Enhancement of ion diffusion by targeted phonon excitation

Gordiz et al. computationally show that, by exciting the top contributing phonons to the ion hop, diffusivity increases without raising the bulk temperature. This observation suggests that diffusion, or more generally any transition, can be enhanced by exciting only the contributing phonons to that transition to elevated temperatures, instead of heating up the entire material.
Enhancement of ion diffusion by targeted phonon excitation

Kiarash Gordiz, 1,* Sokseiha Muy, 2 Wolfgang G. Zeier, 3 Yang Shao-Horn, 1,2,4 and Asegun Henry1,5,*

SUMMARY
Ion diffusion is important in a variety of applications, yet fundamental understanding of the interaction of phonons and the mobile species in solids is still missing. In this work, we introduce two formalisms that determine the individual contributions of phonons to the diffusion of ions through a solid, based on nudged elastic-band calculations and molecular dynamics simulations. The results for a model ion conductor of Ge-substituted Li3PO4 (Li3.042Ge0.042P0.958O4) reveal that more than 87% of the Li+ ion diffusion originates from less than 10% of the vibrational modes between 8 and 20 THz. By deliberately exciting a small, targeted subset of these contributing modes (<1%) to a higher temperature and still keeping the lattice at a low temperature, we observe an increase in diffusivity by several orders of magnitude. This new understanding identifies new avenues for increasing diffusivity by engineering the vibrations in a material without necessarily changing the compound chemistry.

INTRODUCTION
Solid state mass and ion diffusion is central to many applications, ranging from batteries1 and fuel cells2 to sensors3 and filters.4 In many of these applications, the performance is limited by the slow diffusion of mass or ions. Although some families of solid-ion conductors, such as silver iodide and Li10GeP2S12 (LGPS) exhibit high ion conductivities of ~1 S/cm5 and ~0.01 S/cm6,7 at room temperature, respectively, achieving high ion conductivities for divalent ions, including oxides, is challenging. For the conductivity of oxygen ions exhibited by yttria-stabilized zirconia (YSZ)8,9 to reach 0.01 S/cm, an elevated temperature of 1,100 K is needed. Higher oxygen diffusivity at a lower operating temperature could enable a lower system cost, longer life, and greater proliferation.10 Great advances have been made to increase the ion transport by cation and anion substitution to alter the charge carrier density and the diffusion pathways to facilitate low-energy jumps. The classical approaches of chemical modification, such as cation and anion substitution, have led to tunable conductivity over several orders of magnitude in several families of super-ionic conductors, such as the lithium super-ionic conductor (LISICON)7 and the sodium super-ionic conductor (NASICON).11 Thus, regardless of the mobile ion species, understanding the key factors that dictate ion conductivity is needed to devise new strategies to further increase the conductivity beyond tuning of chemical compositions.

One possible avenue to improve diffusivity without modifying the chemistry stems from considering the local ion movement in the structure. At a given temperature, ions thermally vibrate in their specific crystallographic lattice sites until sufficient thermal energy is available for an ion jump. The coupled ion vibration and ion movement that is paramount for ion diffusion leads to the idea that lattice vibrations...
(phonons) may have a role in the ion jump. Such an idea can be further intuited by considering the role of temperature as a spectrum of scalar contributions, not as a single scalar value. Based on this more-detailed view, in solids and rigid molecules, temperature is composed of a summation of individual contributions by modes of vibration, which means that a group of vibrating atoms can be described as a summation of collective vibrations, each with a specific frequency, often termed eigen modes, normal modes, or phonons. According to the traditional description for solid-state diffusion based on the Arrhenius relation $D \propto \exp(-E_a / k_B T)$, diffusion increases exponentially with increasing temperature, where $k_B$ and $T$ denote the Boltzmann’s constant and the bulk temperature, respectively. The term “bulk” temperature, here, denotes the temperature that would be sensed if the material were in thermal equilibrium. Although temperature is generally understood as a proxy for the level of excitation of the phonons or normal modes of vibration within a solid material, it should be noted that each mode has its own time-varying amplitude and individual temperature. It is possible for a material to experience some non-equilibrium in the individual mode temperatures, whereby certain modes are highly excited to an effective temperature above all others (e.g., joule heating of optical modes in a transistor). In such a situation, a subset of modes can behave as though they are at a higher effective temperature than the bulk temperature.

In the classical limit, for an individual mode labeled $n$, the modal temperature $T_n$ is obtained from $k_B T_n = Q_n^2 \omega_n^2$, where $Q_n$ and $\omega_n$ are the modal displacement coordinates (which can be interpreted as the mode amplitude) and frequency of vibration of mode $n$ (see Note S1 for a derivation of this relation and its quantum analog).

