Machine Learning-Assisted Discovery of Solid Li-Ion Conducting Materials

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* Supporting Information

ABSTRACT: We discover many new crystalline solid materials with fast single crystal Li ion conductivity at room temperature, discovered through density functional theory simulations guided by machine learning-based methods. The discovery of new solid Li superionic conductors is of critical importance to the development of safe all-solid-state Li-ion batteries. With a predictive universal structure−property relationship for fast ion conduction not well understood, the search for new solid Li ion conductors has relied largely on trial-and-error computational and experimental searches over the last several decades. In this work, we perform a guided search of materials space with a machine learning (ML)-based prediction model for material selection and density functional theory molecular dynamics (DFT-MD) simulations for calculating ionic conductivity. These materials are screened from over 12 000 experimentally synthesized and characterized candidates with very diverse structures and compositions. When compared to a random search of materials space, we find that the ML-guided search is 2.7 times more likely to identify fast Li ion conductors, with at least a 44 times improvement in the log-average of room temperature Li ion conductivity. The F1 score of the ML-based model is 0.50, 3.5 times better than the F1 score expected from completely random guesswork. In a head-to-head competition against six Ph.D. students working in the field, we find that the ML-based model doubles the F1 score of human experts in its ability to identify fast Li-ion conductors from atomistic structure with a 1000-fold increase in speed, clearly demonstrating the utility of this model for the research community. In addition to having high predicted Li-ion conductivity, all materials reported here lack transition metals to enhance stability against reduction by the Li metal anode and are predicted to exhibit low electronic conduction, high stability against oxidation, and high thermodynamic stability, making them promising candidates for solid-state electrolyte applications on these several essential metrics.

I. INTRODUCTION

All-solid-state Li-ion batteries (SSLIBs) hold promise as safer, longer lasting, and more energy dense alternatives to today’s commercialized LIBs with liquid electrolytes. However, the design of SSLIBs remains a challenge, with the principal technological bottleneck in realizing these devices being the solid electrolyte. A high performance solid electrolyte material must satisfy several criteria simultaneously: it must possess fast Li-ion conduction, negligible electronic conduction, a wide electrochemical window, robust chemical stability against side reactions with the electrodes, and high mechanical rigidity to suppress dendritic growth on the anode. The material should also be cheap and easy to manufacture. Given these many constraints, searching for promising new materials that satisfy all requirements through the trial-and-error searches has yielded slow progress.

The earliest efforts to discover fast Li-ion conducting solids began in the 1970s1 and have continued to present. More recently, density functional theory (DFT) simulation has enabled high-throughput computational searches, essentially automating the process of guess-and-check.2,3 Across these four decades, only several solids with liquid-level Li conduction (>10^{-2} S/cm) at room temperature (RT) have been identified, including notably Li10GeP2S12 (17 mS/cm)4 and Li7P3S11 (25 mS/cm).5 This slow progress suggests that continuing in the guess-and-check paradigm of decades past is unlikely to quickly yield the material innovations we need to unlock the high...
energy density, high cycle life, and unquestionably safe energy storage devices of the future.

Leveraging atomic and electronic structure data from the Materials Project database, we have screened all 12 000+ Li-containing materials for thermodynamic phase stability, low electronic conduction, high electrochemical stability, and no transition metals (to enhance stability against reduction). We also compiled information on the estimated raw materials cost and the earth abundance of the elemental constituents of each material. This identifies 317 materials that may be strong candidate electrolyte materials if they are also fast ion conductors.

Following the guess-and-check paradigm, one would begin to test these materials for fast ion conduction at random, or according to his/her best scientific intuition. To identify the subset of these materials most likely to exhibit fast ionic conductivity, we have instead developed a machine learning (ML)-based model for predicting the likelihood \( P_{\text{superionic}} \) that an arbitrary material exhibits fast Li-ion conduction at RT, based only on features \( x_i \) derived from the atomistic structure of the unit cell. Throughout this work, we define superionic conductivity to be greater than 0.1 mS/cm, based on the approximate minimum electrolyte ionic conductivity required for battery applications. Experimental reports of ionic conductivity for several dozen materials ranging over 10 orders of magnitude were used to train the model. This data-driven prediction takes the form of a logistic function,

\[
P_{\text{superionic}}(x) = \left(1 + e^{-\sum \Theta x_i}ight)^{-1},
\]

where

\[
\sum \Theta x_i = 0.18 \times \text{LLB} - 4.01 \times \text{SBI} - 0.47 \times \text{AFC} + 8.70 \times \text{LASD} - 2.17 \times \text{LLSD} - 6.56
\]

Here, LLB is the average Li–Li bond number (number of Li neighbors) per Li; SBI is the average sublattice bond ionicity; AFC is the average coordination of the anions in the anion framework; LASD is the average equilibrium Li–anion separation distance in angstroms; and LLSD is the average equilibrium Li–Li separation distance in angstroms. Since this model does not require any electronic structure information, it is >5 orders of magnitude faster to evaluate than a DFT simulation of conductivity.

Screening this list of 317 candidate materials identifies 21 crystalline compounds that are predicted to be fast ion conductors with robust structural and electrochemical stability, representing a 99.8% reduction in the entire space of known Li-containing materials. One of these 21 materials, LiCl, has been reported to exhibit poor RT Li conduction \((\sim 10^{-9} \text{ S/cm})\), making it a known false positive prediction. Another material, Li\(_3\)InCl\(_6\), has been reported to have a RT conductivity \((\text{starting at } t = 900 \text{ K})\); if melting is observed, the simulation is restarted at increasingly lower temperatures until no melting is observed. The vacancy concentration ranges from 3 to 17% depending on the unit cell. The simulation temperatures, computational cell size, and Li vacancy concentrations are provided in the Supporting Information, Table S1.

