms r_{ij} 100 126 and their features \mathbf{h}_{i} . This iterative process allows₁₆₁ information to be propagated along the atomic graph₁₆₂ 128 through a series of convolutional layers and can be₁₆₃ 129 viewed as a message-passing scheme [18]. Operating 164 130 only on interatomic distances allows GNN-IPs to be₁₆₅ 131 rotation- and translation-invariant, making both the166 132 output as well as features internal to the network₁₆₇ 133 invariant to rotations. In contrast, the method outlined 168 in this work uses relative position vectors rather than 169 135 simply distances (scalars), which makes internal features instead equivariant to rotation and allows for angular 137 information to be used by rotationally equivariant filters. 138 Similar to other methods, we can restrict convolutions 139 to only a local subset of all other atoms that lie closer 140 to the central atom than a chosen cutoff distance r_c , see₁₇₀ 141 Figure 1, left. 142

A series of related methods have recently been 173 proposed: DimeNet [11] expands on using pairwise 174 interactions in a single convolution to include angular, 175 three-body terms, but individual features are still 176 comprised of scalars (distances and three-body angles are invariant to rotation), as opposed to vectors used in this work. Another central difference to NequIP is that DimeNet explicitly enumerates angles between pairs of atoms and operates on a basis embedding of distances and angles, whereas NequIP operates on relative position vectors and a basis embedding of distances, and thus never explicitly computes three-body angles. Cormorant [19] uses an equivariant neural network for property prediction on small molecules. This method is demonstrated on potential energies of small molecules but not on atomic forces or systems with periodic boundary conditions. Townshend et al. [20] use the framework of Tensor-Field Networks [21] to directly predict atomic force vectors. The predicted forces are not guaranteed by construction to conserve energy since they are not obtained as gradients of the total potential energy. This may lead to problems in simulations of molecular dynamics over long times. None of these three works [11, 19, 20] demonstrates capability to perform molecular dynamics simulations.

In this work we present a deep learning energy-conserving interatomic potential for both molecules and materials built on SE(3)-equivariant convolutions over geometric tensors that yields state-of-the-art accuracy, outstanding data-efficiency, and can with high fidelity reproduce structural and kinetic properties from molecular dynamics simulations.

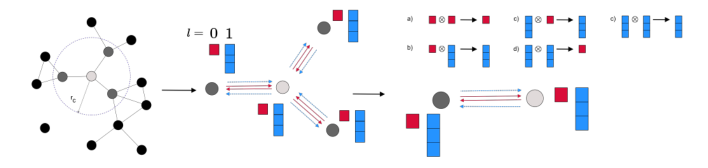


FIG. 1: Left: a set of atoms is interpreted as an atomic graph with local neighborhoods. Middle: every atom carries a set of scalar and vector features with it. Right: atoms exchange information via filters, that are again scalars and vectors. The interactions of features and filters define five interactions.

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RESULTS

Equivariance

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The concept of equivariance arises naturally in²⁰⁰ machine learning of atomistic systems (see e.g. [22]): physical properties have well-defined transformation $_{201}$ properties under translation and rotation of a set of atoms. As a simple example, if a molecule is rotated²⁰² in space, the vectors of its atomic dipoles or forces²⁰³ also rotate accordingly, via equivariant transformation.²⁰⁴ Equivariant neural networks are able to more generally²⁰⁵ represent tensor properties and tensor operations of physical systems (e.g. vector addition, dot products, and cross products). Equivariant neural networks are guaranteed to preserve the known transformation properties of physical systems under a change of coordinates because they are explicitly constructed from equivariant operations. Formally, a function $f: X \to Y$ is equivariant with respect to a group G that acts on Xand Y if:

 $D_Y[g|f(x) = f(D_X[g|x) \quad \forall g \in G, \forall x \in X$ (1)212

where $D_X[g]$ and $D_Y[g]$ are the representations of the group element g in the vector spaces X and Y, respectively. In this work, we focus on equivariance with respect to SE(3), i.e. the group of rotations and translations in 3D space.

Neural Equivariant Interatomic Potentials

Given a set of atoms (a molecule or a material), we aim to find a mapping from atomic positions \vec{r}_i and chemical species (identified by atomic numbers Z_i) to the total potential energy and the forces acting on the atoms:

$$f: \{\vec{r}_i, Z_i\} \to E_{pot} \tag{2}$$

Forces are obtained as gradients of the predicted potential energy with respect to the atomic positions, which guarantees energy conservation:

$$\vec{F}_i = -\nabla_i E_{pot} \tag{3}$$

Gradients can be obtained with relatively low computational overhead in modern auto-differentiation frameworks such as TensorFlow or PyTorch [23, 24]. Following previous work [4], we further define the total

potential energy of the system as a sum of atomic²⁴⁴ potential energies: ²⁴⁵

$$E_{pot} = \sum_{i \in N_{atoms}} E_{i,atomic} \tag{4}_{247}$$

These atomic local energies $E_{i,atomic}$ are the scalar node attributes predicted by the graph neural network. Even though the output of NequIP is the predicted potential energy E_{pot} , which is invariant under translations and rotations, the network contains internal features that are tensors which are equivariant to rotation.

This constitutes the core difference between NequIP and existing scalar-valued invariant GNN-IPs. The remainder of this section will discuss the design of the network in further detail.

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A series of methods has been introduced to realize 226 rotationally equivariant neural networks [13, 21, 25, 26]. Here, we build on the layers introduced in Tensor-Field 228 Networks (TFN) [21], which enable the construction of neural networks that exhibit equivariance to translation,260 230 permutation, and rotation. Every atom in NequIP₂₆₁ is associated with a feature comprised of tensors of 262 232 scalars, vectors, and higher-order²⁶³ different order: 233 tensors. Formally, these features are geometric objects²⁶⁴ 234 that comprise a direct sum of irreducible representations²⁶⁵ 235 of the SO(3) symmetry group. Second, the convolutions²⁶⁶ that operate on these geometric objects are equivariant267 237 functions instead of invariant ones, i.e. if a feature at 268 layer k is rotated, then the output of the convolution²⁶⁹ 239 from layer $k \to k+1$ rotates accordingly. In practice, 270 240 the features are implemented as a dictionary $V_{acm}^{(l)}$ with keys l, where l = 0, 1, 2, ... is a non-negative integer and is called the "rotation order", labeling the irreducible

representations. The indices a, c, m, correspond to the atoms, the channels (elements of the feature), and the representation index which takes values $m \in [-l, l]$, respectively.

