

Lattice-Geometry Effects in Garnet Solid Electrolytes from Lattice-Gas Monte Carlo Simulations

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In many solid electrolytes, ion transport can be approximated as a sequence of hops between distinct lattice sites. Assuming these hops are uncorrelated allows quantitative relationships to be derived between microscopic hopping rates and macroscopic transport coefficients; tracer diffusion coefficients and ionic conductivities. In real materials, hops are uncorrelated only in the dilute limit, where interactions between mobile ions can be neglected. At non-dilute concentrations these interactions can be significant, causing hops to become correlated. In these cases the relationships between hopping frequency, diffusion coefficient, and ionic conductivity deviate from the random walk expressions, with this deviation quantified by single-particle and collective correlation factors, f and f_1 . These factors vary between materials, and depend on the concentration of mobile particles, the nature of the interactions, and the host lattice geometry. Here we study these correlation effects for the garnet lattice using lattice-gas Monte Carlo simulations. This lattice represents diffusion pathways in lithium-garnets—a family of promising solid lithium-ion electrolytes—and has an unusual geometry containing both 4-coordinate and 2-coordinate sites. We find that for non-interacting particles (volume exclusion only) single-particle correlation effects are more significant than for any other known three-dimensional solid-electrolyte lattice. This is attributed to the presence of 2-coordinate lattice sites, which causes correlation effects intermediate between typical three-dimensional and one-dimensional lattices. By including nearest-neighbour repulsion and on-site energies we predict more complex single-particle correlations and also collective correlations. We predict particularly strong correlation effects at $x(\text{Li}) = 3$ (from site energies) and $x(\text{Li}) = 6$ (from nearest-neighbour repulsion). Both effects correspond to ordering of the mobile particles over the lattice. Finally we consider the question of how mobile ion stoichiometry can be tuned to maximise the ionic conductivity, and show that the “optimal” lithium content is highly sensitive to the precise nature and strength of the microscopic interactions.

I. INTRODUCTION

The ability of solid electrolytes to conduct electric charge by transporting ions is central to their use in devices such as fuel cells and solid-state lithium-ion batteries.^{1–4} In both cases, solid electrolytes with high ionic conductivities are desirable. In fuel cells high conductivities allow lower operating temperatures, reducing running costs and increasing operating lifetimes. In solid-state batteries high conductivities allow faster charging rates and higher power outputs. Ionic conductivities depend on a number of factors, including the crystal structure, the chemical composition, and the concentration of mobile ions.⁵ Developing a quantitative understanding of these factors is key to developing high conductivity solid electrolytes for use in high performance electrochemical devices.

Solid electrolytes can be considered to comprise two distinct sets of ions: “fixed” ions that vibrate about their crystallographic sites, and “mobile” ions that diffuse through the system and contribute to ionic conductivity. The fixed ion positions define a network of diffusion pathways through which the mobile ions move. Solid electrolytes with common crystal structures have diffusion networks that are topologically equivalent, while electrolytes with different crystal structures will have topologically distinct diffusion paths. While much research into solid electrolytes focusses on understanding

differences in ionic conductivities within specific structural families, a complementary question considers how differences in crystal structure, and hence diffusion network topology, affect ionic transport.

Crystal structure can be considered a microscopic characteristic, dictating the positions of individual atoms. Ionic transport at scales relevant to device performance, however, is described by macroscopic transport coefficients: diffusion coefficients and ionic conductivities. These describe long-time behaviours, and represent ensemble averages over all microscopic diffusion processes. A meaningful understanding of differences in ionic conductivity between solid electrolytes therefore depends on the quantitative relationships between microscopic diffusion and macroscopic ion transport. This study considers these relationships for the lithium-garnet family of solid lithium-ion electrolytes, using lattice-gas Monte Carlo simulations.

In many solid electrolytes, the microscopic transport of ions can be approximated as a sequence of discrete “hops” between distinct lattice sites.⁶ If these hops are *independent*, every ion follows a random walk. The tracer diffusion coefficient, D^* , and ionic conductivity, σ , can then be expressed in terms of the average hop-rate per atom, $\tilde{\nu}$,⁷ via^{8,9}

$$D^* = \frac{1}{6} a^2 \tilde{\nu}; \quad (1)$$

$$\sigma = \frac{Cq^2}{kT} \frac{1}{6} a^2 \tilde{\nu}; \quad (2)$$

where a is the characteristic hop distance, C is the mobile ion concentration, and q is the charge of the mobile ions. Equations 1 and 2 can be combined to give the Nernst-Einstein relation, which connects D^* and σ :

$$\frac{\sigma}{D^*} = \frac{Cq^2}{kT}. \quad (3)$$

These three equations provide quantitative relationships between the hop-rate, $\tilde{\nu}$, tracer diffusion coefficient, D^* , and ionic conductivity, σ . Their derivation, however, depends on the assumption of independent hops, which holds only in the limit of very low carrier concentrations, or for fully non-interacting mobile ions.¹⁰

Practical solid electrolytes typically have high carrier concentrations, and interparticle interactions can be significant. In these cases, hopping probabilities depend on specific arrangements of nearby ions, and hops are no longer statistically independent. Instead, ion trajectories are *correlated*, and the system dynamics deviates from random walk behaviour.^{8,11–13} Correlations between hops made by any single ion modify the relationship between average hop rate, $\tilde{\nu}$, and tracer diffusion coefficient, D^* , which becomes

$$D^* = \frac{1}{6} a^2 \tilde{\nu} f, \quad (4)$$

where f is a single-particle correlation factor that accounts for the deviations from random walk behaviour. Correlations between hops made by *different* ions modify the relationship between $\tilde{\nu}$ and σ , which becomes

$$\sigma = \frac{Cq^2}{kT} \frac{1}{6} a^2 \tilde{\nu} f_I, \quad (5)$$

where f_I is a collective or “physical” correlation factor.^{10,14,15} The relationship between σ and D^* now differs from Nernst-Einstein behaviour (Eqn. 3) by the ratio of these correlation factors:

$$\frac{\sigma}{D^*} = \frac{Cq^2}{kT} \frac{f_I}{f}. \quad (6)$$

The inverse ratio $\frac{f}{f_I}$ is commonly denoted as the Haven ratio, H_R .^{10,16}

Quantitative relationships between microscopic hopping rates and macroscopic transport coefficients can, in principle, be obtained by combining experimental data for $\tilde{\nu}$, D^* , and σ . Ion hopping rates may be measured from NMR or muon spin-relaxation experiments,^{17–22} diffusion coefficients obtained from tracer diffusion experiments,²³ and ionic conductivities derived via impedance spectroscopy.^{24,25} Computational methods provide an increasingly useful complement to experimental studies. First principles calculations of vibrational partition functions and barrier heights along diffusion pathways

can be used to obtain hopping rates.^{26,27} Molecular dynamics simulations can be used to directly calculate diffusion coefficients and ionic conductivities.²⁸ Often, however, not all members of $\{\tilde{\nu}, D^*, \sigma\}$ are known, and it is necessary to derive one or more from the other, known, properties. In principle, quantitative conversions between $\{\tilde{\nu}, D^*, \sigma\}$ are possible via Eqns. 4–6, providing the correlation factors f , f_I (and hence H_R) are known.

For many simple crystal lattices, the correlation parameters $\{f, f_I, H_R\}$ have been calculated.^{10,29} For more complex crystal structures, however, these parameters are often still unknown. A common approximation, therefore, is to assume correlation effects can be neglected, which allows the simpler Eqns. 1–3 to be used. This approximation is equivalent to assuming dilute-limit non-interacting behaviour. In solid electrolytes ionic motion exhibits strong correlations, however, this can introduce quantitative errors when analysing data.