The idea of modal temperatures have also been introduced in previous theoretical studies, in which, for the first time, the two-temperature model was extended to study the weak coupling between certain groups of phonons and its effect on thermal transport in solid materials. Our focus in this study is the distinction between the bulk and modal temperatures and how that influences solid-state ion transport. As the bulk temperature increases, the amplitudes of all the modes in the system increase, which facilitates ions to hop over activation barriers, thereby enhancing diffusion. There are practical ways, however, to increase the amplitude and temperature of a small subset of normal modes by exciting them with light either directly through photon-phonon coupling or indirectly through electron-photon coupling or THz electric fields and keeping them in non-equilibrium with respect to the rest of the modes, without increasing the bulk temperature. This idea of selective vibrational mode excitation has been used in previous studies, as a novel tool, to modify the electronic properties of the structure through electron-phonon coupling for plasmonics, photovoltaics, and charge-transport applications. Selective modal excitations have also been used to induce vibrational and structural changes in the system for phase-change applications. Inspired by recent experimental observation of solid-state ion-diffusion induced by external THz-electric field illumination, in this study, we aim to examine whether only a small subset of modes in the lattice are responsible for the hopping of ions, which would allow the increase of the diffusivity by exciting those subset of modes without increasing the bulk temperature. Experimental realization of such an idea would have far-reaching implications for chemical reactions, phase transitions, or other related phenomena because it would suggest that migration along a reaction/transition coordinate could be accelerated by exciting only the most important normal modes, rather than all the modes via an increase in the bulk temperature. This approach would mean that a material and its container could remain “cold,” but it could exhibit kinetics associated with a much higher temperature, by only making a subset of the most important modes “hot.” To assess that opportunity, we use the example of ion diffusion, in which we
(1) identify the mode-level contributions to ion diffusion to determine how many modes are responsible for an ion hop in the system, and (2) quantify to what extent diffusivity can be increased by targeted excitation of a small subset of these contributing modes without increasing the bulk temperature.

The role of specific structural modes and phonons in facilitating solid-state diffusion has been reported in other studies.\textsuperscript{30–41} The frequency of longitudinal acoustic phonons was shown to exhibit a strong correlation with the enthalpy of self-diffusion in body-centered cubic metals.\textsuperscript{33} In addition, octahedral rotation has been shown to promote fast oxygen-ion conduction in perovskite-related phases.\textsuperscript{30,31,35,39,40} Li and Benedek\textsuperscript{37} calculated the force constants associated with octahedral rotations in different Ruddlesden-Popper phases and showed that the willingness of a material to go under these octahedral rotations correlates with the migration barrier of oxygen ions. Similarly, anion rotation in closo-borate solid electrolytes has been shown to be important for ion transfer through the lattice.\textsuperscript{32,36,41} Recently, Muy et al.\textsuperscript{38} showed that lowering the computed average vibrational frequency of the lithium sublattice in LISICON by cation and anion substitution is accompanied by a reduced migration barrier. Similar correlations between activation energy and computed average frequency of sodium ions have been noted for sodium ion conductors,\textsuperscript{34} in which activation energy is correlated with the Debye temperature, a proxy for the softness of the lattice, i.e., corresponding to low-energy vibrational modes. However, the average frequency of vibration reflects a weighted sum over all phonon modes, and the Debye temperature only measures the slopes of the acoustic phonon modes at very low wave vectors, neglecting the influence of the optical phonons and including the effect of some irrelevant phonons to ion migration. Most of these insights have been obtained from density-functional theory (DFT) lattice relaxation, \textit{ab initio} molecular dynamics (AIMD), or experimental characterization. However, no direct information about specific phonons and their respective eigenvectors have been employed in a systematic way to obtain a deeper insight into these phenomena. The information for eigenvectors of phonons have been used in several modal-analysis methodologies for thermal transport, including atomistic Green's function (AGF),\textsuperscript{42} Boltzmann transport equation (BTE),\textsuperscript{43} and molecular dynamics (MD) simulations.\textsuperscript{44–47} These methods have shown that, in different materials, certain groups of modes could contribute to heat transfer with higher rates, for instance, the high-frequency, in-plane phonons in graphene\textsuperscript{48} and interfacial modes in semiconductor interfaces.\textsuperscript{49–51} In this study, we aimed to identify modes of vibration that facilitate the ion migration by directly using the knowledge of phonons and their respective eigenvectors, based on the approaches that were recently developed in the context of MD-based phonon analysis.\textsuperscript{44–47}

Here, we chose Ge-substituted Li\textsubscript{3}PO\textsubscript{4} as our model system because Li\textsubscript{3}PO\textsubscript{4} is a well-known parent structure; from which, numerous fast, solid conductors have been developed, including LGPS,\textsuperscript{52,53} and its ion conductivity can be tuned up to \(\sim 10\) orders of magnitude by aliovalent and anion substitution.\textsuperscript{6} To identify the modal contributions to the diffusion of the lithium ion, which is the only diffusing species in the structure, we introduce two methodologies, one based on nudged elastic band (NEB) calculations and another based on MD simulations. Following the previously developed approaches for modal decomposition of thermal transport,\textsuperscript{44–47} we project the atomic displacement or velocity fields onto the normal modes of vibration along the hopping trajectory obtained from a NEB calculation. The magnitude of the projection determines which phonons/normal modes become excited to facilitate that specific ion hop. Our results suggest that only a small subset of the phonons is responsible for ion diffusion in Ge-substituted Li\textsubscript{3}PO\textsubscript{4} and that diffusivity can be
increased by increasing only the temperature of those modes instead of the bulk temperature.