Convolution operations are naturally translation invariant, since their filters act on relative interatomic distance vectors. Moreover, they are permutation invariant since all convolution contributions are summed. Note that while atomic features are equivariant to permutation of atom indices, globally, the total potential energy of the system is invariant to permutation. To achieve rotation equivariance, the convolution filters are constrained to be products of learnable radial functions and spherical harmonics, which are equivariant under SO(3) [21]:

$$F(\vec{r}_{ij}) = R(r_{ij})Y_m^{(l)}(\hat{r}_{ij})$$
 (5)

where if \vec{r}_{ij} denotes the relative position from central atom i to neighboring atom j, \hat{r}_{ij} and r_{ij} are the associated unit vector and interatomic distance, respectively. It should be noted that all learnable weights in the filter lie in the rotationally invariant radial function $R(r_{ij})$. This radial function is implemented as a small neural network with one hidden layer and a shifted softplus activation function [9], operating on interatomic distances expressed in a basis of choice, $R(r_{ij}): \mathbb{R}^{N_b} \to \mathbb{R}^h$, where N_b is the number of basis functions and h is the feature dimension:

$$R(r_{ij}) = W_2 \ln(0.5 \exp(W_1 B(r_{ij})) + 0.5) \tag{6}$$

where $W_1 \in \mathbb{R}^{N_{hidden} \times N_b}$ and $W_2 \in \mathbb{R}^{h \times N_{hidden}}$ are 298
weight matrices, h is the dimension of the feature and 299 N_{hidden} is the dimension of the hidden layer in the 300
feed-forward neural network (in our experiments, we use 301 $N_{hidden} = N_b$, resulting in comparatively small neural 302
networks for the radial function). Radial Bessel functions 303
and a polynomial envelope function f_{env} discussed in 304
recent work [11] are used to expand the interatomic 305
distances:

$$B(r_{ij}) = \sqrt{\frac{2}{r_c}} \frac{\sin(\frac{n\pi}{r_c} r_{ij})}{r_{ij}} f_{env}(r_{ij}, r_c)$$
 (7)

where r_c is a local cutoff radius, restricting interactions₃₁₀ to atoms closer than some cutoff distance and f_{env} is the polynomial defined in [11] with p = 6 operating on the interatomic distances normalized by the cutoff312 283 radius $\frac{r_{ij}}{r_c}$. The use of cutoffs/local atomic environments 284 allows the computational cost of evaluation to scale linearly with the number of atoms. Similar to [11], we initialize the Bessel functions with $n = [1, 2, ..., N_b]$ and subsequently optimize $n\pi$ via backpropagation rather 288 than keeping it constant. For systems with periodic boundary conditions, we use the neighbor_list 290 functionality as implemented in the ASE code [27] to identify appropriate atomic neighbors and then convolve 292 over them.

Finally, in the convolution, the input feature tensor₃₂₄ and the filter have to again be combined in an₃₂₅ equivariant manner, which is achieved via a geometric₃₂₆

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tensor product, yielding an output feature that again is rotationally equivariant. A tensor product of two geometric tensors is computed via Clebsch-Gordan coefficients, as outlined in [21]. Since NequIP deals with force vectors, the network design is simplified by only using scalar (l=0) and vector (l=1) representations. Thus, we can enumerate five distinct products or "interactions" between l=0 and l=1 filters and l=0 and l=1 features that correspond to simple operations between scalars and vectors:

- $0 \otimes 0 \to 0$ (product of two scalars)
- $0 \otimes 1 \to 1$ (scalar multiplication of a vector)
- $1 \otimes 0 \rightarrow 1$ (scalar multiplication of a vector)
- $1 \otimes 1 \rightarrow 0$ (dot product of two vectors)
- $1 \otimes 1 \rightarrow 1$ (cross product of two vectors)

It is trivial to include higher-order interactions, and previous works have increased the rotation order beyond l=1 [20, 28]. However, it should be noted that every interaction comes with additional trainable radial functions and hence additional weights, which thus adds to the model capacity, increasing the number of model weights and the memory footprint of the model. Omitting all higher-order interactions that go beyond the $0 \otimes 0 \to 0$ interaction will result in a conventional GNN-IP with invariant convolutions over scalar features, similar to e.g. SchNet [9]. Finally, as outlined in [21], a full convolutional layer $\mathcal L$ implementing an interaction with filter f acting on an input i producing output o: $l_f \otimes l_i \to l_o$ is given by:

$$\mathcal{L}_{acm_o}^{(l_o)}(\vec{r}_a, V_{acm_i}^{(l_i)}) = \sum_{m_f, m_i} C_{(l_f, m_f)(l_i, m_i)}^{(l_o, m_o)} \sum_{b \in S} R_c^{(l_f, l_i)}(r_{ab}) Y_{m_f}^{(l_f)}(\hat{r}_{ab}) V_{bcm_i}^{(l_i)}$$

$$\tag{8}$$

where a and b index the central atom of the convolution₃₅₇ 327 and the neighboring atom $b \in S$, respectively, and 358 C indicates the Clebsch-Gordan coefficients. As an 359 329 example of this, we write out a full $1 \otimes 1 \rightarrow 1$ operation₃₆₀ (corresponding to a cross-product) in the Methods₃₆₁ 331 section. After every convolution, output tensors of a₃₆₂ 332 rotation order l stemming from different tensor products³⁶³ 333 are concatenated on a per-atom basis. 334 atomic features, the model also leverages self-interaction₃₆₅ 335 layers similar to SchNet [9], corresponding to dense₃₆₆ 336 layers that are applied in an atom-wise fashion with₃₆₇ 337 weights shared across atoms. While different weights₃₆₈ 338 are used for different rotation orders, the same set of 369 weights is applied for all representation indices m of a_{370} 340 given rotation order l. Shifted softplus nonlinearities₃₇₁ 341 [9] are used as rotation-equivariant nonlinearities as₃₇₂ 342 introduced in [21], which are applied to the Euclidean₃₇₃ norm of the input feature, the output of which is in turn₃₇₄ 344 combined with the input tensor, thus preserving overall₃₇₅ 345 equivariance.

The NequIP network architecture, shown in Figure 2, is built on an atomic embedding, followed by a series of interaction blocks, and finally an output block: 37

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- Embedding: following SchNet, the initial feature is generated using a trainable embedding, that operates on the atomic number Z_i (represented via a one-hot encoding) alone, implemented via a trainable self-interaction layer.
- Interaction Block: interaction blocks encode386

interactions between neighboring atoms: the core of this block is the convolution function, outlined in equation 8. For every output rotation order l_o , the features from different tensor product interactions are concatenated to give a new feature, which is in return refined with atom-wise self-interaction layers and equivariant non-linearities. We equip interactions blocks with a ResNet-style update [29] where the input feature \mathbf{x} is updated atom-wise via the output of an interaction block $f(\mathbf{x})$ that gives the final feature $r(\mathbf{x}) = f(\mathbf{x}) + \mathbf{x}$ (features are added element-wise in the m-dimension). Note that this operation is equivariant since the addition of an equivariant feature \mathbf{x} and an equivariant function $f(\mathbf{x})$ preserves equivariance. While later interaction blocks include all five interactions outlined above, the first interaction block operates on the l=0 embedding with a $0\otimes 0\to 0$ and a $0 \otimes 1 \to 1$ only.

 Output Block: the l = 0 feature of the final convolution is passed to an output block, which consists of another atom-wise self-interaction layer, an equivariant non-linearity, and a final atom-wise self-interaction layer.