We simulate the candidate materials for a range of times on the tens to hundreds of picoseconds time scale; see Table S1. To calculate ionic diffusivity, which may be isotropic, we compute the average of the diagonal elements of the second rank Li diffusivity tensor (or equivalently, one-third of the mean squared displacement (MSD) of the Li atoms \(\Delta r^2\) over time (starting at \(t = 0\)) and apply the following formula:

\[
\langle D_i \rangle = \lim_{t \to \infty} \frac{1}{6} \frac{\langle \Delta r^2 \rangle}{\Delta t}
\]

To probe for melting, we also calculate the MSD of the sublattice atoms and assume melting if sustained, nonzero diffusivity is observed in both Li and the sublattice (the smallest diffusivity that can be resolved through DFT-MD at 900 K is approximately 0.01 Å\(^2\)/ps). To assess the degree of convergence in \(\langle D_i \rangle\), we compute the standard deviation in diffusivities when measured from different starting times in the MSD data. We compute the slope of the MSD data for every starting time from \(t = 0\) to up to 75% of the total run time, in 100 fs increments. We compute one standard deviation above and below the mean diffusivity across all starting times to
represent the approximate upper and lower limits of the distribution of diffusivities one may observe under these simulation conditions. The diffusivity as measured from $t=0$ and the mean diffusivity across all starting times are not necessarily equivalent, and thus the upper and lower uncertainties are not necessarily symmetric around the value predicted from $t = 0$.

Ionic transport in crystalline solids is modeled here as a stochastic phenomenon governed by a Boltzmann (Arrhenius) factor that is exponential in a single energy barrier between equilibrium sites, $E_a$:

$$\langle D_{ii} \rangle = D_0 e^{-E_{ii}/k_BT}$$  \hspace{1cm} (3)

Ionic hopping becomes exponentially less likely as the energy barrier increases, and at 900 K ($k_BT = 78$ meV) hopping may not happen on the tens to hundreds of picoseconds time scale if the barrier is much above 0.4 eV ($e^{-0.4/0.078} = 0.006$). Thus, high temperature MD simulations are not well-suited to predict a numerical value for ionic conductivity in medium- to high barrier systems because the statistical behavior is not captured on typical DFT simulation time scales. Numerical values can only be predicted through MD in low barrier systems, and even then are still approximate due to the stochastic nature of hopping. Our approach is most suited for high throughput calculations in which the main goal is identifying whether or not materials are superionic conductors, i.e., for making binary predictions of superionic vs non-superionic conduction.

From the diffusivity $\langle D_{ii} \rangle$, we convert to the average of the diagonal elements of the ionic conductivity tensor $\langle \sigma_{ii} \rangle$ via the Einstein relation:\textsuperscript{12}

$$\langle \sigma_{ii} \rangle(T) = \frac{\langle D_{ii} \rangle(T) n q^2}{k_BT}$$  \hspace{1cm} (4)

where $n$ is the concentration of Li atoms and $q$ is the charge on the Li atoms. We measure the effective Li charge $q$ using the Bader charge analysis methods of Henkelman et al.\textsuperscript{13–16} Given the linear relationship between conductivity and diffusivity, we compute the spread of possible conductivities by applying eq 4 to the computed bounds in diffusivity as well.

It stands to reason that fast Li conduction will not be observed at RT if it is not observed in these favorable conditions. Our approach therefore should not identify false negatives (materials with poor conduction at high temperature but fast conduction at RT), although there is a risk of identifying false positives (materials with fast conduction at high temperature but poor conduction at RT). To ensure favorable scaling to room temperature and guard against the identification of false positives, we simulate DFT-MD again at alternate temperatures if significant Li diffusion has been observed at the initial simulation temperature. The line connecting the two or three diffusivities on an Arrhenius plot of $\log_{10}(\sigma)$ versus inverse temperature is employed here to extrapolate down to RT without direct calculation of $E_a$. This assumes Arrhenius scaling applies and there are no structural phase changes or nonstructural superionic transitions (where

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**Figure 1.** Flowchart of high-throughput density functional theory molecular dynamics (DFT-MD) simulation process. Materials are chosen either according to the machine learning-based model (left stream) or by random selection (right stream). Materials are initially simulated at 900 K for approximately 50–100 ps. If no Li diffusion is observed on this time scale, the materials are immediately considered poor ion conductors and no further simulation is performed. If materials exhibit Li diffusion at 900 K, they are simulated again at alternate temperatures so a simple two- or three-point Arrhenius extrapolation to room temperature can be made. If melting is observed in the ML-selected materials at 900 K, the simulation is restarted at a lower temperature. Randomly chosen materials that exhibit melting at 900 K are discarded immediately for computational efficiency; this occurred in two randomly chosen materials.
conductivity changes abruptly as new ionic pathways become energetically accessible or inaccessible without any significant sublattice rearrangement)\textsuperscript{17} between the simulation temperatures and RT. The spread of possible values in the RT conductivity extrapolations is computed by extrapolating along the upper and lower limits of the deviations in the high temperature conductivity calculations. Using typical values for Li concentration, Bader charge, and site hopping distance, we estimate that the RT conductivity is likely to be $10^{-9}$ S/cm or lower if no Li diffusion is observed on the simulation time scale at 900 K (see Supporting Information, Section S1 for calculation).

