Abstract

Materials design enables technologies critical to humanity, including combating climate change with solar cells and batteries[1,2,3]. Many properties of a material are determined by its atomic crystal structure. However, prediction of the atomic crystal structure for a given material’s chemical formula is a long-standing grand challenge that remains a barrier in materials design. We investigate a data-driven approach to accelerating ab initio random structure search (AIRSS)[4,5], a state-of-the-art method for crystal structure search. We build a novel dataset of random structure relaxations of Li-Si battery anode materials using high-throughput density functional theory calculations. We train graph neural networks to simulate relaxations of random structures. Our model is able to find an experimentally verified structure of Li$_{15}$Si$_4$ it was not trained on, and has potential for orders of magnitude speedup over AIRSS when searching large unit cells and searching over multiple chemical stoichiometries. Surprisingly, we find that data augmentation of adding Gaussian noise improves both the accuracy and out of domain generalization of our models.

1 Introduction

The atomic structure of a material determines its physical and chemical properties. For instance, graphite (a dark, soft, conductor) and diamond (a transparent, hard, insulator) both consist of carbon atoms that are arranged in different structures. In crystalline solids, atoms are arranged in patterns that repeat periodically, called unit cells. Crystal structure search - predictions of unit cells that may be experimentally observed - is a long-standing problem in materials science[6]. The number of possible configurations for a unit cell can be very large. A unit cell with volume $10 \text{ Å}^3$ and 10 atoms already has $10^{25}$ possible configurations (see appendix), and the number of possible structures increases exponentially with the number of atoms and atom types. Materials Project[7], one of the most widely used databases of materials, has 29.5 atoms per cell on average. But among the large number of possible structures, or local minima in the energy landscape, only the global minimum and a few lowest-energy local minima close in energies may be observed in nature.

Crystal structure search usually requires the use of computationally expensive density functional theory (DFT) calculations. Interatomic potentials that give fast approximations to these calculations often generalize poorly, meaning that potentials trained on a few structures fail to describe another structure that is experimentally observed[8,9]. This is in contrast to molecule discovery, where the structure of most small molecules are determined by atomic bonds and interatomic potentials work well for structure search[10].
In this paper, we build a novel dataset of structure relaxations from random initial states in the Li-Si system via high-throughput DFT calculations, and train graph networks to reproduce the relaxation trajectory. The Li-Si system has at least 7 crystal structures\cite{11} with different stoichiometries that have been experimentally verified\cite{12, 13, 14, 15, 11, 16, 17}, and is widely studied for applications in lithium ion batteries\cite{18, 19, 20}. Compared to molecular dynamics simulations of crystalline Li-Si phases, the random initial configurations yield relaxations that cover a wide range of high-stress unit cells not observed in molecular dynamics even at high temperatures (see appendix).

Once trained, our graph networks enable random structure search with a fraction of the computational time used for ab initio random structure search methods, and benefit from better out-of-domain generalization to unseen stoichiometries. We improve out-of-domain generalization, i.e. accuracy on data that is not from the training set distribution, using data augmentation. In computer vision, there has been evidence of Gaussian noise increasing out-of-domain accuracy at the cost of in-domain accuracy\cite{21} whereas another work showed that it improved interpretability\cite{22}. This behavior has been attributed to reduced sensitivity to high frequency noise\cite{23, 24}. In particle-based simulations, Gaussian noise was observed to decrease roll-out error at the cost of in-domain error\cite{25}. In contrast, we show that adding Gaussian noise to atomic positions help in calculating accurate forces and simulating relaxation trajectories for both in-domain and out-of-domain stoichiometries.

We summarize our contributions below:

- We build a novel dataset of random structure relaxations using density functional theory.
- We update the graph network architecture to be able to output stress measurements. We evaluate the effects of Gaussian noise as data augmentation for both positions and stress.
- We show that our approach can generalize to new stoichiometries and speed up random structure search. Compared to DFT-based random structure search, our method is faster by orders of magnitude.
- We ablate several components of our approach and compare to traditional force field baselines to see where the improvements are coming from.