The scalar atomic outputs of the final layer can be interpreted as atomic potential energies which are summed to give the total predicted potential energy of the system (Equation 4). Forces are subsequently obtained as the negative gradient of the predicted total potential energy, thereby ensuring both energy

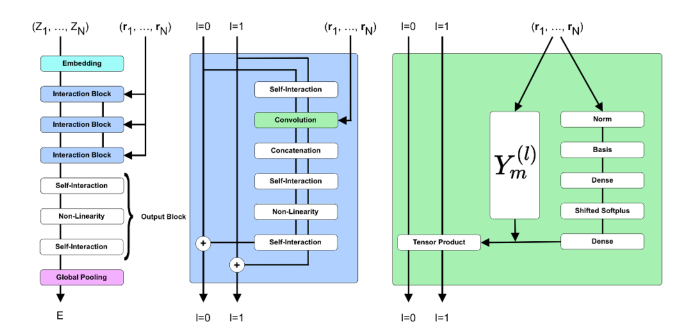


FIG. 2: The NequIP network architecture. Left: atomic numbers are embedded into l=0 features, which are refined through a series of interaction blocks, creating l=0 and l=1 features. An output block generates atomic energies, which are pooled to give the total predicted energy. Middle: the interaction block consists of a series of convolutions, interweaved with self-interaction layers, equivariant nonlinearities and concatenation. Right: the convolution combines the radial function R(r) which operates only on interatomic distances with the spherical harmonics based on unit vector \hat{r} via a tensor product.

conservation as well as rotation-equivariant forces (see₄₀₁ 387 equation 3).

Experiments

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We validate the proposed method on a series of diverse 390 and challenging data sets: first we demonstrate that we 391 improve upon state-of-the-art accuracy on MD-17, a data 392 set of small, organic molecules that is widely used for 393 benchmarking ML-IPs [9, 11, 17, 30, 31]. Next, we show 408 394 that NeguIP can also accurately learn forces obtained on409 395 small molecules at the quantum chemical CCSD(T) level₄₁₀ 396 [31], opening the door to scalable and efficient molecular₄₁₁ 397 dynamics simulations with beyond-DFT accuracy. To₄₁₂ 398 broaden the applicability of the method beyond small₄₁₃ 399 isolated molecules, we explore a series of extended414 number of configurations for validation, and evaluate 400

systems with periodic boundary conditions, consisting of both surfaces and bulk materials: water in different phases [15, 32], a chemical reaction at a solid/gas interface, an amorphous Lithium Phosphate [12], and a Li superionic conductor [13]. Details of the training procedure are provided in the Methods section.

MD-17 small molecule dynamics

We first evaluate NeguIP on MD-17 [17, 30, 31], a data set of eight small organic molecules in which reference values of energy and forces are generated by ab-initio MD simulations with DFT. For training we use N=1,000 structure configurations for each molecule, sampled uniformly from the full data set, the same

Force training at quantum chemical accuracy

the test error on all remaining configurations in the445 The mean absolute error in the force 416 components is shown in Table I in units of [meV/Å]. 446 We compare results using NequIP with those from⁴⁴⁷ 418 published leading ML-IP models that were also trained 448 on 1,000 structures: in particular SchNet [9], DimeNet⁴⁴⁹ 420 [11] (both graph neural networks), sGDML [31], and 450 421 FCHL19/GPR (kernel-based methods) [33]. We find⁴⁵¹ 422 that NequIP outperforms SchNet and sGDML on all 452 423 molecules in the data set, DimeNet on 7 out of 8⁴⁵³ molecules (on par on the remaining one), and performs on 454 425 par with FCHL/19GPR. The consistent improvement in 455 accuracy upon sGDML and the comparable performance $^{456}\,$ 427 to FCHL19/GPR are particularly surprising, as these are 457 based on kernel methods, that typically tend to be more⁴⁵⁸ 429 sample efficient. It should be noted, however, that the 459 430 evaluation cost of kernel methods scales linearly with the 460 431 number of training configurations. Note also that on⁴⁶¹ 432 some molecules, NeguIP trained on 1,000 configurations⁴⁶² 433 even performs as well as SchNet trained on $50,000^{463}$ 434 structures [9]: on aspirin and naphthalene, for example, 464 the NequIP network trained on 1,000 structures produces⁴⁶⁵ 436 mean absolute errors in the forces of 15.1 meV/Å and 4.2^{466} meV/Å, respectively, compared to 14.3 meV/Åand 4.8⁴⁶⁷ 438 meV/Å of SchNet trained on 50x more molecules, hinting⁴⁶⁸ 439 that NequIP exhibits exceptional data efficiency. On⁴⁶⁹ 440 other molecules such as ethanol, however, SchNet trained⁴⁷⁰ with 50,000 molecules still clearly outperforms NeguIP⁴⁷¹ 442 trained with 1,000 molecules (2.2 meV/Å for SchNet for 472 443 N=50,000 vs 9.0 meV/Å for NequIP for N=1,000).

Ability to achieve high accuracy on a comparatively small data set opens the door to training models on expensive high-order ab-initio quantum chemical It has been shown that DFT can fail to methods. capture important subtleties in the potential energy surface, potentially even identifying the wrong ground states [31]. This problem can be remedied through the use of more accurate reference calculations, such as coupled cluster methods CCSD(T), typically regarded as the gold standard of quantum chemistry. However, the high computational cost of CCSD(T) has thus far hindered the use of reference data structures at this level of theory, prohibited by the need for large data sets that are required by available NN-IPs. Leveraging the high data efficiency of NequIP, we evaluate it on a set of molecules computed at quantum chemical accuracy (aspirin at CCSD, all others at CCSD(T)) [31] and compare the results to those reported for sGDML [31]. The training/validation set consists of a total of 1,000 molecular structures which we split into 950 for training and 50 for validation (sampled uniformly), and we test the accuracy on all remaining structures (we use the train/test split provided with the data set, but further split the training set into training and validation sets). We find that NequIP achieves lower errors on four out of five molecules, performing on par with sGDML on the fifth molecule, as shown in Table II.

Liquid Water and Ice Dynamics

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To demonstrate the applicability of NequIP beyond small molecules, we evaluate the method on a series of

Molecule	NequIP	SchNet	sGDML	DimeNet	FCHL19/GPR
Aspirin	15.1	58.5	29.5	21.6	20.7
Benzene [17]	8.1	13.4	n/a	8.1	n/a
Benzene [31]	2.3	n/a	2.6	n/a	n/a
Ethanol	9.0	16.9	14.3	10.0	5.9
Malonaldehyde	14.6	28.6	17.8	16.6	10.6
Naphthalene	4.2	25.2	4.8	9.3	6.5
Salicylic Acid	10.3	36.9	12.1	16.2	9.6
Toluene	4.4	24.7	6.1	9.4	8.8
Uracil	7.5	24.3	10.4	13.1	4.6

TABLE I: MAE of force components on the MD-17 data set, trained on 1,000 configurations, forces in units of [meV/Å]. For the benzene molecule, two different data set exists from [17], [31] with different levels of accuracy in the DFT reference data.