A flowchart describing this process is shown in Figure 1. This extrapolation scheme requires less computational expense than simulating each material at several different temperatures right away and is the most tractable way for us to quantify the ionic conductivity of the materials studied here on a reasonable time scale. Screening for Li superionic RT diffusion by two-temperature extrapolation has been leveraged in recent work to search sulﬁde-based compositional spaces, for example.\textsuperscript{3}

We perform DFT calculations with the Vienna Ab Initio Simulation Package (VASP)\textsuperscript{18} with the generalized gradient approximation (GGA) of Perdew–Burke–Ernzerhof (PBE)\textsuperscript{19} and the project augmented wave (PAW)\textsuperscript{20} method. We use the pseudopotentials and plane wave cutoff energy (520 eV for all structures) as recommended by the Materials Project. The VASP input ﬁles are generated using the pymatgen.io.vasp.sets module of Pymatgen.\textsuperscript{21,22} Given the large unit cells (and DFT VASP input) than simulating each material at several different temperatures right away and is the most tractable way for us to quantify the ionic conductivity of the materials studied here on a reasonable time scale. Screening for Li superionic RT diffusion by two-temperature extrapolation has been leveraged in recent work to search sulﬁde-based compositional spaces, for example.\textsuperscript{3}

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The ultimate metric of the utility of materials selection models is the improvement in likelihood of successfully identifying positive examples over the background probability of these materials. To quantify the superiority of our model to completely random guesswork, we must know the likelihood of discovering a superionic material by chance with no scientiﬁc intuition involved. In 1978, Huggins and Rabenau questioned “whether there is anything fundamentally different in such materials, or whether they merely exhibit extreme values of ‘normal’ behavior.”\textsuperscript{23} Superionic conductors may have remarkably low diffusion barriers, but how do they compare to the distribution of barriers across the space of all the candidate electrolyte materials?

Experience would tell us that the likelihood of chance discoveries is low, given that only a handful of fast Li conductors have been discovered since the search began nearly 40 years ago. However, we are probing for superionic conduction in a very speciﬁc way: we are pulling from a potentially biased group of 317 known materials in the Materials Project database, introducing Li vacancies, and simulating finite computational cells at high temperature. Therefore, this question is best answered by doing a straightforward comparison of high temperature DFT-MD simulations for a similar number of structures chosen at random from the same population.

To accomplish this, we perform a control experiment where we simulate 21 structures chosen uniformly at random from among the 317 materials that satisﬁe all prerequisite screening criteria (band gap > 1 eV, predicted oxidation potential > 4 V, energy above convex hull = 0 eV, no transition metals) and simulate MD under the same procedure. These 21 structures and their predicted superionic likelihoods according to the ML model are the following: LiLa$_2$SbO$_6$ (0%), Li$_2$UO$_6$ (7.7%), LiInF$_4$ (0.6%), LiBiF$_4$ (0.2%), Li$_3$Ho(BO$_3$)$_3$ (10.1%), RbLi$_5$BO$_7$ (4%), Li$_3$Be$_5$As$_3$ClO$_12$ (0.3%), Li$_3$TeO$_6$ (6.4%), Li$_3$Pr$_5$(BO$_3$)$_3$ (9.2%), NaLiS (36%), Li$_2$SiO$_4$ (2.4%), LiCaGa$_6$O$_9$ (0.0%), Li$_2$Te$_5$O$_3$ (15.8%), LiNO$_3$ (8.3%), Ba$_2$Li$_5$(SiO$_3$)$_3$ (0.0%), Rb$_5$Li$_2$SiO$_4$ (4.5%), NaLi$_3$PO$_4$ (0.0%), Cs$_3$Li$_2$(Si$_3$O$_9$)$_3$ (0.1%), RbLi(H$_2$N)$_2$ (24.5%), Cs$_3$Li$_2$TeF$_6$ (0.0%), and Li$_2$SO$_3$F (100%). We chose 29 structures in total but removed eight (Na$_2$Li$_2$NF$_6$, Cs$_3$L$i_2$(HO)$_3$, Li$_2$P$_2$O$_5$, Li$_2$Nd$_2$H$_6$(N$_2$O$_4$)$_3$, Li$_2$La$_2$H$_6$(N$_2$O$_4$)$_4$, Li$_2$P$_2$(H$_2$N)$_2$, Li$_2$H$_2$BrO$_3$, and Li$_2$Pr$_5$(NO$_3$)$_3$) due to either melting at 900 K or due to extremely slow or failed electronic convergence, in some cases due to electronic bandgap closure during the simulation. Of the remaining 21 randomly chosen materials we successfully simulate, 20 have a negative ML-based superionic prediction (below <50%) so we do not expect them to conduct. The only material of these 21 random materials that is predicted by our ML-based algorithm to conduct is Li$_2$SO$_3$F. The computational parameters used in these randomly chosen simulations are also provided in Table S1. This set of randomly drawn materials also provides a test set to explore the ML model performance and provide a test set to explore the ML model performance versus the predictive power of the intuition of human experts; we explore this in Section III, Subsection v.

The ML-chosen materials and randomly chosen materials exhibit notably different distributions in their compositions. For example, compositions containing oxygen are heavily represented in the randomly chosen materials (71.4% of materials) but poorly represented in the ML-chosen materials (14.3%). The former is closely aligned with the distribution of oxygen-containing compositions in the broader pool of 317 candidates (70.0%) that satisfy the constraints including high oxidation potential, which we expect given the randomly chosen materials are drawn from this pool. Similarly, sulfur-containing compositions make up 9.2% of the 317 candidates and 9.5% of the randomly chosen candidates but 57.1% of the ML-chosen materials, indicating a clear preference by the ML model for sulﬁdes over oxides when ionic conductivity is the primary consideration. We note that this work is focused primarily on ﬁnding the best speciﬁc target compounds; a follow-up study\textsuperscript{24} will focus on the identiﬁcation of promising new target systems.