In this study, we describe lattice-gas Monte Carlo simulation of garnet-structured solid electrolytes, to study quantitative correlation effects in these materials. The garnet lattice represents the diffusion pathways in the “lithium-garnets”. This family of solid lithium-ion electrolytes $\text{Li}_x\text{M}_3\text{M}'_2\text{O}_{12}$,^{30,31} have attracted significant attention as candidates for all-solid-state lithium-ion batteries.^{1,32–34} The garnet crystal structure has an unusual three-dimensional network of lithium diffusion pathways, consisting of interlocking rings.³⁵ Each ring comprises 12 alternating tetrahedral and octahedral sites. The tetrahedral sites are coordinated to four octahedral sites, while the octahedral sites are each coordinated to two tetrahedral sites. The tetrahedral sites therefore act as nodal points connecting adjacent rings (Fig. 1). Aliovalent substitution of the M and M' cations allows the lithium stoichiometry to be tuned across a broad range. A lithium stoichiometry of $x_{\text{Li}} = 9$ corresponds to a fully occupied lithium-site lattice, and research has focussed on “lithium stuffed” garnets, typically with $x_{\text{Li}} = 5$ to 7. Ionic conductivities vary enormously as a function of x_{Li} , with σ increasing by $\sim 10^9$ between $\text{Li}_3\text{Ln}_3\text{Te}_2\text{O}_{12}$ and $\text{Li}_{6.55}\text{La}_3\text{Zr}_2\text{O}_{12}$,^{1,31} and it remains an open question precisely how the lithium diffusion coefficient and ionic conductivity vary with lithium stoichiometry. It is also not known to what extent the unusual diffusion pathway topology affects ionic transport. Resolving these questions is critical for the optimisation of ionic conductivity for this family of materials.

Structural considerations and published data both suggest lithium-garnets may exhibit significant correlation effects. The low connectivity of the two-coordinate octahedral sites means blocking effects are expected to be considerable.³⁵ Short distances between neighbouring lattice sites of $\sim 2.4 \text{ \AA}$ suggest strong Li–Li repulsion, with particular significance at high Li stoichiometries.^{36–39} The presence of two non-equivalent sets of lattice sites is also a factor. Non-interacting lithium ions would be expected to occupy octahedral and tetrahedral sites in a 2:1 ratio, reflecting the rela-

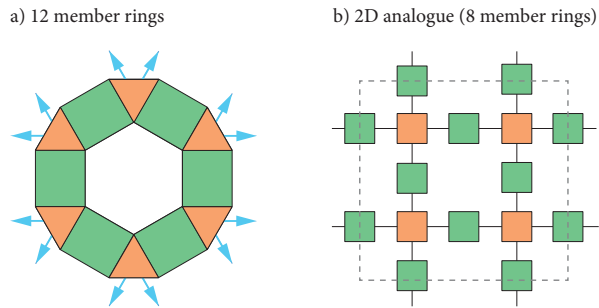


FIG. 1. Schematic of the ring structures that constitute the garnet lithium-diffusion network. a) Each ring consists of 12 alternating tetrahedra (orange) and octahedra (blue). Arrows show connections to neighbouring rings.³⁵ b) A 2D analogue of interconnected 8-membered rings of alternating “tetra-dra” and “octahedra”.

tive site populations. Neutron data, however, show that at low lithium content ($x_{\text{Li}} = 3$) only tetrahedral sites are occupied,⁴⁰ while at higher lithium content ($x_{\text{Li}} = 5 \rightarrow 7$) octahedral sites become preferentially occupied.^{31,38} Experimental conductivities show non-linear dependence on x_{Li} ,⁴¹ and deviate strongly from ideal values predicted (via Eqn. 2) from muon-spin-spectroscopy hopping rates.²¹ Further evidence for correlated transport in lithium garnets comes from computational studies. A variety of correlated diffusion processes have been observed in molecular dynamics simulations,^{42–45} and calculated diffusion coefficients and ionic conductivities show non-Nernst-Einstein behaviour ($H_{\text{R}} < 1$).^{46,47} Together, these results indicate the existence of significant interactions, either between lithium ions or between these ions and the host lattice. The quantitative effects of correlation in lithium garnets, however, are not known, and consequently studies often rely on assumptions of uncorrelated motion to convert between hop rates, diffusion coefficients, and ionic conductivities.^{21,22,24,42,48–56}

Here we present a computational study of these correlation effects, comprising lattice-gas kinetic Monte Carlo simulations of diffusion on a garnet lattice, across a range of model Hamiltonians. We calculate f and f_{I} as functions of lithium stoichiometry (carrier concentration), first for a non-interacting volume-exclusion model,⁵⁷ and then for models that include on-site single-particle energies and/or nearest-neighbour repulsion interactions. In addition to self- and collective-correlation factors, we present site occupation populations, diffusion coefficients, and reduced ionic conductivities for this range of simulation models. Our results demonstrate how different interactions contribute to non-ideal behaviour, and modify the relationships between particle hopping rate, diffusion coefficient, and ionic conductivity.

We find that for non-interacting particles (volume exclusion only) single-particle correlation effects are more significant than for any other 3D solid electrolyte lattice. This is attributed to the presence of 2-coordinate lat-

tice sites, which produces correlation effects intermediate between typical 3D and 1D lattices. Including nearest-neighbour repulsion and on-site energy differences gives more complex single-particle correlation behaviour and introduces collective correlation. In particular, we find strong correlation effects at $x(\text{Li}) = 3$ (due to site energy differences) and $x(\text{Li}) = 6$ (due to nearest-neighbour repulsion). Both effects correspond to ordering of the mobile particles over the lattice, and sharp decreases in diffusion coefficients and ionic conductivities. Finally we consider the question of how mobile ion stoichiometry can be tuned to maximise the ionic conductivity. We find this does not have a straightforward answer, and the optimal stoichiometry is highly sensitive to the choice of interaction parameters.

II. METHODS

Lattice-gas Monte Carlo simulations describe the diffusion of a set of mobile ions populating a host lattice, expressed as a graph of interconnected sites.⁵⁸ Every lattice site is either occupied or vacant, and during a simulation the mobile ions undergo a sequence of hops from site to site. These hops are randomly selected, with relative probabilities that satisfy the principle of detailed balance and represent the underlying model Hamiltonian. The simplest model considered here is a non-interacting volume-exclusion-only model.⁵⁹ Double occupancy of sites is forbidden, and allowed hops are all equally likely. Non-interacting models allow the pure geometric effect of the lattice to be evaluated, but neglect other interactions that may be important in otherwise equivalent experimental systems. Here we extend the non-interacting model to also consider the effect of nearest-neighbour interactions between mobile ions, described by a nearest-neighbour repulsion energy, E_{nn} , and interactions between single ions and the lattice, described by on-site energies for tetrahedral versus octahedral sites, E_{tet} , E_{oct} . The energy of any configuration of occupied sites, j is given by

$$E = \sum_j n_j^{\text{nn}} E_{\text{nn}} + E_{\text{site}}^j, \quad (7)$$

where n_j^{nn} is the number of occupied nearest neighbour sites for (occupied) site j . For interacting systems, the relative probability of hop i depends on the change in total energy if this hop was selected, ΔE_i ,

$$P_i \propto \begin{cases} \exp\left(\frac{\Delta E_i}{kT}\right), & \text{if } \Delta E_i > 0 \\ 1, & \text{otherwise.} \end{cases} \quad (8)$$

For our interacting systems, the change in energy for each candidate hop can depend on the change in number of nearest-neighbour interactions and the change in on-site energy for moving from a tetrahedral to octahedral site (or vice versa):

$$\Delta E_i = \Delta n_{\text{nn}} E_{\text{nn}} \pm \Delta E_{\text{site}}, \quad (9)$$