Molecule	NequIP	sGDML	49
Aspirin	14.7	33.0	_
Benzene	0.8	1.7	4
Ethanol	9.4	15.2	4
Malonaldehyde	16.0	16.0	
Toluene	4.4	9.1	4

TABLE II: Force MAE for molecules at CCSD/CCSD(T) accuracy, reported in units of [meV/Å], with 1,000 reference configurations).

extended systems with periodic boundary conditions. As a first example we use a joint data set consisting of liquid 477 water and three ice structures [15, 32], computed at the PBE0-TS level of theory. This data set contains [15]: 479 a) liquid water, P=1bar, T=300K, computed via pathintegral AIMD, b) ice Ih, P=1bar, T=273K, computed 481 via path-integral AIMD c) ice Ih, P=1bar, T=330K, res 482 computed via classical AIMD d) ice Ih, P=2.13 kbar, 483 T=238K, computed via classical AIMD. The liquid water 506 484 system consists of 64 H₂O molecules (192 atoms), while₅₀₇ 485 the ice structures consist of 96 H₂O molecules (288₅₀₈ 486 atoms). A DeepMD NN-IP model was previously trained 509 487 [15] for water and ice using a joint training set containing 510 488 133,500 reference calculations of these four systems. To511 489 assess data efficiency of the NeguIP architecture, we512 490 similarly train a model jointly on all four parts of the data513 491 set, but using only 133 structures for training, i.e. 1000x514 492

fewer data. The 133 structures were sampled uniformly from the full data set available online, consisting of water and ice structures, made up of a total of 140,000 frames, coming from the same MD trajectories that were used in the earlier work [15]. We also use a validation set of 50 frames and report the test accuracy on all remaining structures in the data set. Table III shows the comparison of the predictive force accuracy of NequIP trained on the 133 structures vs DeepMD trained on 133,500 structures. We find that with 1000x fewer training data, NequIP significantly outperforms DeepMD on all four parts of the data set.

Heterogeneous catalysis of formate dehydrogenation

Next, we demonstrate application of NequIP to a catalytic surface reaction. In particular, we investigate the dynamics of formate undergoing dehydrogenation decomposition (HCOO* \rightarrow H* + CO₂) on a Cu < 110 > surface (see Figure 3). This system is highly heterogeneous, with both metallic and covalent types of bonding as well as charge transfer occurring between the metal and the molecule, making this a particularly challenging test system. Different states of the molecule

System	NequIP, 133 data points	DeepMD, 133,500 data points
Liquid Water	35.9	40.4
Ice Ih (b)	25.9	43.3
Ice Ih (c)	16.6	26.8
Ice Ih (d)	13.5	25.4

TABLE III: Root mean square error (RMSE) of force components on liquid water and the three ices in units of [meV/A]. Note that the NequIP model was trained on < 0.1% of the training data of DeepMD.

Element	MAE
C	55.8
O	86.7
H	42.0
Cu	54.5
Total structure	55.6

TABLE IV: MAE of force components for Formate on Cu system, per-element basis. The training set consists of 2,500 structures, force units are [meV/A]

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also lead to dissimilar C-O bond lengths [34, 35]. Training structures consist of 48 Cu atoms and 4 atoms of the molecule (HCOO* or CO₂+H*). The MAE of the predicted forces using a NequIP model trained on 2,500 structures is shown in Table IV, demonstrating that NequIP is able to accurately model the interatomic forces for this complex reactive system. A more detailed analysis of the resulting dynamics will be subject of a separate study.

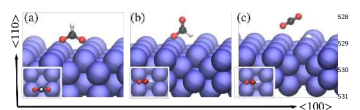


FIG. 3: Perspective view of atomic configurations of (a) bidentate HCOO (b) monodentate HCOO and (c) CO $_{2\,533}$ and a hydrogen adatom on a Cu(110) surface. The blue, red, black, and white spheres represent Cu, O, C, 534 and H atoms, respectively. The subset shown in each subplot is the corresponding top view along the <110> orientation.

Lithium Phosphate Amorphous Glass Formation

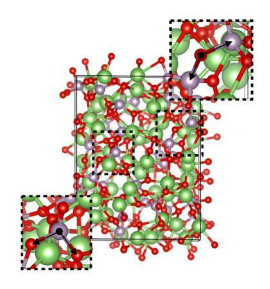


FIG. 4: Quenched glass structure of Li₄P₂O₇. The insets show the P-O-O tetrahedral bond angle (bottom left) as well as the O-P-P bridging angle between corner-sharing phosphate tetrahedra (top right).

To examine the ability of the model to capture dynamical properties, we demonstrate that NequIP can describe structural dynamics in amorphous lithium phosphate with composition $\text{Li}_4\text{P}_2\text{O}_7$. This material is a member of the promising family of solid electrolytes for Li-metal batteries [12, 36, 37], with non-trivial Li-ion transport and phase transformation behaviors. The training data set consists of two 50ps-long AIMD simulations, one of the molten structure at T=3000 K, followed by another of a quenched glass structure at T=600 K. We train NequIP on a subset of 1,000 structures of the molten trajectory, each consisting of

208 atoms, and sampled uniformly from the full data₅₇₀ set of 25,000 AIMD frames. We use a validation set of 571 539 100 structures, and evaluate the model on all remaining₅₇₂ structures. Table V shows the test set error in the force₅₇₃ 541 components on both the test set from the AIMD molten₅₇₄ trajectory and the full AIMD quenched glass trajectory.575 543 To then evaluate the physical fidelity of the trained₅₇₆ model, we use it to run a MD simulation of length 50577 545 ps at T=600 K in the NVT ensemble and compare the578 546 total radial distribution function (RDF) without element₅₇₉ distinction as well as the angular distribution functions₅₈₀ 548 (ADF) of the P-O-O (P central atom) and O-P-P (O₅₈₁ central atom) angles to the ab-inito trajectory at the 582 550 same temperature. The P-O-O angle corresponds to 583 the tetrahedral bond angle, while the O-P-P corresponds 584 552 to a bridging angle between corner-sharing phosphates85 553 tetrahedra (Figure 4). Figure 5 shows that NequIP₅₈₆ 554 can accurately reproduce the RDF and the two ADFs,587 555 in comparison with AIMD, after training on only 1,000588 structures. This demonstrates that the model generates 589 557 the glass state and recovers its dynamics and structure590 almost perfectly, having seen only the high-temperature 591 559 molten training data.

Lithium Thiophosphate Superionic Transport

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To show that NequIP can model kinetic transport properties from small training sets at high accuracy, we study Li-ion diffusivity in LiPS (Li_{6.75}P₃S₁₁) a crystalline superionic Li conductor, consisting of a simulation cell of 83 atoms [13]. MD is widely used to study diffusion; however, training a ML-IP to the accuracy required to accurately predict kinetic properties has in the past required large training set sizes ([38] e.g. uses a

data set of 30,874 structures to study Li diffusion in Li_3PO_4). Here we demonstrate that not only does NequIP obtain small errors in the force components, but it also accurately predicts the diffusivity after training on a data set obtained from an AIMD simulation. Again, we find that very small training sets lead to highly accurate models, as shown in Table V for training set sizes of 10, 100, 1,000 and 2,500 structures. We run a series of MD simulations with the NequIP potential trained on 2,500 structures in the NVT ensemble at the same temperature as the AIMD simulation for a total simulation time of 50 ps and a time step of 0.25 fs, which we found advantageous for reliability and stability of long simulations. We measure the Li diffusivity in ten Nequip-driven MD simulations (computed via the slope of the mean square displacement), all of length 50 ps and started from different initial velocities, randomly sampled from a Maxwell-Boltzmann distribution. We find a mean diffusivity of $1.42 \times 10^{-5} \text{ cm}^2/\text{s}$, in excellent agreement with the diffusivity of $1.38 \times 10^{-5} \,\mathrm{cm}^2/\mathrm{s}$ computed from AIMD, thus achieving a relative error of as little as 3%. Figure 6 shows the mean square displacements of Li for an example run.