In this work we report a total of 4.3 ns of molecular dynamics simulation, with a mean simulation time of 74.9 ps and mean computational cell size of 99.1 atoms. The total volume of data reported here corresponds to approximately 330 000 GPU-hours of simulation.

III. RESULTS AND DISCUSSION

i. Discoveries from ML-Guided Selection. The computed ionic conductivities for the ML-chosen candidate materials and the randomly chosen materials are provided in the Arrhenius diagram on Figure 2a. The computed high temperature diffusivities, ionic conductivities, average Li Bader charge, and extrapolated RT ionic conductivities with predicted deviations (conductivities measured from alternate starting time points) of these candidates are given in Table 1. We do not calculate results for one of the ML-chosen candidates, Li$_2$GePbS$_6$ because of problems with electronic convergence in our simulations.

Our simulations show 10 of the simulated candidate materials exhibit signiﬁcant Li conduction at high temperature. After applying Arrhenius scaling and extrapolating to RT, eight of these materials are predicted to exhibit superionic
Li\(_3\)ErCl\(_6\), Li\(_3\)InCl\(_6\), Li\(_2\)HIO, and LiSO\(_3\)F. Two materials, at 800 K. Of these promising materials that were simulated at 900 K, all have diffusivities significantly above 0.01 Å\(^2\)/ps. The two marginal conductors are above this limit by a factor of two to three, while the best conductors are above this limit by a factor of 10 or more. This provides additional confidence in their fast RT ionic conductivity.

The rightmost column of Table 1 indicates whether the DFT evidence suggests the superionic prediction of the ML model was correct. In total, 8 materials are observed to show Li conduction of 10\(^{-4}\) S/cm or above at RT; 2 materials are near this threshold, and 10 fall far below.

**ii. Discoveries from Random Selection.** The results of the simulation on the randomly selected materials are listed in Table 2 and plotted in Figure 2b. Of the 21 randomly chosen and simulated materials, five structures demonstrated measurable Li conduction at 900 K: LiSO\(_3\)F, LiSbO\(_3\), Li\(_2\)Ho(BO\(_3\))\(_3\), Li\(_2\)Te\(_5\)O\(_9\), and RbLi(H\(_2\)N)\(_2\), with Li diffusivities at 900 K of 0.3, 0.1, 0.05, 0.04, and 0.02 Å\(^2\)/ps, respectively. The remaining 16 structures showed no observable Li conduction on the time scale of tens to hundreds of picoseconds. See Supporting Information for computational parameters and simulation times.

The five materials that exhibited diffusion at 900 K were simulated again at alternate temperatures. After constructing the conductivity extrapolation, the conductivities in three of these materials (LiSO\(_3\)F, Li\(_2\)Ho(BO\(_3\))\(_3\), and LiSbO\(_3\)) were found to extrapolate to above 10\(^{-4}\) S/cm at room temperature. The remaining two, however, showed zero diffusion during simulation at 700 K. Using typical values for the Li concentration and Bader charge, we estimate that zero Li diffusion after 50 ps of simulation at 700 K likely corresponds to an ionic conductivity of 10\(^{-5}\) S/cm or less. Therefore, the RT conductivity of these two materials is predicted to be far below 10\(^{-4}\) S/cm when the assumed 700 K conductivity is factored into the extrapolation.

**iii. Discussion of Discovered Materials.** Between the ML-guided search and the random search, we identify 12 materials total that are predicted by our DFT-MD simulations to exhibit significant Li ion conduction at RT (and the DFT-MD of Zevgolis et al.\(^9\) for Li\(_3\)InCl\(_6\)). Aside from Li\(_3\)InCl\(_6\) and Li\(_2\)Te\(_5\)O\(_9\), none of these materials have reported DFT or experimental conductivity values in the literature to our knowledge. Due to our additional screening steps, these materials also satisfy several other critical criteria beyond fast ionic conductivity to make them useful as solid-state electrolytes in Li-ion batteries: all materials possess DFT-predicted band gaps greater than 1 eV to ensure limited electronic conduction, are free from transition metal elements which may easily reduce in contact with a Li metal anode, exhibit high predicted electrochemical stability against oxidation by cathodes (>4 V vs Li/Li\(^+\)), and sit on the convex hull of their phase diagrams to ensure robust structural (phase) stability.

Two of the materials discovered here (Li\(_2\)B\(_2\)S\(_5\) and Li\(_3\)B\(_2\)S\(_3\)) belong to the Li–B–S system, which appears to be a promising class of materials for realizing fast conducting, electrochemically stable, low mass solid electrolyte materials. The RT ionic conductivity of Li\(_2\)B\(_2\)S\(_5\) in particular is predicted to be an exceptional 74 mS/cm, six times higher than the material Li\(_2\)GeP\(_2\)S\(_12\) (12 mS/cm),\(^{16}\) one of the best known Li-ion conductors. CsLi\(_3\)BS\(_3\) is a Cs-substituted isomorph of Li\(_3\)BS\(_3\), a fast conductor also in the Li–B–S family.\(^{25}\) These three phases were initially identified and characterized as phases observed within Li–B–S (and Cs–Li–B–S) glasses.\(^{26–28}\) The glassy...
Table 1. Density Functional Theory Molecular Dynamics Results for Machine Learning-Guided Material Selection