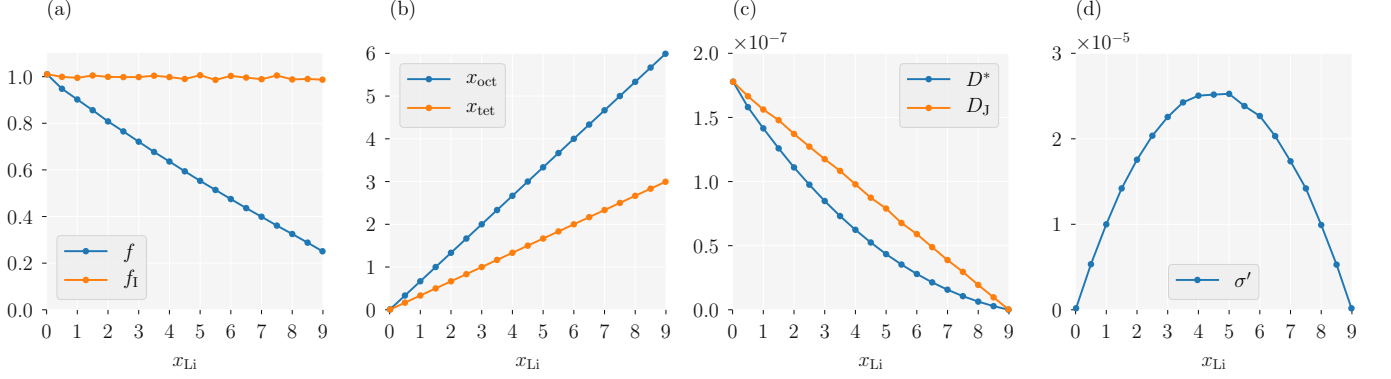


FIG. 2. Non-interacting particles on a garnet lattice: (a) The single-particle correlation factor, f , and collective correlation factor, f_I ; (b) Average octahedral and tetrahedral site occupations per formula unit, x_{oct} and x_{tet} ; (c) Tracer diffusion coefficient, D^* , and “jump” diffusion coefficient D_J . (d) Reduced ionic conductivity, σ' .

where $\Delta E_{\text{site}} = E_{\text{oct}} - E_{\text{tet}}$. At each simulation step, one hop is randomly selected according to the set of relative probabilities. The corresponding ion is moved, and a new set of relative hop probabilities is generated for the subsequent simulation step.

In the limit of a large number of hops, the tracer- and collective-correlation factors can be evaluated as

$$f = \frac{\sum_i \langle R_i^2 \rangle}{Na^2}, \quad (10)$$

where $\langle R^2 \rangle$ is the mean-squared displacement of the mobile ions, and N is the total number of hops during the simulation,²⁶ and

$$f_I = \frac{|\sum_i R_i|^2}{Na^2}, \quad (11)$$

where $\sum_i R_i$ is the *net* displacement of all mobile particles. In both cases the denominators correspond to the limiting behaviour for uncorrelated diffusion.

To allow time-dependent properties to be evaluated, such as average site occupations and transport coefficients, we perform our simulations within a rejection-free kinetic Monte Carlo scheme.⁶⁰ At each simulation step, k , the set of relative hop probabilities, $\{P_{i,k}\}$, are converted to rates, $\{\Gamma_{i,k}\}$, by scaling by a common prefactor $\nu' = 10^{13} \text{ s}^{-1}$. After selecting a hop, the simulation time is updated by $\Delta t = Q_k^{-1} \ln(1/u)$, where Q_k is the “total rate”; $Q_k = \sum_i \Gamma_{i,k}$, and u is a uniform random number $u \in (0, 1]$.

Our lattice-gas kinetic Monte Carlo simulations were performed using the `lattice_mc` code.⁶¹ Simulations were performed for an ideal cubic $2 \times 2 \times 2$ garnet lattice, with 384 octahedral sites and 192 tetrahedral sites. The lattice-site coordinates were generated from the cubic high-temperature $\text{Li}_7\text{La}_3\text{Zr}_2\text{O}_{12}$ (LLZO) structure,³⁵ using the centres of the octahedra and tetrahedra defined by the oxide sublattice. In cubic LLZO, each lithium-octahedron contains a “split” pair of distorted 96h sites,

separated by 0.81 \AA . The construction used here considers each octahedron as a single ideal $48g$ site. The graph of diffusion pathways includes connections between nearest-neighbour sites only, i.e. all connections are between neighbouring tetrahedral–octahedral pairs. For each simulation, n_{Li} mobile ions are randomly distributed across the lattice sites. We perform 1,000 steps for equilibration, followed by 10,000 production steps.

For each set of model parameters, $\{E_{\text{nn}}, \Delta E_{\text{site}}\}$, simulations were performed across the full range of possible lithium stoichiometry. For a $2 \times 2 \times 2$ garnet supercell, the maximum Li content of $x_{\text{Li}} = 9$ corresponds to $n_{\text{Li}} = 576$. For each set of interaction parameters, data were collected as an average over 5,000 independent trajectories.

III. RESULTS

A. Non-Interacting Particles and Geometric Effects

We first examine the pure geometric effect of the garnet lattice by considering non-interacting particles, where any deviations from random walk behaviour are solely due to blocking effects expressed within the specific lattice geometry. Fig. 2 shows for the volume-exclusion-only simulations, as a function of x_{Li} , (a) the calculated self- and collective-correlation factors, f and f_I , (b) average tetrahedral and octahedral site occupations, n_{tet} and n_{oct} , (c) tracer and “jump” diffusion coefficients, D^* and D_J , (d) and a reduced ionic conductivity, σ' (Eqn. 13).

In the single particle limit, $x_{\text{Li}} \rightarrow 0$, both correlation factors equal 1. There are no blocking effects, and particles follow a random walk. With increasing concentration of mobile ions, however, single particle diffusion increasingly deviates from random walk behaviour. The tracer correlation factor, f , decreases from $f = 1$ in the single particle limit $f = 0.25$ in the single vacancy limit $x_{\text{Li}} \rightarrow 9$, showing approximately linear dependence on

x_{Li} .⁶²

The magnitude of the tracer correlation effect can be compared across different lattice geometries by considering f in the limit of a single vacancy, f_v . Table I presents values previously calculated for simple 3D lattices,⁶³ for a 1D chain,¹⁴ and our result for the garnet lattice. The garnet lattice value of $f_v = 0.25$ is smaller than for all of the other 3D lattices listed, and is a factor of two less than the next lowest (the diamond lattice). This indicates that the garnet lattice exhibits particularly strong site blocking effects. This can be understood as a consequence of the garnet lattice geometry. For a general set of 3D lattices, as the number of nearest neighbours of each lattice site, z , decreases, f_v also decreases, and correlation effects become more significant. The garnet lattice has both 4-coordinate (tetrahedral) and 2-coordinate (octahedral) sites, and long ranged diffusion follows an alternating tet→oct→tet→oct sequence. The calculated value of $f_v = 0.25$ is halfway between the values for the 4-coordinate diamond lattice ($f_v = 0.5$) and for a 1-D chain, where every site is 2-coordinate ($f_v = 0$).¹⁴ This suggests that the extreme low value of f_v for the garnet lattice is a consequence of the low coordination of the lattice sites, in particular the local 1D coordination at the octahedral sites, which act as bottlenecks for long-ranged diffusion.

Lattice	z	f_v
Face centered cubic ⁶³	12	0.78146
Body centred cubic ⁶³	8	0.72722
Simple cubic ⁶³	6	0.65311
Diamond ⁶³	4	0.5
Garnet [This work]	4+2	0.25
1D chain ¹⁴	2	0.0

TABLE I. Vacancy correlation factors for some common crystal lattices. z is the number of nearest neighbours for each site in the lattice.

For any non-interacting system, the hops made by *different* particles are uncorrelated, and $f_I = 1$ for all x_{Li} , hence $H_R = f$. There are also no correlations between site occupations, and the mobile particles are randomly distributed over the available octahedral and tetrahedral sites, with a 2:1 population ratio that reflects the underlying lattice geometry.⁶⁴

We also calculate three explicit measures of ionic transport in this system.⁶⁵ Fig. 2(c) shows the tracer diffusion coefficient, D^* (Eqn. 4) and the “jump diffusion coefficient”, D_J ,⁵ calculated as

$$D_J = \frac{|\sum_i R_i|^2}{6Nt}. \quad (12)$$

At a fixed temperature D_J is proportional to the mobility, and measures the ease with which the mobile particles collectively migrate. Both D^* and D_J decrease monotonically from $x_{\text{Li}} = 0$ to $x_{\text{Li}} = 9$ ($x = 0 \rightarrow 1$), as the number of vacancies available to accomodate hopping decreases.