## 2 Related Work

Crystal structure prediction is an optimization problem in the structure-energy landscape. Popular methods include random sampling\cite{5, 4}, evolutionary algorithms\cite{26, 27, 28}, simulated annealing\cite{29} and basin hopping\cite{30, 31}. These methods involve at least hundreds of computationally expensive ab initio calculations\cite{32}, which take hours on a supercomputer. Elemental substitutions from libraries of known crystal structures\cite{33, 34} is another technique that requires only a few calculations. To address the problem of slow ab initio calculations, some researchers build faster approximations to energies and forces\cite{35}, including graph neural networks\cite{36, 37, 38, 39, 40, 41, 42}. A combination of random structure searching and machine learned potentials has been proposed for phosphorus system in\cite{9, 43}. The Li-Si system has been studied with machine learned potentials\cite{44, 45, 46}, genetic algorithms\cite{47}, embedded atom methods\cite{48}, and random sampling\cite{49}.

In computer vision, data augmentation\cite{50, 24, 23} has been used for improving the accuracy of image classification models, and work on adversarial examples discusses model robustness against distribution shifts\cite{51, 52, 53}. Recent work on adversarial examples in physical systems\cite{54} showed that models approximating physical systems move in their own adversarial directions, which might explain why adding Gaussian noise was advantageous in this work.

## 3 Implementation

### 3.1 Dataset of Random Structure Relaxations

We use AIRSS\cite{4, 5} to generate 5000 random unit cells for each of: LiSi, Li$_2$Si, Li$_7$Si$_2$, Li$_{13}$Si$_4$, and Li$_{15}$Si$_4$. AIRSS constructs random structures by choosing a set of random unit cell lengths and angles, and placing the atoms in random locations in the unit cell. Additional constraints on the cell volume, minimum separation between atoms and symmetry may also be imposed on the cell as described in\cite{4, 5} and in the appendix. We use the Vienna Ab initio Simulation Package (VASP)\cite{55} to perform DFT relaxations on the random unit cells. The total number of converged random structure relaxations...
for each stoichiometry were 4680(LiSi), 4365(Li$_{13}$Si$_4$), 4596(Li$_2$Si), 4515(Li$_7$Si$_2$) and 4199(Li$_{15}$Si$_4$). For all stoichiometries except Li$_7$Si$_2$, experimentally reported structures were recovered.

49.5% of the relaxation trajectories match the converged final structure by the time the maximum force acting on the atoms reaches 0.1 eV/Å, where ‘match’ is defined by pymatgen’s StructureMatcher. StructureMatcher compares two structures by reducing them to primitive cells and evaluating whether the maximum RMS displacement is less than a predefined tolerance cutoff (stol), see §57. We split the data into high-force (maximum force on atoms > 0.1 eV/Å) and low-force (maximum force on atoms < 0.1 eV/Å) regions. Since our goal is to quickly screen random structures to find the few lowest energy ones, we focus our discussion on learning energy relaxations from the high-force dataset as computed using a steepest descent algorithm.

We construct our training set by sampling structures from the relaxation trajectory from regions where the maximum force acting on the atoms is larger than 0.1 eV/Å. We sample at most one structure out of every five ionic steps. We only sample structures that do not match the previous structure in the trajectory. Again, we used StructureMatcher with stol=0.05. After sampling, we use 80% of the trajectories for training, 10% each for validation and test. For trajectory predictions, we use all ionic steps until the maximum force in the crystal reaches 0.1 eV/Å.

### 3.2 Dataset of Molecular Dynamics Simulations

We take the experimental crystalline structures of LiSi, Li$_2$Si, Li$_7$Si$_2$, Li$_{13}$Si$_4$ and Li$_{15}$Si$_4$ from the Materials Project and perform molecular dynamics(MD) simulations with VASP. We create 64 to 76-atom supercells of the conventional structure in the Materials Project, and ran the simulations at 800K for at least 47 picoseconds. We used StructureMatcher to discard matching structures, and end up with 17134(LiSi), 16088(Li$_{13}$Si$_4$), 9613(Li$_2$Si), 13079(Li$_7$Si$_2$) and 12749(Li$_{15}$Si$_4$) structures for each stoichiometry.

### 3.3 Graph Neural Networks (GNN)

We use graph neural nets to compute the forces on each atom and the stress tensor on the unit cell, with the architecture based on MEGNet[38]. Each node is initialized to states corresponding to the atomic number of the element, and a 16-dimensional embedding layer is applied. Edges are constructed for every pair of atoms within 4Å, and each edge is initialized to the displacement vector between the atoms, distance, and distance squared ($r_{ij}$, |$r_{ij}$|, |$r_{ij}$|^2), where $r_{ij} = r_i - r_j$. Global states are initialized to 0. Each state is passed through 7 message-passing layers as per in MEGNet, consisting of two MLP layers of size 64 and state updates. To get forces on each atom, we concatenate node features(with two MLP layers of size 64 and 32) with global features and follow with a output layer without activation.