<table>
<thead>
<tr>
<th>MPID (mp-#####)</th>
<th>chemical formula</th>
<th>computed (\langle D_{ii} \rangle) (Å²/ps), (T) (K)</th>
<th>average Li Bader valence charge computed (\langle \sigma_{ii} \rangle) (mS/cm) (T) (K)</th>
<th>extrapolated RT (\langle \sigma_{ii} \rangle) (mS/cm)</th>
<th>ML-predicted superionic?</th>
<th>evidence for correct model prediction?</th>
</tr>
</thead>
<tbody>
<tr>
<td>554076 BaLiBS₃</td>
<td>~0 [900]</td>
<td>–</td>
<td>~0 [900]</td>
<td>&lt;10⁻⁶</td>
<td>yes</td>
<td>no</td>
</tr>
<tr>
<td>532413 Li₄B₁₃S₃</td>
<td>1.05 (1.1, 0.74)  [900], 0.39 (0.65, 0.42) [700], 0.058 (0.065, 0.052) [400]</td>
<td>+0.89</td>
<td>1900 (2000, 1400) [900], 920 (1500, 990) [700], 240 (270, 210) [400]</td>
<td>74 (71, 64)</td>
<td>yes</td>
<td>no</td>
</tr>
<tr>
<td>569782 Sr₂LiCBr₅N₂</td>
<td>0.15 (0.15, 0.047) [1200], 0.038 (0.039, 0.027) [900]</td>
<td>+0.90</td>
<td>80 (84, 26) [1200], 27 (28, 20) [900]</td>
<td>0.0033 (0.0031, 2.3)</td>
<td>yes</td>
<td>marginal</td>
</tr>
<tr>
<td>558219 SrLi(BS₃)₃</td>
<td>~0 [900]</td>
<td>–</td>
<td>~0 [900]</td>
<td>&lt;10⁻⁶</td>
<td>yes</td>
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<td>yes</td>
<td>no</td>
</tr>
</tbody>
</table>

"We simulate 20 materials as identified by the ML model. For each of the simulated materials, we provide the computed values for diffusivity, conductivity, and average Li Bader charge and the extrapolated RT ionic conductivity values. Here, the diffusivity \(\langle D_{ii} \rangle\) represents the average of the diagonal elements of the diffusivity tensor. In parentheses we provide the diffusivities corresponding to one standard deviation above and below the mean when measuring the diffusivity from varying times. We extrapolate these alternate diffusivities to RT to compute expected deviations in the RT conductivity predictions. The second-to-right column gives the ML-based model prediction for superionic likelihood, and the rightmost column communicates whether the ML-based model gave a correct prediction for each material. Eight materials have an extrapolated RT ionic conductivity of \(10^{-4}\) S/cm or higher, two are below but near this threshold, and ten did not show any Li conduction even at high temperature. \(\text{Li}_{3}\text{InCl}_6\) and \(\text{LiCl}\) were not simulated ("N/S") because conductivity values could be found in the literature. \(\text{Li}_{3}\text{InCl}_6\) was only simulated at 400 K due to melting at higher temperatures. This extrapolation to RT assumes a diffusion barrier of 0.35 eV, a typical diffusion barrier in fast conducting systems. \(\text{CsLi}_2\text{BS}_3\) could not be simulated at lower temperatures and thus the extrapolated ionic conductivity at RT was taken to be of the same order of magnitude as that of \(\text{Li}_2\text{BS}_3\)."
We simulate 21 materials as chosen at random and provide the computed values for diffusivity $\langle D_\sigma \rangle$ (average of diagonal elements in the diffusivity tensor), conductivity, and average Li Bader charge and the extrapolated RT ionic conductivity values. In parentheses, we provide the deviations from the mean when measuring the di ionic conductivity.