RESULTS AND DISCUSSION

Modal contributions obtained from NEB calculations of Ge-substituted Li3PO4

The modal contributions to one example hop in Ge-substituted Li3PO4 shows that a small subset of modes (around 10 THz) contributed much greater than the rest of the modes (Figure 1A). The dominant hopping mechanism in the Ge-substituted Li3PO4 structure, is the concerted hop54 of two Li ions, one of which is the interstitial, as shown in Figure 2C, and it is compared with two other hopping mechanisms in Figures 2A and 2B. The concerted hop mechanism is known as the interstitialcy mechanism,55,56 which has been reported57,58 as the dominant mechanism for lithium ion diffusion in Li3PO4 and is used as the migration mechanism for the NEB-based modal analyses presented in this study. The normalized integrations of the contributions with respect to frequency (the accumulation) are shown as a continuous curve in Figure 2B for all the 619 hops in the structure (the process of identifying those hops is explained in the Experimental procedures section). Most of the contributions to all the hops in the system come from modes between 8 and 20 THz (Figure 1B). Although that range is centered around the attempt frequency of the lithium ion (≈14 THz; see Figure S1 and Note S2 for the calculation procedure), the large range of frequencies for the contributing phonons shows that more-intricate phonon processes are operative, which differs from the simpler picture described by traditional approaches,59 wherein only one value of vibrational frequency (the attempt frequency) enters the formulation. By averaging over all the modal contributions calculated for all the hops in the system (Figures S2), we confirmed that, on average, >87% of the lattice energy during the ion hop in the structure came from <10% of the modes in the system (Figure S3). In addition, as is explained and shown in Figure S4, in each hopping event, there is at least one mode that has a contribution more than 10 times the average contribution by all the modes. Thus, the results in Figure 1 show that in Ge-substituted Li3PO4, only a few modes are responsible for the interstitialcy diffusion mechanism in Figure 2C.

Because of the larger vibration amplitudes that localized modes attribute to the interstitial and other neighboring ions than that of the delocalized modes60
localized modes might be the most strongly contributing modes to each ion hop. However, inspection of the eigenvectors for the top two contributing modes in Figure 1A (at 9.64 THz and 10.94 THz) revealed that highly contributing modes can be either delocalized (Figure 3A) or localized (Figure 3B) in nature. Because the modal analysis formalism is based on the supercell lattice dynamics (SCLD) calculations (see Experimental procedures), all types of vibrational modes in the system with no inherent assumptions about their character were identified; from which, the contributions to ion diffusion, even for localized modes, were computed. Having access to such a formalism (see Experimental procedures) is crucial because, by breaking the symmetry in the system (e.g., by inclusion of vacancies and interstitial and substitutional ions, which is common in highly conductive solid electrolytes\textsuperscript{61}), the conventional picture of propagating phonons in pure crystalline solids breaks down as other non-propagating and localized modes must be included.\textsuperscript{50,62,63} Figure 5 shows the spectral distribution of all the localized modes of vibration for all the equilibrium configurations in our Ge-substituted Li\textsubscript{3}PO\textsubscript{4} structure.

Diffusivity enhancement by excitation of the highly contributing modes

Figure 4 shows that increasing the energy of the top five contributing modes among all the possible hops belonging to the equilibrium configuration present at each time step during an MD simulation—by increasing their temperature to 500 K and 700 K—increases the diffusivity by 2 ($\sim 10^{-10}$ cm$^2$/s) and 4 ($\sim 10^{-8}$ cm$^2$/s) orders of magnitude respectively, compared with its value ($\sim 10^{-12}$ cm$^2$/s) in the unperturbed (natural) simulation at a lower temperature of 400 K. In fact, when the lattice is kept at 400 K and only the top five most-contributing modes are excited to a particular temperature, the diffusivity becomes what it would have been if the entire material had been heated to that temperature. Unexpectedly, even if we excite only the single highest-contributing mode in each of the equilibrium configurations, almost the same increase in the diffusivity is observed. The small difference between the increase in diffusivity by exciting the five highest-contributing modes and the one by exciting only the highest-contributing mode can be explained by noting that more kinetic energy is input to the hopping ion by exciting five modes. How these excited modes interact with the rest of the unexcited modes in the system is an interesting topic for a future study, which can be investigated using the two-temperature model that was presented in a recent study.\textsuperscript{15}
To illustrate that the increase in diffusivity from excitation only happens when the specific modes that matter most are excited, Figure 4 shows that exciting five random modes in the 8–20 THz frequency range did not lead to significant enhancement in the diffusivity. The novelty of discovering that only a small subset of vibrational modes is responsible for the orders-of-magnitude increase in diffusivity by modal excitation can be understood by noting the theoretical definition of the diffusion coefficient in solid materials,

\[ D = D_0 \exp \left( -\frac{E_A}{k_B T} \right) \]

In that definition, \( E_A \) and \( D_0 \) are known as the activation energy and the prefactor. According to the transition state theory, \( E_A \) and the passing of an ion through the transition state are solely influenced by one mode of vibration in the system. However, \( D_0 \) is theoretically influenced by many modes of vibration in the system, which can be understood by noting its dependence on several other parameters,

\[ D_0 \propto a^2 n v \exp \left( -\frac{\Delta S_m}{k_B} \right) \]

where \( a \), \( n \), \( v \), and \( \Delta S_m \) are the lattice constant, the number of the nearest hopping sites, the average frequency of vibration for the hopping atom, and the entropy of migration, respectively. Among these parameters, \( \Delta S_m \) is a function of all the frequencies of vibration in the system, and because of its exponential influence, it can affect the diffusion coefficient by orders of magnitude, to a degree comparable to the influence of the migration barrier, as was shown in recent studies by Krauskopf et al. and Muy et al. Therefore, because, theoretically, many modes of vibration influence \( \Delta S_m \) and hence, \( D_0 \), the observation that by exciting only a few modes in the system, the diffusivity can change by several orders of magnitude, is, indeed, unexpected.