For stress tensor predictions, we put fictitious non-interacting atoms on (100), (010) and (001) planes. We used the vector distances from the planes to each atom to compute edge features. Each fictitious atom outputs three stress tensor components - e.g. the atom on the x-axis outputs $\sigma_{xx}$, $\sigma_{yx}$, $\sigma_{zx}$.

The force and stress inputs were scaled to have mean 0 and standard deviation 1. We used the sum of mean squared errors for forces and stresses for our loss function. For data augmentation, we rotate 50% of the structures at each epoch by 90 degrees, and perturb the atom positions and lattice vectors with Gaussian noise for 30% of the structures. The models were trained with ADAM optimizer with learning rate $10^{-3}$, and the learning rate was halved every 15 steps. Using the force and stress outputs, we simulate relaxations using the velocity verlet equations as implemented in VASP with details in the appendix.

As a baseline, we train another model on predicting crystal energies with a similar architecture, and differentiate the energies to get forces and stresses. After the message-passing layers, the node and edge states are aggregated with set2set[58] and concatenated with the global state, and passed through two MLP layers of sizes 64 and 32. The final predictions are produced by an output layer without activation. The energy inputs were also scaled to have mean 0 and standard deviation 1, and sum of

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1. we investigated three architectures and chose the one with best performance. Appendix section Architectures for stress calculations

3
mean squared errors was used as loss function. We used rotation augmentation for training the energy model.

4 Experiments

We train the models below on the Li-Si system, holding out Li$_{15}$Si$_4$ for out-of-domain generalization experiments.

4.1 Impact of data augmentation

We investigate the effects of adding Gaussian noise to atom positions and lattice vectors. When simulating relaxation trajectories, errors may accumulate over multiple time steps. Adding small perturbations during training may make the simulations more robust over longer trajectories.[25]

As a baseline, we use the second nearest-neighbor modified embedded atom method (2NN MEAM) from ref.[59]. We evaluate the force and stress prediction accuracy by mean absolute error on the test set sampled as described above. We simulate random structure relaxations with predicted force and stress outputs using the same algorithm as VASP and use pymatgen’s StructureMatcher to compare the final relaxed structures. Additionally, we match structures at each step in the VASP and GNN trajectories and report the fraction of structures that match.

We evaluate model performance using metrics for force and stress prediction accuracy, and for producing accurate relaxation trajectory in Table 1. All GNN models outperform MEAM by an order of magnitude difference in both force and stress MAE. In contrast to ref.[25] where Gaussian noise helped simulate longer trajectories at the cost of force errors at each step (one-step MSE in their paper), in our system adding Gaussian noise leads to more accurate force and stress outputs and better relaxation trajectories. While the force and stress outputs are the most accurate with noise of 0.01 eV/Å, the trajectory prediction improves with a larger noise of 0.02 eV/Å. The advantages of Gaussian noise decrease with larger amounts of noise.

Table 1: Gaussian noise improves both force predictions and relaxation trajectories. The units for Gaussian noise standard deviation, force and stress are Å, eV/Å and kBar. We averaged the results from three random seeds. Quantities in parentheses are the standard deviations from different random seeds. The last row uses MEAM from [48]. *: Fraction of relaxed final structures that match VASP **: Fraction of matching structures in each trajectory

| Model | Gaussian noise (std. dev) | Force MAE | Stress MAE | Relaxed structure match* | Matches in trajectory** |
|-------|---------------------------|-----------|------------|--------------------------|-------------------------|
| GNN 0  | 0.033 (2.1 × 10$^{-4}$)   | 1.05 (0.021) | 0.40 (0.015) | 0.65 (0.015) |
| GNN 0.01 | **0.031** (4.5 × 10$^{-4}$) | **0.98** (0.047) | 0.42 (0.007) | **0.67** (0.006) |
| GNN 0.02 | 0.032 (1.4 × 10$^{-4}$) | 1.01 (0.023) | 0.43 (0.005) | **0.67** (0.000) |
| GNN 0.03 | 0.034 (1.9 × 10$^{-4}$) | 1.06 (0.050) | 0.42 (0.007) | 0.66 (0.006) |
| MEAM 0 | 0.55 | 9.11 | 0.18 | 0.39 |

4.2 Ablation experiments on dataset and model architecture

Compared to the MEAM baseline, our approach uses a more expressive model family (GNN) and a richer dataset. In order to determine which factors contribute most to our model’s superior performance, we perform two ablation experiments.