System	Data Set Size	MAE
LiPS	10	157.1
LiPS	100	50.0
LiPS	1,000	25.1
LiPS	2,500	24.1
$\overline{\text{Li}_4\text{P}_2\text{O}_7,\text{melt}}$	1,000	63.2
$Li_4P_2O_7$, quench	1,000	36.9

TABLE V: Force MAE for LiPS and $\text{Li}_4\text{P}_2\text{O}_7$ for different data set sizes in units of [meV/A]. The model for $\text{Li}_4\text{P}_2\text{O}_7$ was trained exclusively on structures from the melted trajectory, the reported test errors show the MAE on both the test set of the melted trajectory as well as the full quench trajectory.

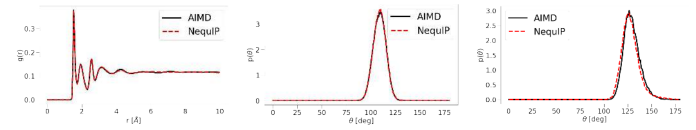


FIG. 5: Left: Radial Distribution Function, middle: Angular Distribution Function, bridging oxygen, right: Angular Distribution Function, tetrahedral bond angle. All are defined as probability density functions.

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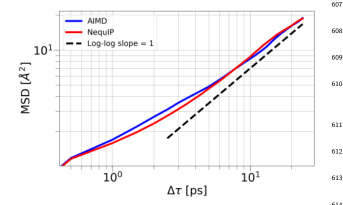


FIG. 6: Comparison of Lithium mean square displacement of AIMD and NequIP trajectories.

Data Efficiency

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exhibits₆₂₀ the above experiments, NeguIP exceptionally high data efficiency, i.e. it can be₆₂₁ trained successfully to state-of-the-art accuracy from 622 unexpectedly small training sets. It is interesting to 623 consider the reasons for such high performance and verify₆₂₄ that it is connected to the equivariant nature of the625 model. First, it is important to note that each training 626 configuration contains multiple labels, thus increasing 627 the total number of labels available beyond just the 628 potential energy label associated with each structure.629 In particular, for a training set of M first-principles₆₃₀ calculations with structures consisting of N atoms, the 631 total number of labels available is M(3N+1) since every₆₃₂

force component on every atom constitutes a label and so does the total energy of the reference calculation (we only train to atomic forces and not energies, thus using 3MN force components as labels).

In order to gain insight into the reasons behind increased accuracy and data efficiency, we perform a series of experiments with the goal of isolating the effect of using equivariant convolutions of geometric tensors compared to invariant convolutions over scalars. particular, we run a set of experiments for a system with a fixed number of training configurations in which we explicitly turn on or off interactions of higher order than l=0. This defines two settings: first, we train the network with both l = 0 and l = 1 features and all five interactions as previously outlined in this work. Second, when all interactions involving l=1 are turned off, this turns the network into a conventional invariant GNN-IP, involving only invariant convolutions over scalar features in a SchNet-style fashion. As a test system we chose bulk water: in particular we use the data set introduced in [39], consisting of 1,593 bulk liquid water structures with 64 water molecules each. We train a series of networks with identical hyperparameters, but vary the training set sizes between 10 and 1,000 structures, sampled uniformly from the full data set, as well as a validation set consisting of 100 structures. We then evaluate the error on all

remaining structures for a given training set size. As655 shown in Figure 7, we find that the equivariant setting 656 634 (using l = 0 and l = 1) significantly outperforms the 657 invariant setting (using only l=0) for all data set sizes₆₅₈ 636 as measured by the MAE of force components. This 659 637 suggests that it is indeed the use of tensor (in our specific660 638 case vector) features and equivariant convolutions that 661 639 enables the high data efficiency of NequIP. We further 662 640 note, that in [39], a Behler-Parrinello Neural Network 663 641 (BPNN) was trained on 1303 structures, yielding a664 642 RMSE of $\approx 120 \text{ meV/Å}$ in forces when evaluated ones 643 the remaining 290 structures. We find that NequIP models trained with as little as 50 and 100 data points 645 obtain RMSEs of 122.9 meV/Å and 93.3 meV/Å on their respective test sets (note that Figure 7 shows the MAE).₆₆₇ 647 This provides further evidence that NequIP exhibits₆₆₈ 648 significantly improved data efficiency in comparison with $_{669}$ 649 existing methods. 650

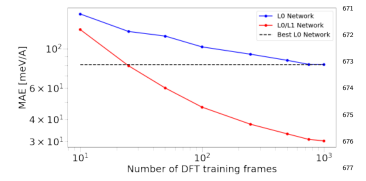


FIG. 7: Log-log plot of the predictive error in forces of ⁶⁷⁸ NequIP with l=0 vs. l=0/l=1 interactions as a function of data set size, measured via the force MAE.

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Computational Efficiency

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Finally, we report the computational efficiency of 684

NequIP and compare it to that of the *ab-inito* methods 685

on two examples shown in this work: for a molecular 686

system, we choose the Toluene molecule, computed at the CCSD(T)-level of theory [31]; for a material with periodic boundary conditions, we choose the Formate on Cu system, in which reference data were obtained with DFT. For both systems, we report the time required for a single force call on a CPU node with 32 cores. The results are shown in Table VI. In both cases, NequIP gives a large speed-up over the *ab-initio* methods. In the case of the Toluene system, this means that 58.4 minutes of a NequIP simulation can obtain the simulation time equaling one century of a CCSD(T) simulation.

DISCUSSION

We demonstrate $_{
m that}$ $_{
m the}$ Neural Equivariant Interatomic Potential (NeguIP), a new type of graph neural network built on SE(3)-equivariant convolutions exhibits state-of-the-art accuracy and exceptional data efficiency on data sets of small molecules and Furthermore, we find that we periodic materials. can reproduce structural and kinetic properties from molecular dynamics simulations with very high fidelity in comparison to ab-initio simulations. The ability to both learn from small numbers of reference samples, while retaining high computational efficiency opens the door to performing simulations of large systems over long time-scales at quantum mechanical accuracy, using DFT or higher order methods such as coupled-cluster or quantum Monte Carlo data as reference. We expect the new method will enable researchers in computational chemistry, physics, biology, and materials science to conduct molecular dynamics simulations of complex reactions and phase transformations at increased accuracy and efficiency.

System	Number of atoms	NequIP	Ab-initio	Speed-up
Toluene	15	16 ms	4 hours*	900,000
Formate on Cu	52	58 ms	$1045.6~\mathrm{s}$	18,028

TABLE VI: Time required for a single force call for NequIP in comparison to CCSD(T) for Toluene and DFT for Formate on Cu; * personal communication with Stefan Chmiela and Alexandre Tkatchenko.

METHODS

Reference Data Sets

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718 MD-17 [17, 30, 31] is a data set of₇₁₉ *MD-17*:

eight small organic molecules, obtained from MD_{720} at T=500K and computed at the_{721} simulations PBE+vdW-TS level of electronic structure theory,722 resulting in data set sizes between 133,770 and 993,237 structures. The data set was obtained from $_{\scriptscriptstyle{724}}$ http://quantum-machine.org/gdml/#datasets.