To check whether traditional parameters for diffusivity can explain the enhanced diffusion rates caused by modal excitation in Figure 4, we quantified the changes in attempt frequency (Figure S1), jump rate (Table S1), radial distribution function (RDF) (Figure S7), amplitude of vibration (Figure S8), Haven ratio (the ratio of the tracer diffusion coefficient to the conductivity [total] diffusion coefficient, which can be interpreted as an index for the strength of correlated hops in the system) (Table S2), and the migration barrier (Figure S9). However, the very small changes in those parameters were not able to explain the orders-of-magnitude increase in diffusivity observed by modal excitation (Figure 4). This also highlights the novelty of the mechanism discovered herein, illustrating the importance of modal temperatures.
and the concept of directionality brought into the analysis by including the eigenvectors of vibration in our phonon-based methodologies (see Experimental procedures). Overall, the 2–4 orders-of-magnitude increase in diffusivity by modal excitation (Figure 4) can be attributed to increasing the energy of the highly contributing modes in the structure, and those modes can push the ion forward along its migration pathway. These results not only show that the identified modes of vibration using our proposed methodologies are correct and are indeed responsible for the ion diffusivity but also offer a new strategy to increase the diffusivity of known solid-state ionic conductors by using techniques such as direct\textsuperscript{16,17,19} and indirect\textsuperscript{20–22} coupling to phonons without increasing the temperature of the lattice.

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**Figure 4. Effect of targeted phonon excitation on the ion diffusion in the Ge-substituted Li\textsubscript{3}PO\textsubscript{4} structure**

By exciting the five highest-contributing modes or even the single highest-contributing mode to ion diffusion in the Ge-substituted Li\textsubscript{3}PO\textsubscript{4} structure to 500 K or 700 K and keeping the lattice at 400 K, the diffusivity increases by around two and four orders of magnitude compared with the unperturbed/natural simulation at 400 K. Exciting five random modes to 700 K, however, does not lead to a noticeable increase in diffusivity. The reported values are tracer diffusivity, but the same increase was also observed in the conductivity (total) diffusivity\textsuperscript{47} (see Table S2). The activation energy obtained from the linear fitting of the Arrhenius equation\textsuperscript{97} (0.68 eV) is in reasonable agreement with the one obtained from the NEB calculations for the interstitialcy hop (Figure 2). In addition, the orders of magnitude of our obtained diffusivity values are in agreement with the ones reported for similar structures in previous studies.\textsuperscript{84,65} Error bars represent the standard deviation for each data point. For some data points, error bars cannot be seen because they are smaller than the marker size.

**Modal contributions obtained from MD simulations of Ge-substituted Li\textsubscript{3}PO\textsubscript{4}**

The modal contributions to the diffusivity for the Ge-substituted Li\textsubscript{3}PO\textsubscript{4} structure, calculated from the mass diffusivity modal analysis (MDMA) method (see experimental procedures), are shown in Figure 5B. The increase in the diffusivity accumulation function in Figure 5B shows that the modes with frequencies between \(\sim 8\) and \(20\) THz are responsible for ion diffusivity. This observation is in excellent agreement with the modal contributions obtained from the NEB-based approach (Figure 1B), which is also averaged and shown in Figure 5B. The accumulation of the partial
The MDMA formalism is able to provide the degrees of both harmonic and anharmonic contributions to the ion diffusion in the lattice.

(A) Partial phonon DOS for different elemental components of the structure in the form of a stacked area chart.

(B) Modal contributions in the form of an accumulation function to the conductivity diffusivity obtained from the MDMA method (red curve) and to the ion hop from the NEB-based modal analysis (black curve). In addition, the accumulation function for the lithium partial DOS (blue curve) has also been shown.

(C) Two-dimensional (2D) map showing the magnitudes of the pairwise correlations/interactions contributing to the conductivity diffusivity on the plane of two frequency axes.

(D) Similar to (C), except the values on the diagonal (harmonic interactions) have been zeroed for better visualization of the off-diagonal terms (anharmonic interactions). The shaded gray and red regions in (B) represent the statistical distributions in NEB (see Figure 1B) and MD-based modal decompositions, respectively. MD results shown on this plot are obtained from natural simulation at 800 K.

density of states (DOS) for lithium ions shown in Figure 5B, obtained from the partial DOS in Figure 5A, also shows reasonable agreement with the modal contributions obtained from the NEB-based and MDMA approaches. The generality of this
agreement should be investigated more in future studies; however, NEB and MDMA reveal the exact individual contributing modes to each ion hop in the system.