Baseline 1. We train the same GNN architecture on a separate dataset of structures from a molecular dynamics trajectory.

Baseline 2. We train the GNN on the same dataset of random structure relaxations, but train on energies instead of forces and stresses. The model trained on random structure energies achieve a MAE of 15 meV/atom with more details in the appendix.

The results are in Table 2. Our architecture trained on random structure data shows an order of magnitude improvement against both baselines. The MD baseline suggests that the distribution of
data generated by MD is quite different (see appendix) and training on random structures is crucial for crystal structure search.

Table 2: Using random structure relaxations with our architecture for directly predicting forces and stresses (GNN) leads to an order of magnitude improvement in accuracy over the physics-based interatomic potential MEAM. First row is the result from Table 1 without Gaussian noise augmentation. RR = random relaxations.

| Model (baseline) | Trained on | Training data | Test data | Force MAE | Stress MAE |
|------------------|------------|---------------|-----------|-----------|------------|
| GNN              | forces, stresses | RR | RR | 0.033 | 1.05 |
| GNN (1)          | forces, stresses | MD | MD | 0.067 | 1.63 |
| GNN (1)          | forces, stresses | MD | RR | 0.61 | 32.3 |
| GNN (2)          | energies    | RR | RR | 0.24 | 18.7 |
| MEAM             | -           | - | RR | 0.55 | 9.11 |

4.3 Out-of-domain generalization and structure search of Li₁₅Si₄

We trained the models on four stoichiometries in the Li-Si system - LiSi, Li₂Si, Li₇Si₂, Li₁₃Si₄. We tested the model on 4199 random structure relaxations of Li₁₅Si₄ to see if it recovers the experimentally reported unit cell of Li₁₅Si₄. The experimental structure of Li₁₅Si₄ has 38 atoms in the unit cell, which is more than the other stoichiometries used in training. This makes it a challenging task as the number of possible configurations scales exponentially with the number of atoms in the cell. Again, we compare it to the 2NN MEAM baseline. Out of the 4199 random structure relaxations using DFT, three of the structures relaxed into the same experimental structure reported in [13], and the structures relaxed to the force cutoff of 0.1 eV/Å match the final relaxed structures. All GNN models reproduced the experimental structure trajectories, and MEAM did not find the experimental structure, as shown in Table 3. This is surprising since Li₁₅Si₄ was included in the data used to fit MEAM. It suggests that for random structure relaxations, the behavior of the force field model at high-stress regions may relax the structure into the wrong basin in the energy landscape and training on this region may be crucial for random structure search.

Table 3: Generalization of the model on Li₁₅Si₄. *: whether relaxations with the model found the experimental structure of Li₁₅Si₄

| Model   | Gaussian noise (std. dev) | Relaxed structure match | Matches in trajectory | experimental* |
|---------|---------------------------|-------------------------|-----------------------|---------------|
| GNN     | 0                         | 0.35                    | 0.64                  | yes           |
| GNN     | 0.01                      | 0.38                    | 0.66                  | yes           |
| GNN     | 0.02                      | **0.39**                | **0.67**              | yes           |
| GNN     | 0.03                      | **0.39**                | 0.66                  | yes           |
| MEAM    | -                         | 0.19                    | 0.39                  | no            |

VASP simulations of Li₁₅Si₄ structures took 38 hours on average on Google Cloud’s standard N1 machine with 8 cores. GNN simulations take less than a minute on a laptop (MacBook Pro (15-inch, 2017)), and is still able to find the experimental structure of Li₁₅Si₄.

5 Conclusion

We build a novel dataset of DFT random structure relaxations, and show that graph network models trained on this data has significant advantages for crystal structure search. Moreover, we show that using Gaussian noise in training force field models improve accuracy in force predictions, simulating trajectories and out-of-domain generalization. It would be interesting to draw on the literature on adversarial defense and explore how it can be used to improve physical models.