For the non-interacting system there are no correlations between hops made by different particles, and the jump diffusion coefficient is proportional to $(1-x)$ (in the garnet lattice, $x = 1$ corresponds to $x_{\text{Li}} = 9$).^{5,59} The tracer diffusion coefficient, however, is affected by correlations between hops made by individual particles, and varies as $D^* \propto (1-x)f$.⁶⁶ The ionic conductivity of a system depends on both the charge-carrier concentration, and the ionic mobility, which is proportional to D_J . We consider the relative effect of carrier concentration on ionic conductivity by considering a reduced conductivity, σ' ,⁶⁷ given by

$$\sigma' = xD_J. \quad (13)$$

For any non-interacting system, $\sigma' \propto x(1-x)$, giving a maximum at $x = 0.5$, corresponding to $x_{\text{Li}} = 4.5$ for the garnet lattice (Fig. 2(d)).

B. Interacting Particles

The conceptual simplicity of the non-interacting system makes it a useful starting point for understanding the factors affecting ionic transport in different lattices. In real Li-garnet materials, however, interactions between lithium ions, or between lithium ions and the host lattice, can be significant. Lithium ions carry positive charge, and can be expected to experience mutual Coulomb repulsion. The different oxygen-coordination environments of the octahedral and tetrahedral sites can be expected to produce a preference for occupation by lithium at one site versus the other.⁶⁸ Within the lattice-gas Monte Carlo scheme, we consider these two factors by introducing, first, nearest-neighbour repulsion, and second, an octahedral versus tetrahedral site preference.

1. Nearest-neighbour repulsion

To examine the effect of Li-Li repulsion, we consider a simplified model with only nearest-neighbour repulsion. The energy of Li at each specific site now depends on the number of occupied neighbouring sites. Individual hop probabilities therefore now depend on whether they increase or decrease the total number of nearest-neighbour pairs. Fig. 3 presents results from simulations performed for $E_{\text{nn}} = 0.0$ – $3.0 kT$. Repulsive nearest-neighbour interactions disfavour simultaneous occupation of adjacent pairs of sites, which promotes ordering of particles across alternating occupied–vacant–occupied–vacant sites. This ordering causes the single-particle correlation behaviour to deviate from that of the non-interacting system, and also introduces collective correlations between the mobile ions.¹⁰ f and f_I both have their non-interacting values in the empty and fully-occupied lattice limits: $x \rightarrow 0$ and $x \rightarrow 1$. In a lattice with only one crystallographic site complete ordering would occur at half-site-occupancy,

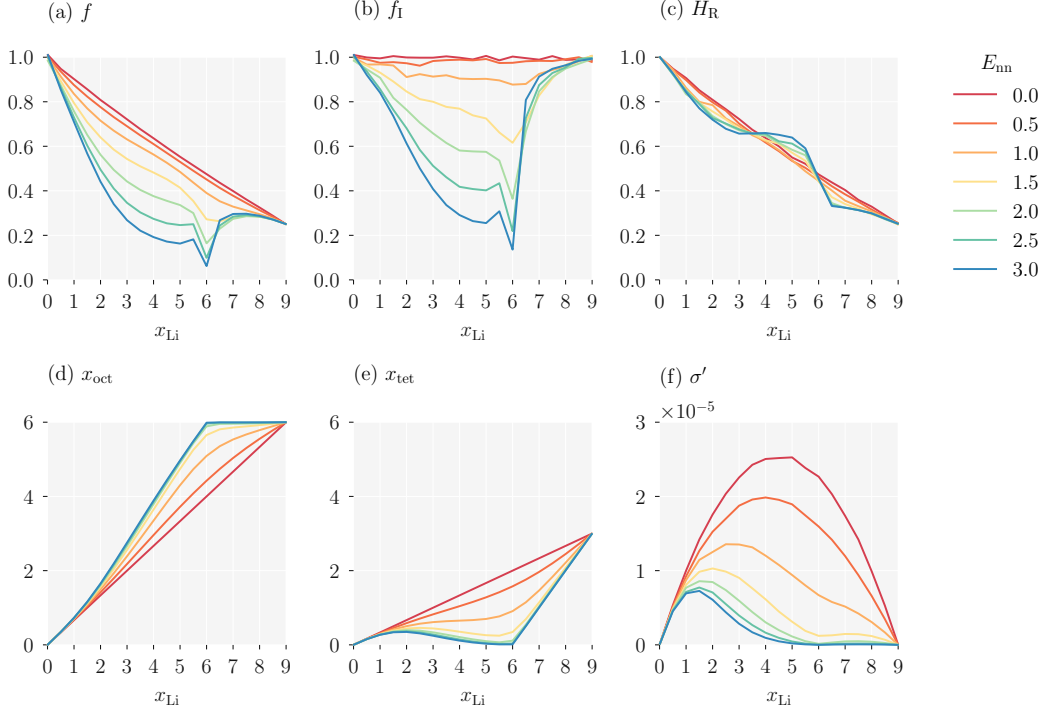


FIG. 3. \Rightarrow x_{tet} and x_{oct} are labelled (a) (and in other plots?) The effect of nearest-neighbour repulsion between mobile particles on a garnet lattice: (a) single-particle correlation factor, f ; (b) collective correlation factor, f_{I} ; (c) Haven ratio, H_{R} ; (d) average octahedra occupation, x_{oct} ; (e) average tetrahedra occupation, x_{tet} ; (f) reduced ionic conductivity, σ' . E_{nn} is in multiples of kT .

corresponding to $x_{\text{Li}} = 4.5$ for the garnet lattice. f and f_{I} approximately follow this general trend (Fig. 3(a,b)), both decreasing at intermediate x_{Li} values as E_{nn} increases. Superimposed on this general trend, at moderate E_{nn} values, both correlation factors sharply decrease at $x_{\text{Li}} = 6$, i.e. two-thirds occupancy. Because f and f_{I} do not change uniformly as E_{nn} is increased, the Haven ratio H_{R} develops a more structure. Perhaps most relevant to the lithium-stuffed garnets, above $x_{\text{Li}} = 6$, the introduction of nearest-neighbour repulsion reduces H_{R} even further from the already low non-interacting value.

The garnet lattice contains octahedral and tetrahedral sites in a 2:1 ratio. In the non-interacting system, the average site occupancies follow this ratio across the full x_{Li} range (Fig. 2(b)). Introducing repulsive nearest-neighbour interactions increases the probability that octahedra are occupied relative to tetrahedra. Because octahedral sites are two-coordinate, versus the four-coordinate tetrahedral sites, preferentially occupying octahedral sites minimises the number of disfavoured nearest-neighbour interactions. This effect is strongest at two-thirds site occupation ($x_{\text{Li}} = 6$) where a sufficiently large E_{nn} drives the system into a fully ordered arrangement with all the octahedral sites filled and all the tetrahedral sites empty. In this fully ordered system, correlation effects are maximised: a single ion hopping from octahedron to tetrahedron is blocked, and must return to its starting position. Diffusion becomes possible

only for groups of particles undertaking highly correlated collective movement.⁴⁵ Both tracer diffusion and ionic conductivity are stringently reduced compared to their corresponding non-interacting system values, as seen in the changes in f and f_{I} . The collective correlations ($f_{\text{I}} < 1$) are also visible in the reduced conductivity, σ' , which decreases relative to the non-interacting system across the full x_{Li} range, with a notably strong decrease at $x_{\text{Li}} = 6$.

2. Asymmetric site-occupation energies

In the non-interacting model, not only do mobile ions not interact with each other (excepting volume exclusion), but there are no relevant interactions between the mobile ions and the host lattice. Identifying a site as octahedral or tetrahedral only has relevance for defining the connectivity of the lattice graph. Mobile ions show an equal preference for octahedral and tetrahedral sites, with average occupations following a simple 2:1 ratio. This behaviour contrasts with experimental observations. Neutron data for lithium-garnets such as $\text{Li}_3\text{Y}_3\text{Te}_2\text{O}_{12}$ reveal that at $x_{\text{Li}} = 3$ the lithium ions exclusively occupy the tetrahedral sites.^{40,69} This suggests that at relatively low lithium concentrations, there is an energetic penalty for occupying octahedral rather than tetrahedral sites.⁶⁸ We model this difference in site-occupation energy by including a term $\Delta E_{\text{site}} = E_{\text{oct}} - E_{\text{tet}}$. To investigate the

effect of this ion–lattice interaction on ion dynamics and site occupations we performed a series of simulations for otherwise non-interacting particles, with $\Delta E_{\text{site}} = 0 - 5kT$.