Molecules@CCSD/CCSD(T): The data set of_{727} small molecules at CCSD and CCSD(T) accuracy₇₂₈ 699 [31] contains positions, energies, and forces for five₇₂₉ 700 different small molecules: Asprin (CCSD), Benzene,730 Malondaldehyde, Toluene, Ethanol (all CCSD(T)).731 702 Each data set consists of 1,500 structures with the₇₃₂ 703 exception of Ethanol, for which 2,000 structure are₇₃₃ For more detailed information, we direct₇₃₄ the reader to [31]. The data set was obtained from₇₃₅ 706 http://quantum-machine.org/gdml/#datasets.

708 Liquid Water and Ice: The data set of liquid waters⁷³⁸ 709 and ice structures [15, 32] was generated from classical⁷³⁹ 710 AIMD and path-integral AIMD simulations at different⁷⁴⁰ 711 temperatures and pressures, computed with a PBE0-TS₇₄₁ 712 functional [15]. The data set, obtained from http:742 713 //www.deepmd.org/database/deeppot-se-data/,

contains a total of 140,000 structures, of which 100,000 are liquid water and 20,000 are Ice Ih b),10,000 are Ice Ih c), and another 10,000 are Ice Ih d).

Formate decomposition on Cu: The decomposition process of formate on Cu involves configurations corresponding to the cleavage of the C-H bond, initial and intermediate states (monodentate, bidentate formate on Cu < 110 >) and final states (H ad-atom with a desorbed CO₂ in the gas phase). Nudged elastic band (NEB) method was first used to generate an initial reaction path of the C-H bond breaking. ab initio molecular dynamics, starting from different NEB images, were run to collect a total of 6855 DFT structures. The CP2K [40] code was employed for the AIMD simulations. Each trajectory was generated with a time step of 0.5 fs and 500 total steps. We train NeguIP on 2,500 reference structures sampled uniformly from the full data set of 6,855 structures, use a validation set of 250 structures and evaluate the mean absolute error on all remaining structures. Due to the unbalanced nature of the data set (more atoms of Cu than in the molecule), we use a per-element weighed loss function in which atoms C, O₁, O₂, and H and the sum of all Cu atoms all receive equal weights.

 $Li_{1}P_{2}O_{7}$ glass: The $Li_{4}P_{2}O_{7}$ ab-initio data were generated using an ab-initio melt-quench MD simulation, starting with a stoichiometric crystal of 208 atoms (space

group P21/c) in a periodic box of $10.4 \times 14.0 \times 16.0_{778}$ Å. The dynamics used the Vienna Ab-Initio Simulation⁷⁷⁹ 745 Package (VASP) [41-43], with a generalized gradient₇₈₀ PBE functional [44], projector augmented wave (PAW)₇₈₁ 747 pseudopotentials [45], a NVT ensemble and a Nosé-782 Hoover thermostat, a time step of 2 fs, a plane-wave₇₈₃ 749 cutoff of 400 eV, and a Γ-point reciprocal-space mesh.₇₈₄ The crystal was melted at 3000 K for 50 ps, then, $_{785}$ 751 immediately quenched to 600 K and run for another, 86 752 50 ps. The resulting structure was confirmed to be_{787} amorphous by plotting the radial distribution function₇₈₈ 754 of P-P distances. The training was performed only₇₈₉ on the molten portion, and the MD simulations for a₇₉₀ 756 quenched simulation. 791

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LiPS: Lithium phosphorus sulfide (LiPS) based₇₉₃ 759 materials are known to exhibit high lithium ion₇₉₄ 760 conductivity, making them attractive as solid-state $_{795}$ electrolytes for lithium-ion batteries. Other examples of₇₉₆ 762 known materials in this family of superionic conductors₇₉₇ are LiGePS and LiCuPS-based compounds. The training 708 764 data set is taken from a previous study on graph neural₇₉₉ network force field [13], where the LiPS training data₈₀₀ were generated using ab-initio MD of an LiPS structure₈₀₁ 767 with Li-vacancy (Li_{6.75}P₃S₁₁) consisting of 27 Li, 12 P,802 768 and 44 S atoms respectively. The structure was first₈₀₃ 769 equilibrated and then run at 520 K using the NVT_{sod} ensemble for 50 ps with a 2.0 fs time step. The full $\mathrm{data_{805}}$ 771 set contains 25,001 MD frames. We set aside 10,000 frames as a fixed test set. From the remaining frames, 773 we choose training set sizes of 10, 100, 1,000, and 2,500 774 frames with a fixed validation set size of 100. In order to generate a diverse training set, we sample both the 806 training and validation sets in a way such that 30% of 807 both of them are comprised of the structures with the shortest interatomic distances out of all frames not in the test set and the remaining 70% of the training and validation set are uniformly sampled.

Liquid Water, Cheng et al.: The training set used in the data efficiency experiments on water consists of 1,593 reference calculations of bulk liquid water at the revPBE0-D3 level of accuracy, with each structure containing 192 atoms, as given in [39]. Further information can be found in [39]. The data set was obtained from https://github.com/BingqingCheng/ab-initio-thermodynamics-of-water.

Molecular Dynamics Simulations. To run MD simulations, NequIP force outputs were integrated with the Atomic Simulation Environment (ASE) [27] in which we implement a custom version of the Nosé-Hoover thermostat. We use this in-house implementation for the both the Li₄P₂O₇ as well as the LiPS MD simulations. The thermostat parameter was chosen to match the temperature fluctuations observed in the AIMD run. The RDF and ADFs for Li₄P₂O₇ were computed with a maximum distance of 10 Å (RDF) and 2.5 Å (both ADFs).

Training. Networks are trained using a loss function based on atomic forces:

$$\mathcal{L} = \frac{1}{3N} \sum_{i=1}^{N} \sum_{\alpha=1}^{3} \left| \left| -\frac{\partial \hat{E}}{\partial r_{i,\alpha}} - F_{i,\alpha} \right| \right|^2 \tag{9}$$

where N is the number of atoms in the system and \hat{E} is the predicted potential energy. Note that we do

not train on energies since atomic forces are the only842 quantities required to integrate Newton's equations of 843 809 motion. Since the predicted forces are computed as the844 810 gradient of a scalar potential, they are still conservative.845 811 If energies are of interest, however, one can add them to₈₄₆ 812 the loss function and determine the relative weighting847 813 via a trade-off parameter as done in previous works848 814 [9, 11]. In a similar fashion, it is trivial to add others49 815 quantities of interest to the loss function (e.g predicting850 816 atomic charges or multipole tensors can be of interest851 817 for modeling long-range interactions), where they may 852 818 be scalar fields, vector fields, or higher-order tensor fields.853