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2 visualization of the unit cell is in the appendix
3 number of atoms in the unit cell of experimental structures - LiSi: 16, Li₂Si:6, Li₇Si₂:36, Li₁₃Si₄:34
6 Acknowledgements

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Appendix

7.1 Estimation of number of configurations in unit cells

The number of configurations for placing \( N \) atoms in a cell of volume \( V \), discretizing the cell by length \( \delta \), is

\[
\frac{1}{N!} \left( \frac{V}{\delta^3} \right)! \left( \frac{V}{\delta^3} - N \right)!.
\]

For \( N = 10 \), \( V = 10\,\text{Å}^3 \) and \( \delta = 0.1\,\text{Å} \), this number is \( 2.7 \times 10^{25} \).

7.2 VASP simulation details

**Random structures.** We used AIRSS to generate random structures with up to 6 symmetry operations, and vary the number of formula units in the cell up to 38 atoms in the cell. We estimate the volume per element from the experimental data of Li–Si structures in the Materials Project, and set the target volume of AIRSS structure outputs to be within 80% 150% of the estimated volume. We set minimum separation between atoms to be 2.28 Å, as small values of atomic separation may cause forces to diverge. The random structures were relaxed using Kohn-Sham density functional theory with the projector augmented-wave method\(^{[61]}\) as implemented in VASP. Relaxations were performed with steepest descent algorithm (IBRION=3, SMASSE=2, POTIM=0.4) to relax both the ions and the unit cell(ISIF=3). The calculations were converged to energy convergence of \( 10^{-6} \) eV and force convergence of \( 10^{-2} \) eV/Åwith self-consistent, periodic DFT. A plane wave basis set with
the kinetic energy cutoff of 520 eV was used, and Γ-centered k-point mesh with spacing 0.25 Å⁻¹ was used to sample the Brillouin zone. Generalized gradient approximation Perdew-Burke-Ernzerhof (PBE) functional was used to treat the exchange-correlation energy.

**Molecular Dynamics.** We used the structures from Materials Project flagged as having an experimentally characterized structure, and created 64 to 78-atom supercells of the conventional cells of these structures to ensure the unit cells are sufficiently large for molecular dynamics. (Li₁₅Si₄: mp-569849, 76 atoms. Li₁₃Si₄: mp-672287, 68 atoms. Li₂Si: mp-27705, 72 atoms. Li₇Si₂: mp-27930, 72 atoms. LiSi: mp-795, 64 atoms) The simulations were run at a constant temperature of 800K with 2 femtosecond time steps with a single k-point at Gamma. All other details are the same as random structures.

### 7.3 Distribution of random structure relaxations data and MD data

In the experiments in the main text, models were trained on all stoichiometries except for Li₁₅Si₄, which was held out for generalization experiments. The distributions below are also plotted excluding Li₁₅Si₄. We see that the random structures have higher stresses than MD data, which is expected from random structures. The forces for random structures, however, are smaller than MD data. We enforced a moderately large minimum separation between atoms for random structure generation to ensure that none of the random structures diverged during relaxations, and moved atoms in the direction of forces during the relaxations. This led to a narrower distribution of forces in the random structure dataset.

![Figure 1: Force(left) and stress(right) distribution of MD data.](image1)

![Figure 2: Force(left) and stress(right) distribution of random structure relaxations data.](image2)

### 7.4 Architectures for stress calculations

We investigated three different architectures below:

**Stress tensor outputs as an global outputs.** The 6 independent components of the stress tensor($\sigma_{xx}, \sigma_{yy}, \sigma_{zz}, \sigma_{xy}, \sigma_{yz}, \sigma_{zx}$) are predicted from the graph. The node and edge states are aggregated with set2set and concatenated with the global state, and passed through two MLP
Table 4: Details of the random structure dataset. The first two rows correspond to the components of force vectors and stress tensors. Max |force| refers to the maximum force magnitude in each structure. Max stress component is the maximum value of the stress tensor components (σ_{xx}, σ_{yy}, σ_{zz}, σ_{xy}, σ_{yz}, σ_{zx}) in each structure.

|                      | mean  | standard deviation | min   | max   |
|----------------------|-------|--------------------|-------|-------|
| Force (eV/Å)         | 0.000 | 0.208              | -4.797| 4.797 |
| Stress (kBar)        | 0.733 | 12.318             | -192.012| 439.783|
| max |force| (eV/Å)     | 0.453 | 0.442 | 0.000  | 4.797  |
| max stress component (kBar) | 8.662 | 23.555             | 0.000  | 439.783|