We use the MDMA formalism to, for the first time, quantify the anharmonic interactions that lead to ion diffusion in a solid lattice. Individual values of $D_{a,n}$ (see Equation 11 and the experimental procedures section) are shown on maps of correlation in Figures 5C and 5D. By summing over all the pairwise correlations in Figure 5C, we confirmed that >92% of the contributions to diffusivity arise from the correlations along the diagonal (iso-frequency modes), indicating that most of the contributions by phonons to the diffusivity in the Ge-substituted Li$_3$PO$_4$ system originate from the harmonic interactions. The remaining contributions (~8%) came from off-diagonal terms (anharmonic interactions), which can be observed better in Figure 5D after omitting the diagonal terms (i.e., artificially setting them to zero). Even after removing the diagonal terms, all the large contributions are still close to the diagonal, which shows that all the anharmonic terms are also between the modes that have relatively similar frequencies. Quantifying anharmonic interactions is particularly important for fast ion conductors because the lower migration barriers in those ion conductors have been shown to be correlated with softer lattices, which typically allow for greater degrees of anharmonic interactions. In a recent study, using AIMD simulations and Raman polarization-orientation measurements, Brenner et al. attributed the large broadening of vibrational peaks in AgI superionic conductor to the strong anharmonic interaction of the Ag$^+$ ion and the rigid iodine lattice. Our chosen Ge-substituted Li$_3$PO$_4$ compound is not a superionic conductor, which could be the reason for the low degrees of anharmonic contributions that we calculated for ion conduction from the MDMA method. Overall, our analysis provides a quantitative way to measure the effect of anharmonicity on the solid-state diffusion, which has already been recognized in earlier works, but little quantitative evidence has been provided, and that is, with the exception of a few compounds, such as CuCrSe$_2$ and AgCrSe$_2$, where recently, inelastic neutron-scattering measurements have revealed a softening of acoustic modes, which are associated with the superionic transitions in those materials. Ultimately, more studies are needed to compare the inelastic interactions observed for our Ge-substituted Li$_3$PO$_4$ compound with other ion conductors with more-polarizable anions in which the anharmonicity is expected to be more important.

We have introduced two modal decomposition formalisms (see experimental procedures) that determine the individual normal mode contributions to ion diffusion. Our results for a model structure of Ge-substituted Li$_3$PO$_4$ show that a small group of modes (<10% of the vibrational modes in the system) are responsible for >87% of the diffusion. Of greatest significance is that, when we externally excited a targeted subset of the highest-contributing modes, the diffusion increased to the value associated with the effective temperature of the targeted modes, even if the lattice is kept at a lower temperature. This observation is intriguing because a high diffusivity can be achieved while the bulk temperature of the material remains at a much lower temperature.

Although the selected Ge-substituted Li$_3$PO$_4$ composition in this study comprises a rather slow ion conductor, we think the presented, rigorous theoretical frameworks in this report can be used to study the fundamental concept of phonon-ion interaction, which is important in the field of superionics. Recent reports have, so far, only linked the idea that phonons contribute to ion transport, but this is the first study showing that specific-mode excitations can be used to influence diffusion in solids, which is influential to the field of lattice dynamics affecting ion transport.
and will most likely lead to more work on optoionics. In addition, the idea that a diffusion hop can be enhanced directly by phonon excitation has far-reaching implications. If such an effect can be confirmed experimentally (see Note S3 for a possible approach to experimental evaluation of the reported observations in this study), it would open the door to many new possibilities. For example, it may be possible to operate systems or devices that have portions that are diffusion limited at lower temperatures, which could lead to major cost savings and performance improvements. For instance, hypothetically, if oxygen diffusion through YSZ were limiting in solid oxide fuel cells (SOFCs), and the diffusivity it exhibits at a bulk temperature of 1,100 K could be achieved at a much lower temperature of 300–400 K, via some other means, there might be a dramatic effect on lowering the cost, enhancing the efficiency, and increasing the life and proliferation, at the system level by using cheaper materials and seals for the piping/infrastructure. Furthermore, it is likely that this effect can be generalized to reactions, enabling a type of phonon catalysis, whereby only the modes that matter are targeted and deliberately excited to accelerate a reaction. Similarly, this effect may be generalizable to phase transitions and other transformations, or possibly even the reverse, i.e., the suppression of a reaction or phase transition by reducing the amplitude of the modes that facilitate the transition(s). The findings reported in this study, therefore, provide a gateway to thinking about new ways that specific vibrations can be used to enhance or potentially inhibit diffusion, reactions, and/or phase transitions, etc.

EXPERIMENTAL PROCEDURES

Resource availability

Lead contact

Further information and requests for resources should be directed to and will be fulfilled by the lead contact, Asegun Henry (ase@mit.edu).

Materials availability

This study did not generate new unique reagents.

Data and code availability

All data are available from the lead contact upon reasonable request.

Modal decomposition based on NEB calculations

Ion diffusion in solids typically involves the translation of an ion from one location to another; in which, the starting and final locations usually correspond to ion configurations that are local minima in the potential energy surface (see Figure 6). Consequently, each of the equilibrium ionic configurations at the beginning and end of a hop could have a different set of normal modes of vibration. Thus, although we may have one minimum energy pathway (MEP) between equilibrium configurations 1 and 2 (see Figure 6), because of the different vibrational modes at these two energy minima, the modes that initiate an ion hop from site 1 to site 2 may be different from the ones that initiate a hop from site 2 to site 1. More explanation on how two different equilibrium configurations in one structure possess different normal modes of vibration is provided in Note S4. To determine which modes of vibration in a local equilibrium contribute to the ion hop along a specific pathway, we combine NEB and lattice dynamics (LD) calculations.