\[
\begin{array}{cccccccc}
\text{MPID (mp-##)} & \text{chemical formula} & \text{computed } \langle D_\sigma \rangle (\text{Å}^2/\text{ps}) & T (K) & \text{computed } \langle \sigma_{ii} \rangle (\text{mS/cm}) & T (K) & \text{extrapolated RT } \langle \sigma_{ii} \rangle (\text{mS/cm}) & \text{ML-predicted superionic?} & \text{evidence for correct model prediction?} \\
6674 & \text{Li}_{14}\text{Sr}_{4}\text{SbO}_{6} & \sim [900] & \sim [900] & \sim [900] & [900] & [900] & \text{no} & \text{yes} \\
8609 & \text{Li}_{4}\text{UO}_{3} & \sim [900] & \sim [900] & \sim [900] & [900] & [900] & \text{no} & \text{yes} \\
28567 & \text{Li}_{4}\text{BiF}_{6} & \sim [900] & [900] & \sim [900] & [900] & [900] & \text{no} & \text{yes} \\
12160 & \text{Li}_{5}\text{H}_{5}\text{(BO}_{3})_{3} & 0.10 (0.12, 0.10) [1200], 0.048 (0.05, 0.044) [900] & [1200], 300 & [900] & [1200], 300 & [900] & \text{no} & \text{no} \\
6787 & \text{Rb}_{2}\text{Li},\text{O}_{2} & \sim [900] & \sim [900] & [900] & [900] & [900] & \text{no} & \text{yes} \\
560072 & \text{Li}_{4}\text{Be}_{5}\text{As}_{2}\text{ClO}_{12} & \sim [900] & \sim [900] & \sim [900] & [900] & [900] & \text{no} & \text{yes} \\
7941 & \text{Li}_{2}\text{TeO}_{3} & \sim [900] & \sim [900] & \sim [900] & [900] & [900] & \text{no} & \text{yes} \\
13772 & \text{Li}_{2}\text{Pr}_{2}(\text{BO}_{3})_{3} & \sim [900] & \sim [900] & \sim [900] & [900] & [900] & \text{no} & \text{yes} \\
8452 & \text{NaLiS} & \sim [900] & \sim [900] & \sim [900] & [900] & [900] & \text{no} & \text{yes} \\
770932 & \text{Li}_{2}\text{SbO}_{3} & 0.18 (0.19, 0.16) [1200], 0.065 (0.16, 0.078) [900] & [1200], 210 & [900] & [520, 240] & [900] & 0.67 (520, 5.3) & \text{no} & \text{no} \\
12829 & \text{Li}_{2}\text{CaGaF}_{4} & \sim [900] & \sim [900] & \sim [900] & [900] & [900] & \text{no} & \text{yes} \\
27811 & \text{Li}_{2}\text{TeO}_{3} & 0.050 (0.14, 0.065) [1200], 0.044 (0.057, 0.042) [900], \sim [700] & [1200], 130 & [900], \sim [700] & [900], \sim [700] & \text{no} & \text{no} \\
8180 & \text{LiNO}_{3} & \sim [900] & \sim [900] & \sim [900] & [900] & [900] & \text{no} & \text{yes} \\
7971 & \text{Ba}_{2}\text{Li}_{2}\text{SbO}_{4} & \sim [900] & \sim [900] & \sim [900] & [900] & [900] & \text{no} & \text{yes} \\
8449 & \text{Rb}_{2}\text{Li}_{2}\text{SiO}_{4} & \sim [900] & \sim [900] & \sim [900] & [900] & [900] & \text{no} & \text{yes} \\
558045 & \text{NaLi}_{2}\text{PO}_{4} & \sim [900] & \sim [900] & \sim [900] & [900] & [900] & \text{no} & \text{yes} \\
562394 & \text{Cs}_{4}\text{Li}_{4}(\text{SiO}_{4})_{3} & \sim [900] & \sim [900] & \sim [900] & [900] & [900] & \text{no} & \text{yes} \\
510073 & \text{RbLi}_{2}(\text{H}_{2} \text{N})_{2} & 0.088 (0.12, 0.03) [1100], 0.04 (0.036, 0) [900], \sim [700] & [1100], 84 (76, 0) [900], \sim [700] & [1100], 84 (76, 0) [900], \sim [700] & [1100], 84 (76, 0) [900], \sim [700] & \text{no} & \text{no} \\
7744 & \text{Li}_{2}\text{SO}_{4} & 0.06 (0.14, 0.065) [1200], 0.20 (0.02, 0.17) [900], 0.072 (0.08, 0.059) [700] & [1200], 130 & [900], \sim [700] & [900], \sim [700] & \text{no} & \text{no} \\
352 & \text{Cs}_{4}\text{Li}_{2}\text{TeF}_{6} & \sim [900] & \sim [900] & \sim [900] & [900] & [900] & \text{no} & \text{yes} \\
8892 & \text{Li}_{2}\text{F}_{6} & \sim [900] & \sim [900] & \sim [900] & [900] & [900] & \text{no} & \text{yes} \\
\end{array}
\]

We simulate 21 materials as chosen at random and provide the computed values for diffusivity $\langle D_\sigma \rangle$ (average of diagonal elements in the diffusivity tensor), conductivity, and average Li Bader charge and the extrapolated RT ionic conductivity values. In parentheses, we provide the deviations from the mean when measuring the di ionic conductivity.

The second-to-right column gives the ML-predicted superionic conductivity, and the rightmost column communicates whether the ML-based model gave a correct prediction for each material. The Bader charge on the Li atoms in the control group is taken to be +1.0 for computational efficiency. Additional randomly chosen materials that melted at 900 K or whose simulations did not converge were discarded and are not listed here. The success rate of random selection is computed to be $3/21 = 14.3\%$.

Li–B–S–I system was reported to be an exceptional Li ion conductor in 1981, but to our knowledge these phases within the crystalline Li–B–S space have not been studied as solid electrolytes.

Li$_3$ErCl$_6$, an Er-substituted analogue to the fast-conducting Li$_3$InCl$_6$, appears to be a promising Li conductor with a predicted RT ionic conductivity of 3 × 10$^{-4}$ S/cm, although the high atomic mass and low earth abundance of Er makes it a less attractive candidate for battery applications. Li$_3$ErCl$_6$ was first studied and characterized in 1996 as a potential fast Li ion conductor, but to our knowledge the ionic conductivity was not reported. We find another Er-based compound appears to be a marginal RT Li ion conductor, LiErSe$_2$. LiErSe$_2$ is a Li-intercalated two-dimensional layered material that was initially studied in 1987 for its phase transitions and magnetic properties.

We discover three nitride-based fast ion conductor materials with quite complex compositions: Sr$_2$LiCBr$_3$N$_2$, RbLi(H$_2$N)$_2$, and LiMgB$_4$(H$_3$N)$_2$. Sr$_2$LiCBr$_3$N$_2$ is a solid-state carbodiimide material that was first characterized in 2005. RbLi(H$_2$N)$_2$ was originally synthesized in 1918 and subsequently characterized in 2002. LiMgB$_4$(H$_3$N)$_2$ was initially synthesized and studied in 2014 as a candidate solid-state hydrogen storage material. The exotic stoichiometries of these materials underscores the utility of the ML-based model, as it seems unlikely that such complex materials would be discovered by chance to be fast ion conductors. A potential drawback in using these materials with many elements as solid electrolytes is that the electrochemical stability is likely to be low, given the high number of stable interfacial phases that could form from a subset of these elements.
incorrectly predicted to be a good ion conductor by DFT-MD. It is also possible that the bulk ionic conductivity of the material is fast, which is accurately modeled by DFT-MD, but it becomes a poor ion conductor under the experimental conditions of ref 40, e.g., due to defects, stoichiometric variations, or microstructural effects. The calculations performed here suggest that this material warrants a revisit of the experimental results.