Table 5: Methods for predicting cell stress. Corner and plane refer to stress outputs from putting fictitious atoms on cell corners and cell planes, and MEAM is the model from ref.[59]. Force and stress units are eV/Å and kBar. *: Fraction of relaxed final structures that match VASP **: Fraction of matching structures in each trajectory

| Method         | Force MAE | Stress MAE | Relaxed structure match* | Matches in trajectory** |
|----------------|-----------|------------|--------------------------|-------------------------|
| corner         | 0.03      | 3.09       | 0.15                     | 0.34                    |
| planes         | 0.03      | 1.05       | **0.41**                 | **0.66**                |
| global output  | 0.04      | 0.96       | 0.36                     | 0.61                    |
| MEAM           | 0.55      | 9.11       | 0.23                     | 0.39                    |

layers of sizes 64 and 32. The final predictions are produced by an output layer without activation. For this model, the distance from each atom to (100), (010) and (001) planes were concatenated to the node feature inputs.

Stress tensor outputs from fictitious atoms on cell corners. We put fictitious non-interacting atoms on the corners of the cells, and predict the stress tensor components from the fictitious atoms. Each fictitious atom outputs three stress tensor components - e.g. the atom on the x-axis outputs σ_{xx}, σ_{yx}, σ_{zx}.

Stress tensor outputs from fictitious atoms on cell planes. We put fictitious non-interacting atoms on (100), (010) and (001) planes, and use distances from the planes to each atom for edge features.

Table 5 shows that stress predictions from fictitious atoms on cell planes and as global output perform similarly on force and stress MAE, but fictitious atoms on cell corners performs poorly on stress MAE. These two models outperform MEAM in force predictions by an order of magnitude difference and by a factor of three in stress MAE. Moreover, fictitious atoms on cell planes is the best at predicting the relaxation trajectory, with 77% improvement over the MEAM baseline. We used fictitious atoms on cell planes for all experiments in the main text.

7.5 Energy predictions on random structure data

The model trained on random structure energies achieved a MAE of 15 meV/atom. This is higher than other machine learning potentials on the Li-Si systems, such as [45, 44, 46]. As random structure data is quite different from what [45, 44, 46] are trained on, we trained and tested the same model on our MD data. The test MAE for MD energies is 3.8 eV/atom, similar to what’s reported in these works.

7.6 Relaxation algorithm affects noise in relaxation data

We used the steepest descent algorithm as implemented in VASP (IBRION=3, SMASS=2, ISIF=3) and adjusted the time step parameter (POTIM) to a small value (0.4, default is 0.5) and checked all relaxations have a monotonic decrease in energy over ionic steps. But in some cases, we see large upward spikes in force or stress trajectory, as in Figure 3. To investigate whether these spikes introduce noise in our simulations, we quantified the amount of upward spikes in the relaxation trajectories that are successfully simulated by our model and those that fail. We calculated the maximum amount of increase in forces (maximum force applied to the atoms in the structure) and
Figure 3: Sample trajectory of Li2Si in which GNN model did not reproduce the VASP relaxation trajectory. The red dot corresponds to where the maximum force on the atoms reach 0.1 eV/Å and the simulation terminates. The green dot is the ionic step until the GNN simulation matches VASP. There are large spikes in atomic forces in this trajectory.

Figure 4: The distribution of upward spikes in relaxation trajectory for maximum forces on atoms (top) and maximum component of stress tensor (bottom). The trajectories where our graph network simulations match the VASP trajectory are plotted in blue (match), and the trajectories where our simulations do not match are plotted in yellow (no match).

stresses (maximum component of the stress tensor) within 5 ionic steps for Li2Si structures in the test set. We plot the distribution of those spikes in Figure 4. We see that for both forces and stresses, the distribution of spikes is generally larger for the trajectories that our model fails to simulate correctly. The distribution suggests that some of our model’s failures may attributed to unphysical behavior in DFT simulations with VASP used to construct the ground-truth data.

7.7 Experimental structure of Li15Si4

The experimental structure of Li15Si4 [13] is visualized in Figure 5.
Figure 5: Experimental structure of Li$_{15}$Si$_4$