An NEB calculation determines the MEP between two equilibrium configurations. Each NEB calculation results in a desired number of ionic configurations (snapshots of displaced ions) that are distributed along the MEP between the two end-equilibrium configurations. According to the LD formalism, any atomic displacement field
can be projected onto any complete set of normal modes of vibration to determine how the normal modes superpose to recreate that exact displacement field. The magnitude of projection, as has been recently learned from modal decomposition of thermal transport,\textsuperscript{44–47} shows the degree to which each normal mode is contributing to that displacement field, which can be quantified using the following expression\textsuperscript{14,47},

\[
Q_n = \sum_{i=1}^{N} \sqrt{m_i} \mathbf{e}_{i,n}^* \cdot \mathbf{u}_i \quad \text{(Equation 1)}
\]

where \(\mathbf{u}_i\) is the displacement of atom \(i\) from its equilibrium position in the configuration at the hopping origin; \(\mathbf{e}_{i,n}\) is the eigenvector for mode \(n\), assigning the direction and displacement magnitude of atom \(i\) obtained from an SCLD calculation\textsuperscript{45,47,63} for the structure at the hopping origin; \(*\) denotes the complex conjugate operator; and \(m_i\) is the mass of atom \(i\). \(Q_n\) is the modal displacement coordinate, the square of which is proportional to the mode potential energy \(E_n\) according to Equation 2:

\[
E_n = \frac{1}{2} \mathbf{u}_i^2 Q_n^2 \quad \text{(Equation 2)}
\]

The total energy of the system \(E\) is equal to the summation over all modal energy values, \(E = \sum E_n\). Here, \(E_n\) is the contribution by mode \(n\) to the potential energy of the displaced lattice during the ion migration, which can be interpreted as the contribution by mode \(n\) to the ion hop along its migration pathway. It should be noted that, in the presented NEB-based modal analysis, the definition for modal kinetic energy\textsuperscript{14} could not be used as a metric for analyzing the modal contributions to the ion hop because the atomic velocities are not available in an NEB calculation. To perform the projection of the displaced atoms onto the normal modes, a choice must be made as to which configuration along the MEP should be used to perform the projection. Although that is somewhat arbitrary, for simplicity, we selected the configuration that is halfway between the hopping origin and the transition state (see Figure 6). We avoided projection on points closer to the saddle point based on the practical consideration that the structure would likely exhibit modes with imaginary frequencies near the transition state, which might introduce undesirable

\begin{figure}[h]
\centering
\includegraphics[width=0.5\textwidth]{ MEP_schematic.png}
\caption{MEP schematic between three minimum energy sites (sites 1, 2, and 3). Migration barrier (MB) is the maximum energy difference along the MEP (e.g., \(MB = E_{\text{transition state}} - E_1\) for the hop from site 1 to site 2) that the hopping ion needs to overcome. The dots on the MEP between sites 1 and 2 show the ionic configurations (displacement fields) obtained from the NEB calculation. The displacement-field halfway points to the transition state, shown by the solid and dashed squares, are used to obtain the modal contributions to the ion hops from sites 1 to 2 and from sites 2 to 1, respectively.}
\end{figure}
complexity, the effect of which will be analyzed in follow-on studies. Nevertheless, we examined the projections closer to the beginning stages of the MEP, compared them to those at the halfway point, and noticed almost no change in the contributing modes (Figure S10), which suggests the choice of the halfway point is satisfactory, at least for the system under consideration. To calculate the modal contributions for all the possible hops in the structure, we applied the above procedure to (1) all the distinct equilibrium configurations in the system because each of them would have different vibrational modes, and (2) all the ion hops in each of those equilibrium configurations.

**NEB calculation details**

To test our formulations, we chose Ge-substituted Li$_3$PO$_4$ as an example structure. The interatomic potentials for NEB calculations and MD simulations were obtained from a well-established form potential, which has been successfully tested in a number of previous studies, including with Li-conducting compounds. This form potential comprises a long-range coulomb term, a short-range Morse function, and a repulsive term. The parameters were obtained from the library of potentials developed by Pedone et al., which has been successfully tested in previous MD simulations of silicates and polyanion type materials, including Li$_3$PO$_4$. The exact formulas and parameters of which are provided in Table S3. To further check the accuracy of this interatomic potential used, we calculated the lattice constants for the perfect γ-Li$_3$PO$_4$ crystal from the isobaric-isothermal MD relaxation at zero pressure and room-temperature and compared them with the existing first-principles and experimental values (Table S4). The comparison showed that the lattice constants in our simulations have <3% difference with other reported values. The pristine Li$_3$PO$_4$ structure was chosen to be a 3 × 2 × 1 supercell containing 192 atoms, (γ-Li$_3$PO$_4$, space group Pnma) with the lattice dimensions given is Table S4. In the pristine structure, one P was substituted by one Ge, and one Li was added in the form of an interstitial to maintain charge neutrality; hence, the structure contains 193 ions (Li$_{3.042}$Ge$_{0.042}$P$_{0.958}$O$_4$). This composition is a simple case to show the effectiveness of the modal excitation approach in this article. If the number of Ge-substitute atoms increases, for charge neutrality, the number of Li interstitials should also increase. More interstitial sites complicate the modal-excitation experiment because of the possibility of simultaneous hops in the simulation and the consequent challenge in deciding which modes to excite during the simulation. There are 36 available, unique interstitial sites to add this Li interstitial. However, among those 36 interstitial sites, only 31 resulted in a stable configuration with positive vibrational frequencies (Figure S5 shows schematics of those distinct configurations). Each hopping event brings the structure from one of those equilibrium configurations to another.