The effect of ion–lattice interactions qualitatively mirrors the effect of nearest-neighbour interactions (Fig. 4). Both single-particle and collective correlation factors are lower than their non-interacting values, average site occupancies deviate from those in the ideal system, and the reduced ionic conductivity decreases. Here, however, the strongest correlations emerge at $x_{\text{Li}} \approx 3$. As ΔE_{site} increases, tetrahedral sites are preferentially occupied with respect to octahedral sites, in contrast to the opposite behaviour observed when increasing nearest-neighbour repulsion. In the limit $T \rightarrow 0$ this again corresponds to a fully ordered arrangement of ions, now with all the tetrahedral sites filled and all the octahedral sites empty. The Haven ratio, H_R , shows less change compared to the non-interacting result, with only a small decrease for $x_{\text{Li}} < 3$.

3. Combined site inequality and nearest-neighbour repulsion

In real garnet electrolytes, lithium ions can be expected to interact with the host lattice, and also with each other. To explore the behaviour when both nearest-neighbour and site-occupation interactions exist we performed simulations to map the $\{x_{\text{Li}}, \Delta E_{\text{site}}, E_{\text{nn}}\}$ parameter space. The full data from these calculations is presented in Figs. A1–A4. With both interactions present, the ion dynamics and site occupation statistics are more complex, with specific details that depend on the precise values of both interaction terms. The general features, however, are illustrated by considering the subset for which $E_{\text{nn}} = \Delta E_{\text{site}}$ (Fig. 5). The correlation factors, f and f_I , both show sharp decreases at $x_{\text{Li}} = 3$ and $x_{\text{Li}} = 6$, in both cases corresponding to ordered arrangements of Li ions throughout the lattice. As in the previous single-interaction models, the ordering at $x_{\text{Li}} = 3$ corresponds to filled tetrahedra and empty octahedra (due to ΔE_{site}), and the ordering at $x_{\text{Li}} = 6$ corresponds to filled octahedra and empty tetrahedra (due to E_{nn}). The average site occupation switches sharply from pure tetrahedral occupation to pure octahedral occupation in the range $x_{\text{Li}} = 3 \rightarrow 6$. The reduced ionic conductivity, σ' , is depressed most strongly at lithium stoichiometries corresponding to the ordered arrangements of ions, again, mirroring the results for single interactions.

4. Tuning lithium stoichiometry to maximise ionic conductivity

In the context of the lithium-garnet solid electrolytes, a recurring question is the design of specific compositions with high ionic conductivities. For garnets with stoichiometries $\text{Li}_x\text{A}_3\text{B}_2\text{O}_{12}$, the lithium content may be continuously varied by appropriately selecting the

A and B cations, or by substituting Li^+ with small hypervalent cations such as Al^{3+} or Ga^{3+} . Lithium garnets with different lithium stoichiometries can exhibit very different ionic conductivities. For $x_{\text{Li}} = 3$ (e.g. $\text{Li}_3\text{Y}_3\text{Te}_2\text{O}_{12}$) room temperature conductivities are typically too small to measure⁴⁰ while for $x_{\text{Li}} \approx 6.5$ (e.g. $\text{Li}_{6.55}\text{Ga}_{0.15}\text{La}_3\text{Zr}_2\text{O}_{12}$) conductivities as high as 1.3Scm^{-1} have been reported.^{70,71} One approach to optimising the ionic conductivity, therefore, is to try to identify an “optimal” lithium stoichiometry.^{36,50,72–78} Other studies have considered how the conductivity varies with the distribution of lithium ions between tetrahedral and octahedral sites.^{30,36,76,79} As discussed above, this distribution is itself a function of the lithium stoichiometry, and also of the interactions experienced by the lithium ions.

In general, results here emphasise that ordered phases (typically at integer Li stoichiometries / commensurate with site stoichiometries) should be avoided.

From equations(?) (and assuming fixed ion mobilities?) the ionic conductivity varies with carrier mole fraction, x , as

$$\sigma_{\text{dc}} \propto x(1-x)f_I \quad (14)$$

In the non-interacting case this gives parabolic behaviour, with a maximum as $x = 0.5$ ($x_{\text{Li}} = 4.5$).⁵⁹ The interactions present in real systems modify this equation by introducing collective correlations. Interactions (between particles or with the lattice) introduce an additional scaling f_I which describes the efficiency with which diffusive jumps contribute to collective transport.

σ_{dc} is proportional to three terms: the \Rightarrow rewrite this in notation consistent with what appeared earlier The $c(1-c)$ factor gives a parabolic dependence with a maximum ionic conductivity as $x = \frac{9}{2}$, as seen (above?) for the non-interacting / equivalent sites system.

By “switching on” additional physics: on-site energies and interactions between mobile ions, $f_I \neq 1$, with f_I a function of the interaction strengths, the lattice topology, and the carrier concentration. The optimal x_{Li} for a given material therefore depends on the behaviour of $f_I(x_{\text{Li}})$.

The low computational cost of the LGMC simulations have allowed us to calculate a reduced ionic conductivity σ' as a function of $\{x_{\text{Li}}, \Delta E_{\text{site}}, E_{\text{nn}}\}$. Considering this data set, we can pose the following question: what is the value of x_{Li} that maximises the (reduced) ionic conductivity, at each combination of $\Delta E_{\text{site}}, E_{\text{nn}}$

At any combination of $\Delta E_{\text{site}}, E_{\text{nn}}$ we can identify the x_{Li} that gives the maximum ionic conductivity, and then map how $\arg \max \sigma'(x_{\text{Li}})$ depends on the scale of interactions. This defines a surface in model parameter space, which is shown as a contour plot in Fig. 6. A wide range of “optimal” x_{Li} values are identified. With regard to real materials, the question is then, which regions of parameter space correspond to real materials, and to what degree the simple parameters considered here vary across possible chemistries of the garnet family. For example \Rightarrow discussion of effect of cation size / polarizabilities on

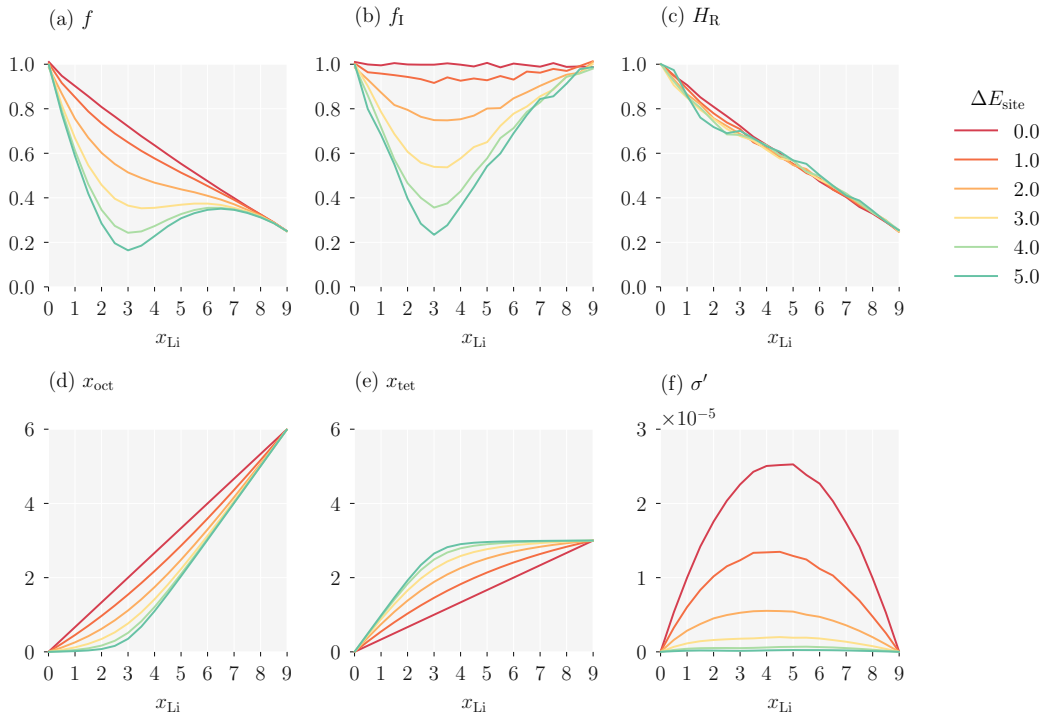


FIG. 4. The effect of unequal site occupation energies for mobile particles on a garnet lattice: (a) single-particle correlation factor, f ; (b) collective correlation factor, f_I ; (c) Haven ratio, H_R ; (d) average octahedra occupation, x_{oct} ; (e) average tetrahedra occupation, x_{tet} ; (f) reduced ionic conductivity, σ' . ΔE_{site} is in multiples of kT .