iv. Model Performance. Quantifying the accuracy of the original ML-based model should be done based on its performance on a randomly sampled test set, not on a test set consisting only of positive predictions. The 21 randomly chosen materials that were simulated here provide such a set. On this set, the model correctly predicts the DFT-validated material label 19/21 times, yielding an overall predictive accuracy of 90.5%. This value is similar to the 90.0% model accuracy predicted via leave-one-out cross-validation in the original model building process.7 Only one material (LiSO3F) was predicted to be a superionic conductor by the ML-based model, and our DFT simulation confirmed the prediction was correct; this gives a model precision of 1.0 on this particular test set. Of the three materials that DFT-MD found to be RT superionic conductors (LiSO3F, LiShO3, LiH6O3(BO3)3), only one was accurately predicted by the ML model; this gives a model recall of 1/3 = 0.333 on the test set. Taken together, these values give the model an F1 score of 0.50. For comparison, the baseline F1 score of fast ion conductors on the test set (i.e., the F1 score of completely random guesswork) is simply the fraction of positive examples in the test set, which gives a baseline F1 score of 0.143. Therefore, the F1 score of the ML model is 0.50/0.143 = 3.5 times higher than the F1 score associated with completely random trial-and-error.

As an alternative means of quantifying the expected improvement in experimental outcomes of Li conductivity measurements in the ML-guided versus random searches, we average the predicted RT ionic conductivities across all randomly chosen materials and all ML-chosen materials. Given that ionic conductivity varies over many orders of magnitude, we average the base-10 logarithm of conductivity. This gives a log-average RT ionic conductivity of the randomly chosen materials of 6.8 × 10−9 mS/cm, while the log-average conductivity in the ML chosen materials is 3.0 × 10−4 mS/cm, 44 times higher. This comparison assumes the average RT ionic conductivity in the materials with no observed Li motion is 10−8 mS/cm; as discussed in Supporting Information Section S1, this value likely serves as an upper bound in most cases on the conductivity in these materials, and thus the 44× improvement in conductivity is a lower bound on the true improvement the model offers, e.g., if the average conductivity of the nonconductors is taken to be 10−9 mS/cm, the improvement in conductivity increases to over 350×.

As an additional performance metric, we compare the likelihood of discovering a RT superionic conductor under the ML-guided screening versus random searching. The random search yields 3 fast conductors and 18 nonconductors. This yields a baseline superionic probability of 3/21 = 14.3%. To make a fair comparison to the ML-guided case where materials that melted at 900 K were not discarded, we do not consider in the statistics those ML-chosen materials that melted at 900 K. Two of the eight ML-chosen RT superionic conductors melted at 900 K (LiMgB3(H9N)2 and Li2HIO). Removing these two materials from the count, the ML model identifies six fast conductors, two marginal conductors, and ten nonconductors. Counting the marginal cases as one-half, this gives an ML-guided conductor discovery rate of 7/18 = 38.9%. Therefore, applying the ML-based model to screen candidate materials before experimentation yields an expected 0.389/0.143 = 2.7× improvement in discovery rate. In Figure 3 we provide a histogram showing the ionic conductivity distribution in the ML-chosen and randomly chosen materials; the ML model discovers over five times more materials with ionic conductivity above 1 mS/cm, while the number of nondiffusive materials falls from 86% to 56%. We refer the reader to the Supporting Information Section S2 for further discussion.

v. Comparison to Human Intuition. We perform a test of the model's speed and predictive power against human intuition, which is likely to identify fast conducting materials at a higher rate than the background distribution of ion conductors. Several design principles for predicting superionic diffusion have been reported recently,41–43 but human-guided search efforts may or may not take these principles into account. To understand the precision of human intuition, we polled six Ph.D. students in the Department of Materials Science & Engineering at Stanford University to test their ability to predict superionic conduction in the 21 randomly chosen materials. The students are all actively engaged in research involving ion conductors. The students were allowed to take as much time as necessary to make predictions and could access all structural information about the candidate materials provided on the Materials Project database or any other source. This included structure (CIF) files, space and point groups, and properties like formation energy and band gap. This poll sought to test how quickly and accurately the
students would naturally predict superionic conduction in the absence of eq 1, and so they were not given access to any of the pretabulated features employed in eq 1, nor were they encouraged to make predictions quickly. After making predictions, we calculated their average precision and recall on the data set; the average precision was 0.25 and the average recall was 0.222, giving an overall average F1 score of 0.235. The baseline F1 score for random guessing is 0.143. The students took approximately one minute to make each individual prediction. For comparison, the ML model made each prediction in approximately one millisecond.

In Figure 4 we compare the performance in speed and F1 score of the Ph.D. students and the ML model. The ML model exhibits more than double the F1 score of the students and is more than 3 orders of magnitude faster. As a reference, we provide the performance of DFT-MD, which we assume has an F1 score of 1.0 and requires approximately 2 weeks to make a reliable prediction.

While DFT-MD is taken here to be ground truth, the logistic regression model utilized in this work is trained on experimental measurements, where grain boundaries, contaminants, and other uncharacterized factors may have altered the result. This may mean the model is best suited to guide experimental searches, as the model has a built-in bias toward expected results under experimental conditions. In contrast, our DFT-MD simulates conductivity in single crystal bulk systems with small Li vacancy concentrations. Although DFT-MD is well-suited to make predictions under these conditions, its predictions have potential to deviate from the ground truth for the ML model; for example, the conflicting DFT-MD and experimental reports for RT Li ion conductivity in Li₆Ho(BO₃)₃ between this work and ref. 40. This work predicts RT ion conductivity of approximately 5.1 mS/cm, while ref 40 reports RT ion conductivity orders of magnitude lower (see Section III, Subsection iii). It is possible that the model may recognize that experimental efforts on Li₆Ho(BO₃)₃, are likely to yield a poor conductor even though the single crystal conductivity is fast. We look forward to future experimental reports of the ionic conductivities of some of the materials presented here, to both advance the state of the art in ion conductors and to further quantify the performance of the data driven approach.