As is shown in the results section, the dominant hopping mechanism in our Ge-substituted Li$_3$PO$_4$ is the interstitialcy mechanism. To determine the phonon contributions to all the possible interstitialcy hops in the system, we need to account for all the interstitialcy hops in each of the equilibrium configurations. To do so, we first counted the nearest-neighbor Li lattice sites to the interstitial ion. Then, for each of those nearest-neighbor Li lattice sites (tetrahedral sites), we counted the nearest-neighbor interstitial sites (octahedral sites), with the constraint that the nearest-neighbor interstitial site be among the 31 stable interstitial sites. Counting for all those combinations, resulted in 619 distinct hops in our structure that follow the interstitialcy mechanism. The SCLD calculations determined the normal modes of vibration for all 31 equilibrium configurations and were performed using the general utility lattice program (GULP) package, whereas the MD simulations and the NEB
calculations were conducted using the large-scale atomic/molecular massively parallel simulator (LAMMPS)\textsuperscript{93} package, following the formulations by Henkelman et al.\textsuperscript{83} to ensure the correct determination of the saddle point. The reasons for why our calculations are based on classical interatomic potentials and LAMMPS package are discussed in \textit{Note S5}.

**Targeted excitation of modes to enhance diffusivity**

After finding the strongest contributing modes to the ion diffusion using the NEB approach, we examined whether external excitation of a small subset of those modes in an MD simulation had any measurable effect on the tracer and conductivity ion diffusivities. At each time step during the simulation, the equilibrium configuration was found by checking where the interstitial ion was located (i.e., which octahedral site was occupied) because the distinction between equilibrium configurations was based only on the occupied interstitial site and because the relative position of non-interstitial ions remained unchanged across different equilibrium configurations. By knowing the equilibrium configuration, we then excited the five highest-contributing modes among all the contributing modes to all the possible hops belonging to that configuration and kept the bulk temperature fixed. After a hop happened, a new equilibrium configuration was detected, and the process continued by exciting the five highest-contributing modes among all the possible hops associated with the new configuration.

Because ion diffusivity increases with temperature, we ensured that the bulk lattice temperature remained constant at a low temperature, so that any change in the diffusivity could be attributed only to the excitation of the targeted modes and not to the bulk temperature. To do so, we kept the total kinetic energy of the system constant via a velocity-rescaling scheme, in which, the addition of energy to the top five modes was complimented by a uniform reduction in the kinetic energy of all other modes in the system. To change the temperature of mode \( n \) to a desired temperature \( T_{d} \), we modified the atomic velocities in the system according to the following formula:

\[
v_{i} = v_{i} + \frac{1}{\sqrt{m}} \left( \sqrt{2k_{B}T_{d}} - \tilde{Q}_{n}(t) \right) \mathbf{e}_{i,n}
\]

(Equation 3)

where \( v_{i} \) is the velocity of atom \( i \), and \( \tilde{Q}_{n} \) is the modal velocity coordinate defined by\textsuperscript{14}

\[
\tilde{Q}_{n} = \sum_{i=1}^{N} \sqrt{m} \mathbf{e}_{i,n} \cdot \mathbf{v}_{i}
\]

(Equation 4)

The derivation of Equation 3 is provided in \textit{Note S6}. This rescaling procedure induces only a slight perturbation to the system, given that the top-five modes comprise only 0.86\% of the modes in the system, and thus, the reduction in energy for all other modes is only <1\% of their energy (see \textit{Note S7} and Table S5 for discussions about the needed thermalization power). In this study, the above-mentioned rescaling procedure was applied every five-times steps (every 5 fs). Although a velocity-perturbation scheme has been used in this study, other methods for modal excitation, such as atomic position perturbation\textsuperscript{94} or simultaneous perturbation of atomic positions and velocities,\textsuperscript{95} have also been employed in previous studies, particularly those with the goal of investigating the phonon-phonon interactions in MD simulations. However, the chosen method for modal excitation should not change the reported observations in this study because modal excitation (increasing the energy of a mode) can be achieved either by increasing the potential (position perturbation) or the kinetic (velocity perturbation) energy of the mode; the choice of which should not matter.
Modal decomposition based on MD simulations

To directly obtain the contributing modes to the ion diffusion from MD simulations, we used the definition of diffusivity based on the fluctuation-dissipation theorem\(^ {96}\) and followed a modal-decomposition approach similar to that employed in MD-based thermal-transport studies.\(^ {44–47}\) Consistent with prior work on phonon transport,\(^ {44–47}\) we have termed this formalism “mass diffusivity modal analysis” (MDMA). In MDMA, the modal contributions to the atomic velocities are obtained and then used to decompose the following definition for conductivity diffusivity (\(D_s\)):\(^ {67,75}\)

\[
D_s = \frac{N_c}{3} \int_0^\infty \langle v_c(\tau)v_c(0) \rangle \, d\tau \quad \text{(Equation 5)}
\]