conductivity trends? Larger cations: larger lattice parameter \rightarrow decreased nearest-neighbour repulsion? Possibly modify balance of ΔE_{sites} ? High polarisability? \rightarrow increased screening? and reduced E_{nn} ? Poses an interesting challenge: to what degree can effective parameters describing a simplified Hamiltonian (such as those employed here) be extracted from e.g. electronic structure calculations on real garnet materials?

note: reported decrease in activation energy with lattice parameter for the series $\text{Li}_5\text{La}_3\text{M}_2\text{O}_{12}$ ($M = \text{Ta}, \text{Nb}, \text{Sb}, \text{Bi}$)⁷⁶[CHECK THIS PAPER]

IV. DISCUSSION

[MOVE THIS COMMENT TO THE INTRODUCTION OR/AND THE DISCUSSION?: SIMULATIONS OF OTHER LATTICE GEOMETRIES WHERE NEAREST-NEIGHBOUR REPULSION HAS BEEN MODELLED PREDICT PARTICLE ORDERING AT SPECIFIC LATTICE OCCUPATION FRACTIONS (DEPENDENT ON THE LATTICE GEOMETRY UNDER CONSIDERATION) AND IS ASSOCIATED WITH STRONG SELF- AND COLLECTIVE-CORRELATION EFFECTS.]

Note. No evidence for a fully ordered $x_{\text{Li}} = 6$ phase with pure octahedral site occupation, suggesting that at typical operating temperatures, the Li–Li interaction is not sufficiently strong to drive complete ordering. Note: Xu *et al.* from DFT predict at $x_{\text{Li}} = 7$ that octahedral sites are 90% occupied, and tetrahedral sites 50% occu-

pied (which material at what temperature?)⁷⁷ (c-LLZO. 0 K? is this a single geometry optimisation?)

[KOZINSKY *et al.* HAVE ALSO PROPOSED A FULLY ORDERED PHASE AT $x_{\text{Li}} = 6$, FROM FIRST-PRINCIPLES CALCULATIONS AND GROUP THEORY ANALYSIS, THAT DIFFERS FROM THE PURE OCTAHEDRAL OCCUPATION DESCRIBED HERE⁸⁰. THE FULLY-ORDERED CONFIGURATION OBSERVED HERE AT $x_{\text{Li}} = 6$ IS DIFFERENT FROM THAT PROPOSED BY KOZINSKY *et al.*⁸⁰. THAT WORK PREDICTED MIXED OCCUPATION OF OCTAHEDRAL AND TETRAHEDRAL SITES. IN THAT CASE CONNECTED TO SYMMETRY BREAKING / IN THIS CASE THE CUBIC LATTICE SYMMETRY IS MAINTAINED]

Summary. What was the motivation (A \rightarrow B). What have we done to explore / solve this problem? What are notable results?

Summary \rightarrow How well does this meet the goals laid out in the introduction?

\Rightarrow To add above: nearest-neighbour interactions and relative site energies are possibly manipulated through the choice of lattice cations, because this determines lattice-site separations, and also the dielectric screening between adjacent Li ions.

Can come back to the point mentioned in the introduction that there is a $\times 10^9$ difference in ionic conductivity between $\text{Li}_3\text{Ln}_3\text{Te}_2\text{O}_{12}$ and $\text{Li}_{6.55}\text{La}_3\text{Zr}_2\text{O}_{12}$, which fits with the ordering behaviour seen here when the model include on-site preference for tetrahedral sites. [CEDER DISCUSSES THE (TYPICAL?) LOWER ENERGY FOR LI IN

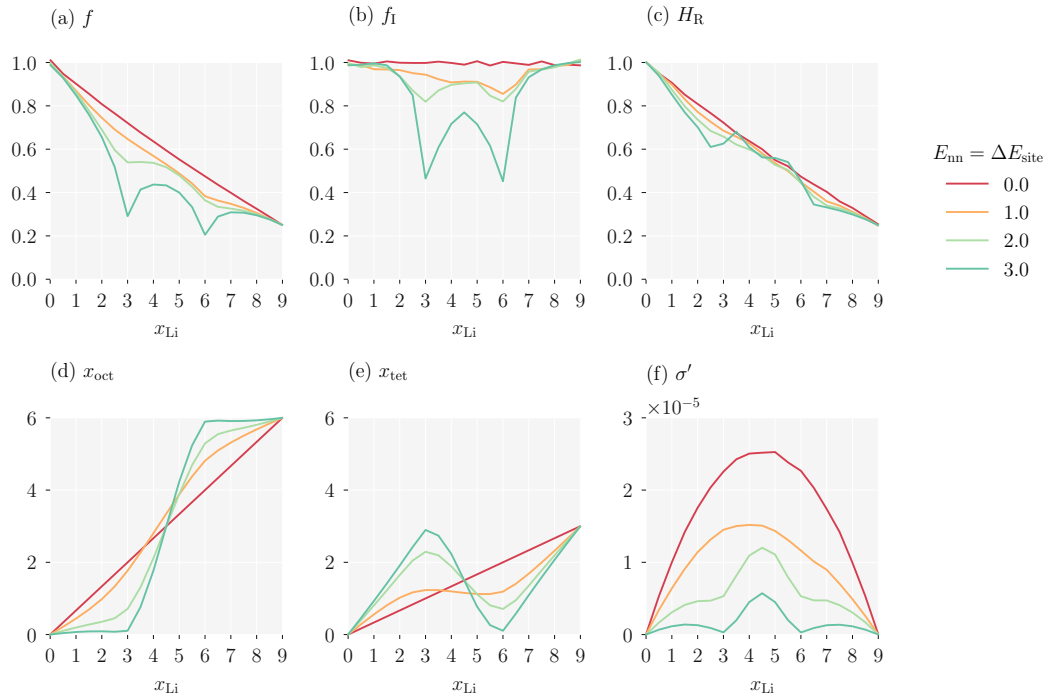


FIG. 5. The effect of combined nearest-neighbour repulsion and site-occupation energy differences on a garnet lattice, for $E_{nn} = \Delta E_{\text{site}}$: (a) single-particle correlation factor, f ; (b) collective correlation factor, f_I ; (c) Haven ratio, H_R ; (d) average octahedra occupation, x_{oct} ; (e) average tetrahedra occupation, x_{tet} ; (f) reduced ionic conductivity, σ' . E_{nn} and ΔE_{site} are in multiples of kT .

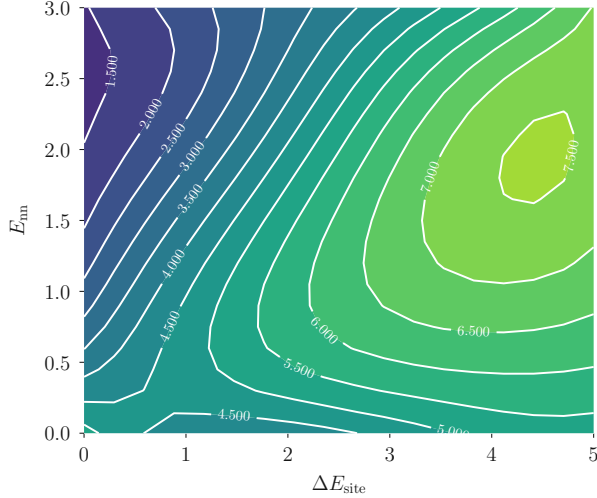


FIG. 6. Contour plot of the value of x_{Li} that maximises the reduced ionic conductivity, σ' , as a function of nearest-neighbour interaction, E_{nn} , and on-site energy difference, ΔE_{site} .