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Hyperparameters. Training of models was^{855} 821 performed on NVIDIA Tesla V100 GPUs. Throughout⁸⁵⁶ 822 all experiments shown in this work, we use a feature⁸⁵⁷ 823 dimension of $h=64,\,6$ interaction blocks, $N_b=8$ Bessel*58 824 basis functions and radial neural networks with one859 hidden layer, also of hidden dimension $N_{hidden} = 8,800$ 826 giving light-weight radial functions with a comparatively⁸⁶¹ small number of parameters. The final interaction block⁸⁶² 828 is followed by the output block, which first reduces⁸⁶³ the feature dimension to 16 through a self-interaction⁸⁶⁴ 830 An equivariant non-linearity is applied and 865 831 finally through another self-interaction layer the feature866 832 dimension is reduced to a single scalar output value867 833 associated with each atom that is then summed over⁸⁶⁸ to give the total potential energy. Weights were⁸⁶⁹ 835 initialized with the uniform Xavier initialization in the 870 radial networks and orthogonal initialization in the871 837 self-interaction layers, biases were initialized with a872 838 constant value of 0. In all experiments, we use the Adam₈₇₃ 839 optimizer [46] with the TensorFlow 1.14 default settings₈₇₄ of $\beta_1 = 0.9$, $\beta_2 = 0.999$, and $\epsilon = 10^{-8}$. We decrease₈₇₅ the initial learning rate of 0.001 by a decay factor of 0.8 whenever the validation RMSE in the forces has not seen an improvement for a given number of epochs: for the small molecule tasks, we set this learning rate patience to 1,000, for all other tasks we use 100. We continuously save the model with the best validation RMSE and use the model with the overall best RMSE for evaluation on the test set and MD simulations. We stop the training if either a maximum number of 50,000 epochs (one epochs equals a full pass over the training set) has been reached, or the validation force RMSE has not improved for 2,500 epochs, or the maximum training time has been exceeded, whichever occurs first. All systems were trained for a maximum of 8 days (consisting of four runs of 48-hour time-limited compute jobs, which are restarted from the best saved model, i.e. potentially including repeats in the training) with the exception of the Li₄P₂O₇, which was trained for 12 days (six 48-hour compute jobs) and the LiPS systems, which were trained for 4 days (two 48-hour compute jobs). We use a batch size of 5 structures for all small molecule tasks, and a batch size of 1 structure for all other tasks. We found small batch sizes to be important for obtaining high predictive accuracy. We also found it important to choose the radial cutoff distance r_c appropriately. A list of the cutoff radii in units of [Å] that were used for the different systems is given in Table VII.

Example of a tensor product interaction. To illustrate that the interactions outlined in this work reduce to a set of five simple operations, we write out the example of a full $1 \otimes 1 \to 1$ interaction, i.e. a convolution that uses a l=1 filter to operate on a l=1 feature, yielding again a l=1 output. This corresponds to

 $l_i = l_f = l_o = 1$, facilitating a cross-product interaction₈₇₈ Gordan coefficients reduce to the Levi-Civita symbol [21]:

between two l=1 tensors. In this case, the Clebsch-

$$C_{(l_f=1,m_f),(l_i=1,m_i)}^{(l_0=1,m_o)} \propto \epsilon_{ofi} = \begin{cases} 1 & (o,f,i) \in \{(1,2,3),(2,3,1),(3,1,2)\} \\ -1 & (o,f,i) \in \{(1,3,2),(2,1,3),(3,2,1)\} \\ 0 & \text{else} \end{cases}$$
(10)

879 relationship $Y^{(1)}(\hat{r}) \propto \hat{r}$, we recognize the output as₈₈₂ positions and the input feature element $V_{bc}^{(l=1)}$:

Evaluating equation 8 as well as ϵ_{ofi} and using the the vector cross product, taken here between the relative

$$\mathcal{L}_{ac}^{(l_o=1)}(\vec{r}_a, V_{ac}^{(l_i=1)}) = \begin{pmatrix} \sum_{b \in B} R_c(r_{ab}) \hat{r}_2 V_{bc3}^{(l=1)} - \sum_{b \in B} R_c(r_{ab}) \hat{r}_3 V_{bc2}^{(l=1)} \\ \sum_{b \in B} R_c(r_{ab}) \hat{r}_3 V_{bc1}^{(l=1)} - \sum_{b \in B} R_c(r_{ab}) \hat{r}_1 V_{bc3}^{(l=1)} \\ \sum_{b \in B} R_c(r_{ab}) \hat{r}_1 V_{bc2}^{(l=1)} - \sum_{b \in B} R_c(r_{ab}) \hat{r}_2 V_{bc1}^{(l=1)} \end{pmatrix}$$
(11)

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Data Set	Cutoff	== 888
MD-17 [17, 30, 31]	4.0	889
Molecules, CCSD/CCSD(T) [31]	4.0	009
Water+Ices, DeepMD [15, 32]	6.0	890
Formate on Cu	5.0	891
$\text{Li}_4\text{P}_2\text{O}_7$ [12]	5.0	031
LiPS [13]	5.0	892
Water, data efficiency tests [39]	4.5	893

TABLE VII: Radial cutoff in units of [Å].

DATA AVAILABILITY

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The code and data sets will be made available upon $_{_{900}}$ 884 publication. 885

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AUTHOR CONTRIBUTIONS

S.B. initiated the project, conceived the NeguIP model, implemented the software and conducted all software experiments under the guidance of B.K. T.E.S. contributed to the conception of the model, guidance of computational experiments, and the software implementation. L.S. created the data set for formate/Cu, guided work on training and MD simulations on this system, and contributed to development of the software implementation. J.P.M. guided the work on the LiPS conductor and implemented the thermostat for MD simulations together with S.B.. M.K. created the AIMD data set of Li₄P₂O₇, wrote software for the analysis of MD results and guided the benchmarking on this system. N.M. wrote software for the computation of the diffusion results and guided discussions on the interpretation of results. B.K. supervised the project from conception to design of experiments, implementation, theory, as well as analysis of data. All authors contributed to the manuscript and the discussion of results.

COMPETING INTERESTS

The authors declare no competing interests.

Figures

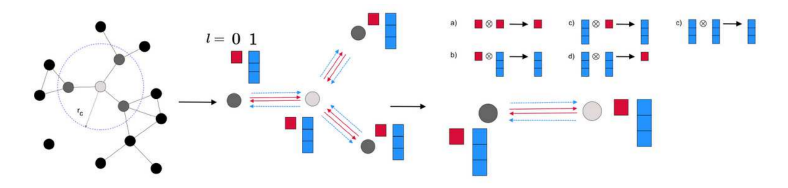


Figure 1

Left: a set of atoms is interpreted as an atomic graph with local neighborhoods. Middle: every atom carries a set of scalar and vector features with it. Right: atoms exchange information via filters, that are again scalars and vectors. The interactions of features and filters define five interactions.