Conductivity diffusivity (\(D_s\)) not only includes the self-correlation information embedded in the tracer-diffusivity definition (Equation S11), but it also includes the many-particle-correlation information. In Equation 5, \(N_c\) is the number of hopping particles in the system, \(\langle \cdot \rangle\) denotes the auto-correlation operator, and \(v_c\) is the center of mass velocity for the hopping particles:

\[
v_c = \frac{1}{N_c} \sum_{i=1}^{N_c} v_i \quad \text{(Equation 6)}
\]

To calculate the individual contribution by mode of vibration \(n\) to \(v_i (v_{i,n})\), first modal velocity coordinate \(\hat{Q}_n\) is calculated (Equation 4),\(^ {14}\) from which, \(v_{i,n}\) is determined by\(^ {14}\)

\[
v_i = \frac{3N}{N_c} \frac{\hat{Q}_n(t)}{\sqrt{m_i}} e_{i,n} = \frac{3N}{N_c} \sum_{n=1}^{N_c} v_{i,n} \rightarrow v_{i,n} = \frac{\hat{Q}_n(t)}{\sqrt{m_i}} e_{i,n} 
\quad \text{(Equation 7)}
\]

Replacing \(v_i\) in Equation 6 with its modal contributions (\(v_{i,n}\)) results in the modal contributions to the carriers’ center of mass:

\[
v_c = \sum_{n=1}^{N_c} \frac{1}{N_c} \sum_{i=1}^{N_c} v_{i,n} \quad \text{(Equation 8)}
\]

which is substituted in one of the \(v_c\) terms in Equation 5 to extract the modal contributions to diffusivity (\(D_{s,n}\)),

\[
D_s = \sum_{n=1}^{3N} \frac{1}{3} \int_0^\infty \langle \sum_{i=1}^{N_c} v_{i,n}(\tau) v_c(0) \rangle \, d\tau = \sum_{n=1}^{3N} D_{s,n} 
\quad \text{(Equation 9)}
\]

Similarly, the modal contributions to ion conductivity (\(\sigma_{i,n}\)) can be obtained; the details of which are provided in Note S8.

By substituting for both center-of-mass velocities in Equation 5 with their modal contributions from Equation 7, the above formulations can be extended to capture the modal contributions to diffusivity in more detail:

\[
D_s = \sum_{n=1}^{3N} \sum_{n'=1}^{3N} \frac{1}{3} \int_0^\infty \langle \sum_{i=1}^{N_c} v_{i,n}(\tau) \sum_{j=1}^{N_c} v_{j,n'}(0) \rangle \, d\tau = \sum_{n=1}^{3N} \sum_{n'=1}^{3N} D_{s,n,n'} 
\quad \text{(Equation 10)}
\]

where individual contributions from the correlation/interaction of eigen mode pairs \(n\) and \(n'\) (\(D_{s,n,n'}\)) are given by,

\[
D_{s,n,n'} = \frac{1}{3} \int_0^\infty \langle \sum_{i=1}^{N_c} v_{i,n}(\tau) \sum_{j=1}^{N_c} v_{j,n'}(0) \rangle \, d\tau 
\quad \text{(Equation 11)}
\]
which provides additional insight into the degree to which each pair of modes interact, i.e., the anharmonicity via terms in which \( n \neq n' \), to facilitate diffusion.

The combination of the MD- and NEB-based modal decomposition methods presented herein provides a detailed picture for the ion diffusion. Although both methods are based on projecting the trajectory of the hopping ion on the eigenvectors of vibration, the way they sample the phonon contributions to the ion hop is different. In the NEB-based approach, the contributions are calculated based on the atomic displacement field, whereas in the MD-based method, the contributions are obtained based on the atomic velocity field. In addition, MD samples the dominant mechanisms that are present in a natural simulation of the material, whereas NEB can be used to study specific mechanisms and the ion hop along specific migration pathways, regardless of the likelihood that they would occur at a given temperature.

**Diffusivity calculations**
The total mean squared displacement method (TMSD) was used to measure the diffusivity of Li ions in our structure; the details of which are explained in the existing literature\(^9\) and in Note S9. First, the structure was relaxed under the isobaric-isothermal ensemble for 1 ns at zero pressure and \( T = 400 \) K and under the Nosé-Hoover canonical ensemble for another 1 ns at \( T = 400 \) K; then, the structure was simulated for another 10 ns under the same ensemble to obtain the needed displacement data for diffusivity calculations. The time step was chosen to be 1 fs, and statistical uncertainty was reduced by considering five independent ensembles.\(^9\) For calculating the conductivity-diffusion coefficients, total conductivity was calculated following the existing literature;\(^67\) the details of which are also explained in Note S9.

**SUPPLEMENTAL INFORMATION**
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**AUTHOR CONTRIBUTIONS**
K.G., Y.S.-H., and A.H. conceived the idea. K.G. and A.H. derived the decomposition formulations. K.G. and S.M. performed the simulations. All authors contributed to the analyses, discussions, and writing of the manuscript.

**DECLARATION OF INTERESTS**
Y.S.-H. is a board member at *Cell Reports Physical Science*. The remaining authors declare no competing interests.

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