TETRAHEDRAL VS. OCTAHEDRAL SITES IN GENERIC FCC & HCP LATTICES⁶⁸.]

Discuss the benefits / problems with this kind of model Hamiltonian analysis (speed / sampling of parameter space / direct interpretation of simple conceptual interac-

tions) Shortcomings: barrierless kMC (all barriers could be quantified, but is there a straightforward way to map across interactions away from “known” materials? => cite relevant VdV / Ceder papers discussing e.g. variation in barriers with composition)

Other issues: assumption of a perfect lattice? (cf. Kozinsky / t-LLZO) Only single particle jumps (again t-LLZO shown to have many-particle jumps⁴⁵ also Sci. Rep. paper if published?)

The result that the Li preferentially order for specific numbers of occupied sites, and that this is associated with strong single-particle and collective correlation effects is consistent with equivalent results obtained for other lattices for which the effect of alternating site energies have been modelled => e.g. refs and examples? 2D hexagonal lattice

Sharp decrease at $x_{\text{Li}} = 3$ in ionic conductivity associated with ordering of Li across occupied tetrahedral sites, reproduced here by including a site-occupation energy preference for the tetrahedral site, which can be explained as the preferred site at low Li stoichiometries due to the small size of the lithium ion.

=> Discuss: Kozinsky *et al.* have also proposed a fully ordered phase at $x_{\text{Li}} = 6$, from a combined first-principles calculations and group theory analysis⁸⁰, with mixed octahedral and tetrahedral occupation.

Criticism of this approach: Assumes transport is well-described by discrete hops. Not valid for “superionic”

materials, where diffusion is a more liquid-like process: can have concerted motion of groups of atoms. Also possible that for highly ordered (low ionic conductivity) systems, there may be concerted diffusion mechanisms that become favoured over the (slow) individual hopping process modelled here (e.g. the low temperature LLZO tetragonal phase: ordered, and at low temperatures transport is predicted to proceed by a highly-concerted “cyclic” mechanism⁴⁵)

V. SUPPLEMENTARY MATERIAL

Supplementary material for this study is available as a GitHub repository.⁸¹ This repository contains (1) the complete data set used to support the findings of this study, (2) example scripts for running `lattice_mc` simulations on a garnet lattice and collating output data,

and (3) a Jupyter notebook containing the code used to generate Figs. 2–A4. The `lattice_mc` code is available under the MIT license.⁶¹

VI. ACKNOWLEDGEMENTS

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VII. APPENDIX

Note that Murch describes physical correlation effects as “a manifestation of order. For example, an ion on a “right” position in an ordered structure which migrates to a “wrong” position will tend to reverse that jump”.¹⁰

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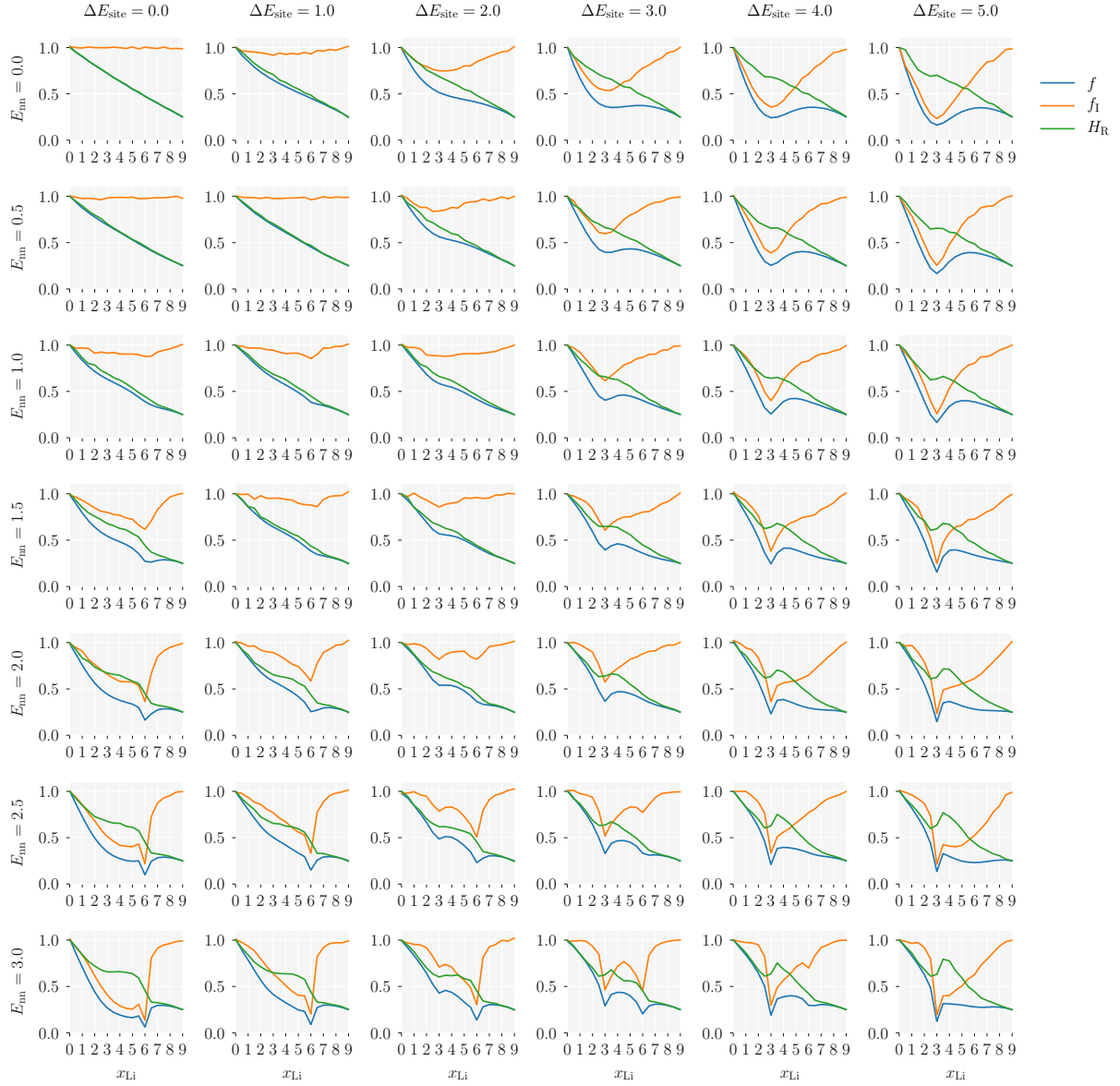


FIG. A1. Correlation factors for the garnet lattice as a function of mobile-ion stoichiometry, x_{Li} , nearest-neighbour repulsion, E_{nn} , and site-occupation energy difference, ΔE_{site} . Each subplot shows the single particle correlation factor, f , the collective correlation factor, f_1 , and the Haven ratio, H_R .