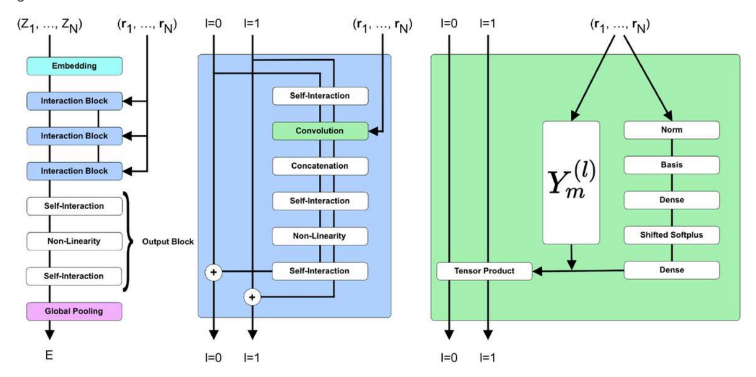


Figure 2

The NequIP network architecture. Left: atomic numbers are embedded into I = 0 features, which are refined through a series of interaction blocks, creating I = 0 and I = 1 features. An output block generates atomic energies, which are pooled to give the total predicted energy. Middle: the interaction block consists of a series of convolutions, interweaved with self-interaction layers, equivariant nonlinearities and concatenation. Right: the convolution combines the radial function R(r) which operates only on interatomic distances with the spherical harmonics based on unit vector \mathbf{r} via a tensor product.

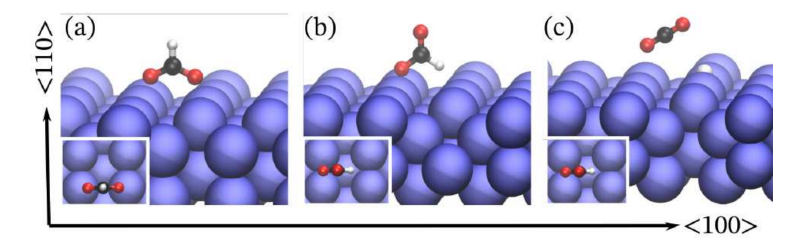


Figure 3

Perspective view of atomic configurations of (a) bidentate HCOO (b) monodentate HCOO and (c) CO2 and a hydrogen adatom on a Cu(110) surface. The blue, red, black, and white spheres represent Cu, O, C, and H atoms, respectively. The subset shown in each subplot is the corresponding top view along the < 110 > orientation.

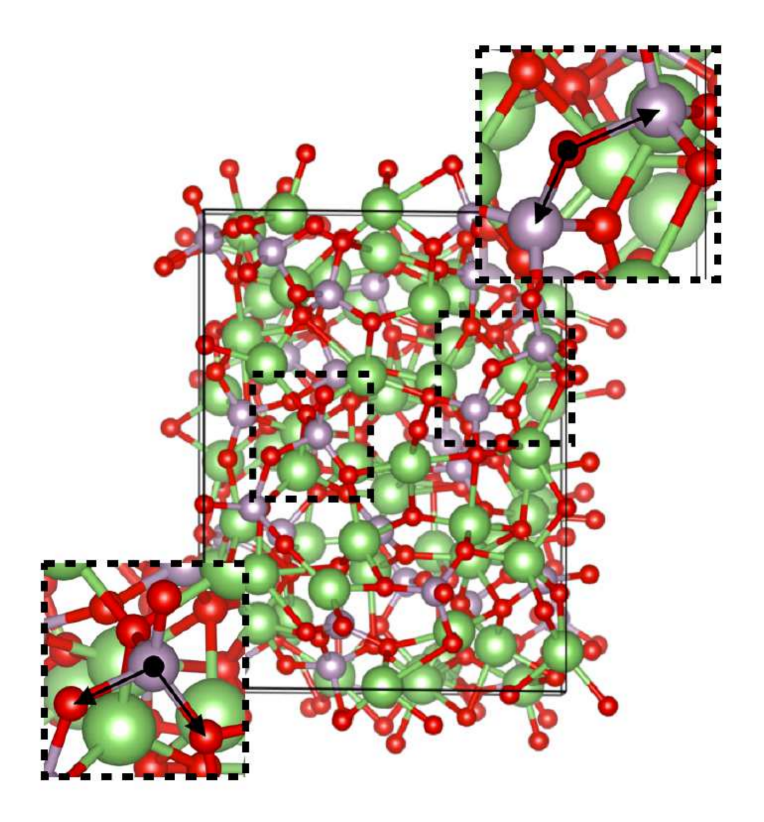


Figure 4

Quenched glass structure of Li4P2O7. The insets show the P-O-O tetrahedral bond angle (bottom left) as well as the O-P-P bridging angle between corner-sharing phosphate tetrahedra (top right).

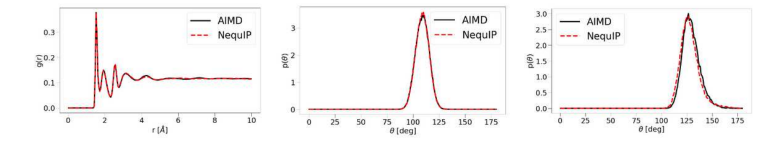


Figure 5

Left: Radial Distribution Function, middle: Angular Distribution Function, bridging oxygen, right: Angular Distribution Function, tetrahedral bond angle. All are defined as probability density functions.

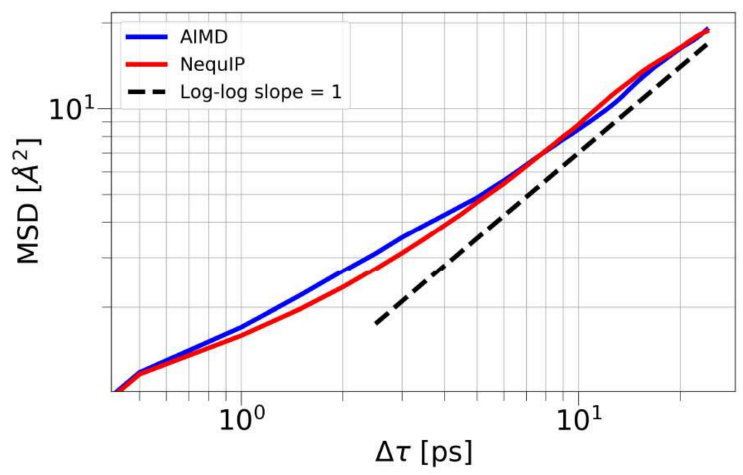


Figure 6

Comparison of Lithium mean square displacement of AIMD and NequIP trajectories.

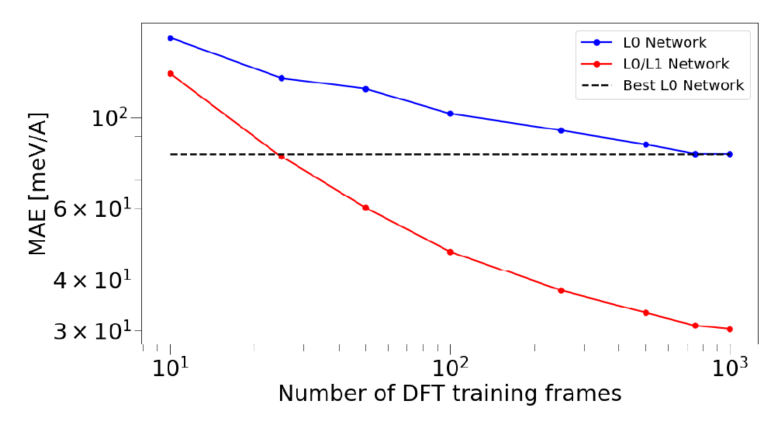


Figure 7

Log-log plot of the predictive error in forces of NequIP with I = 0 vs. I = 0/I = 1 interactions as a function of data set size, measured via the force MAE.

Supplementary Files

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