IV. CONCLUSIONS

Guided by machine learning methods, we discover many new solid materials with predicted superionic Li ion conduction ($>10^{-4}$ S/cm) at room temperature: Li₅B₇S₁₃, Li₃B₂S₅, Li₃ErCl₆, Li₅SO₃F, Li₃InCl₆, Li₃HIO, LiMgB₃(H₉N)₂, and CsLi₃BS₅. Two additional materials show marginal RT conductivity: Sr₂LiCBr₃N₂ and LiErSe₂. One of these materials, Li₅B₇S₁₃, has a DFT-MD predicted RT Li conductivity (74 mS/cm) many times larger than the fastest known Li ion conductors. A search over randomly chosen materials identifies two additional materials with promising predicted RT ionic conductivities: Li₆Ho(BO₃)₃ and LiSbO₃. In addition to high ionic conductivities, all these materials have high band gaps, high thermodynamic stability, and no transition metals, making them promising candidates for solid-state electrolytes in SSLIBs. These materials represent many exciting new candidates for solid electrolytes in SSLIBs, and we encourage 

![Figure 4. Comparison of human and machine. Here we plot the predictive accuracy (F1 score) of the ML model and the predictive accuracy of a group of six Ph.D. students working in materials science and electrochemistry. We chose 22 materials at random from a list of 317 highly stable Li containing compounds in the Materials Project database and queried both Ph.D. students and the ML model to predict which materials are superionic Li conductors at room temperature. Validation of superionic conduction is performed with density functional theory molecular dynamics simulation. The Ph.D. students average an F1 score of 0.24 for superionic prediction while the model produces an F1 score of 0.5. For reference, we provide the baseline F1 score expected for random guessing (0.14) as a dashed vertical line. Additionally, the students average a rate of approximately one minute per prediction while the trained ML model makes over 100 predictions per second, a thousand-fold increase in speed.](image)

Table 3. Summary of Model Performance

<table>
<thead>
<tr>
<th>model</th>
<th>precision</th>
<th>recall</th>
<th>F1 score</th>
<th>fast Li ion conducting materials identified (in this work)</th>
</tr>
</thead>
<tbody>
<tr>
<td>machine learning (logistic regression on 40 examples)</td>
<td>1.0</td>
<td>0.33</td>
<td>0.5</td>
<td>Li₅B₇S₁₃, Li₃B₂S₅, Li₃ErCl₆, Li₃HIO, Li₅SO₃F, Li₃InCl₆, CsLi₃BS₅, LiMgB₃(H₉N)₂, Sr₂LiCBr₃N₂ (marginal), LiErSe₂ (marginal)</td>
</tr>
<tr>
<td>random selection</td>
<td>0.14</td>
<td>0.14</td>
<td>0.14</td>
<td>Li₆Ho(BO₃)₃, LiSbO₃, Li₅SO₃F</td>
</tr>
<tr>
<td>Ph.D. student screening</td>
<td>0.25</td>
<td>0.22</td>
<td>0.24</td>
<td>-</td>
</tr>
</tbody>
</table>

*Here we provide the performance metrics (precision, recall, F1 score) and list of discovered fast Li ion conducting materials for the three models explored in this work. The results of the Ph.D. student screening represent the average performance of six students on the same test set of randomly chosen materials as used to compute the “random selection” statistics. The machine learning-based approach to predicting ion conductivity from a small data set of 40 materials significantly outperforms both random chance and the average polled Ph.D. student. This highlights the promise of applying machine learning-based approaches to materials screening before performing computationally expensive simulation techniques like DFT-MD or time consuming experimental tests.*

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subsequent experimental investigations into the properties of these materials.

Compared to the machine learning-guided search, the control experiment of searching for fast ion conductors through random guesses yields significantly lower quality results. Screening with our ML-based model improves the likelihood of superionic discovery by nearly three times, with a more than 44X improvement in log-average of RT ionic conductivity. Additionally, the model outperforms the intuition of Ph.D. students actively working on ion conductors by more than a factor of 2. This is a significant acceleration beyond trial-and-error research and provides confidence in our ML-based approach to materials screening. These results are summarized in Table 3. Such data-driven models are expected to improve with every new data point, and with the data we report here we expect to drive significant improvements to ML-based models for Li conduction.

The improvement over random guessing provided by our ML-based model for predicting Li ion conductivity underscores the importance of thorough data reporting, centralized data collection, and careful data analysis for materials. Although significant improvements have been made in the last several years thanks to incentives provided by the Materials Genome Initiative, there are still many materials properties, especially experimentally expensive properties like ionic conductivity, without any comprehensive data repository. This work demonstrates that learning on even small sets of materials data (40 samples) can offer a significant advantage in screening efforts. To that end, we encourage efforts to continue centralizing and learning on diverse types of data on materials.

# ASSOCIATED CONTENT

## Supporting Information
The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/acs.chemmater.8b03272.

Table of the computational parameters for the DFT simulations; calculations for the ionic conductivity upper bound in the nondiffusive materials; and discussion on the impact on the model performance metrics of the removal of the randomly chosen materials that melted in simulation at 900 K (PDF)

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## Notes
The authors declare no competing financial interest.

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