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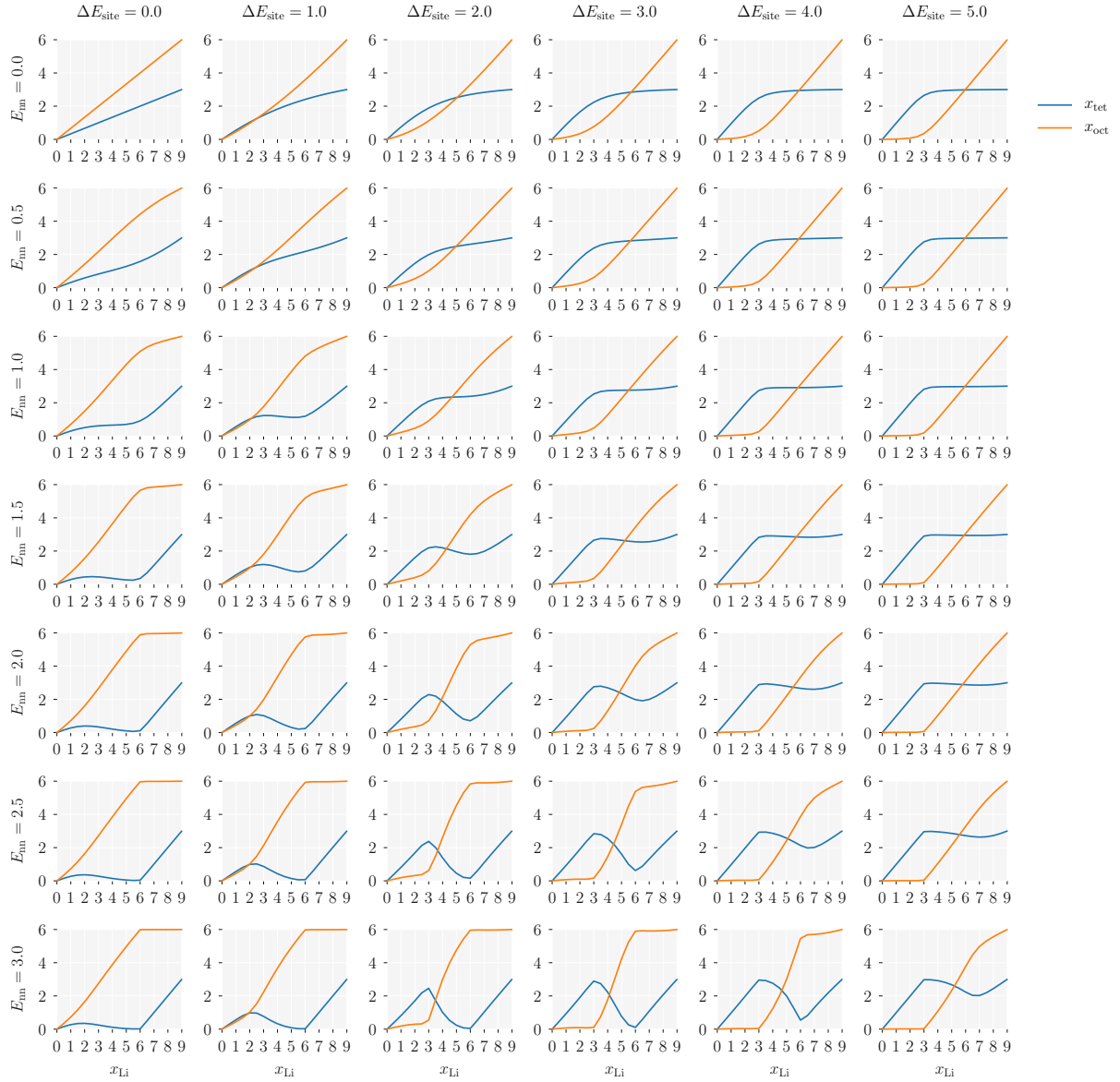


FIG. A2. Average site occupations for the garnet lattice as a function of mobile-ion stoichiometry, x_{Li} , nearest-neighbour repulsion, E_{nn} , and site-occupation energy difference, ΔE_{site} . Each subplot shows the time-averaged number of occupied tetrahedral, x_{tet} , and octahedral, x_{oct} , sites per formula unit.

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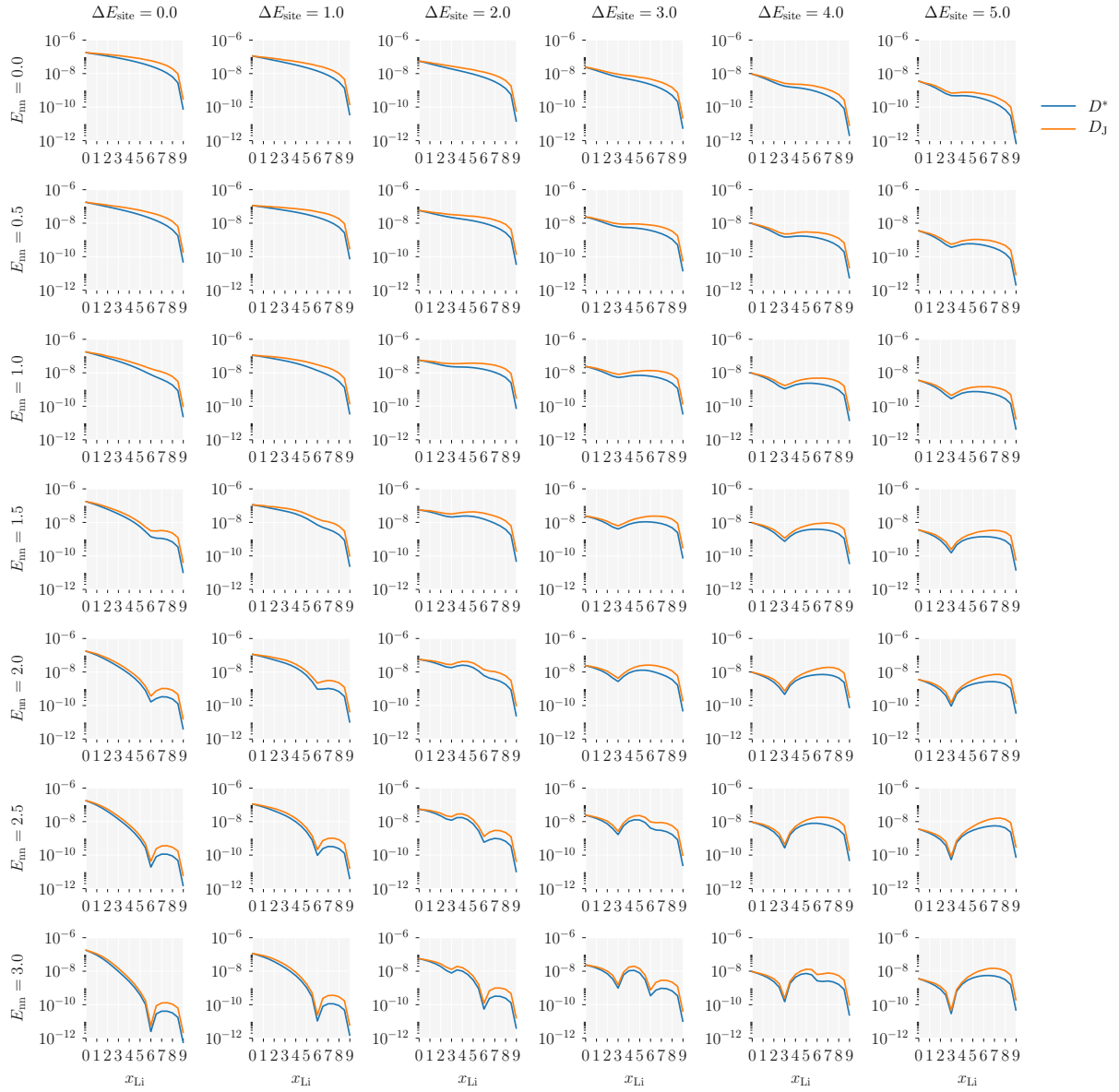


FIG. A3. Diffusion coefficients for the garnet lattice as a function of mobile-ion stoichiometry, x_{Li} , nearest-neighbour repulsion, E_{nn} , and site-occupation energy difference, ΔE_{site} . Each subplot shows the tracer diffusion coefficient, D^* , and the collective “jump” diffusion coefficient, D_J .

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⁶⁵ Because the lattice-gas model used here considers hops as barrierless, where hopping probabilities only depend on energy differences between initial and final states, the ef-

fective transport coefficients calculated here cannot be directly compared to experimental values. Introducing fixed barrier heights for tet↔oct hops is equivalent to scaling the hopping prefactor ν' , which preserves *relative* differences in the transport coefficients presented here. A more realistic model would need to account for the influence of local site occupations on individual hopping barriers, see e.g. Ref.⁸², and would give quantitative deviations from the trends presented here.

⁶⁶ D_J is related to the ionic conductivity (via Eqns. 5 and 11) and also to the chemical diffusion coefficient, \tilde{D} , via

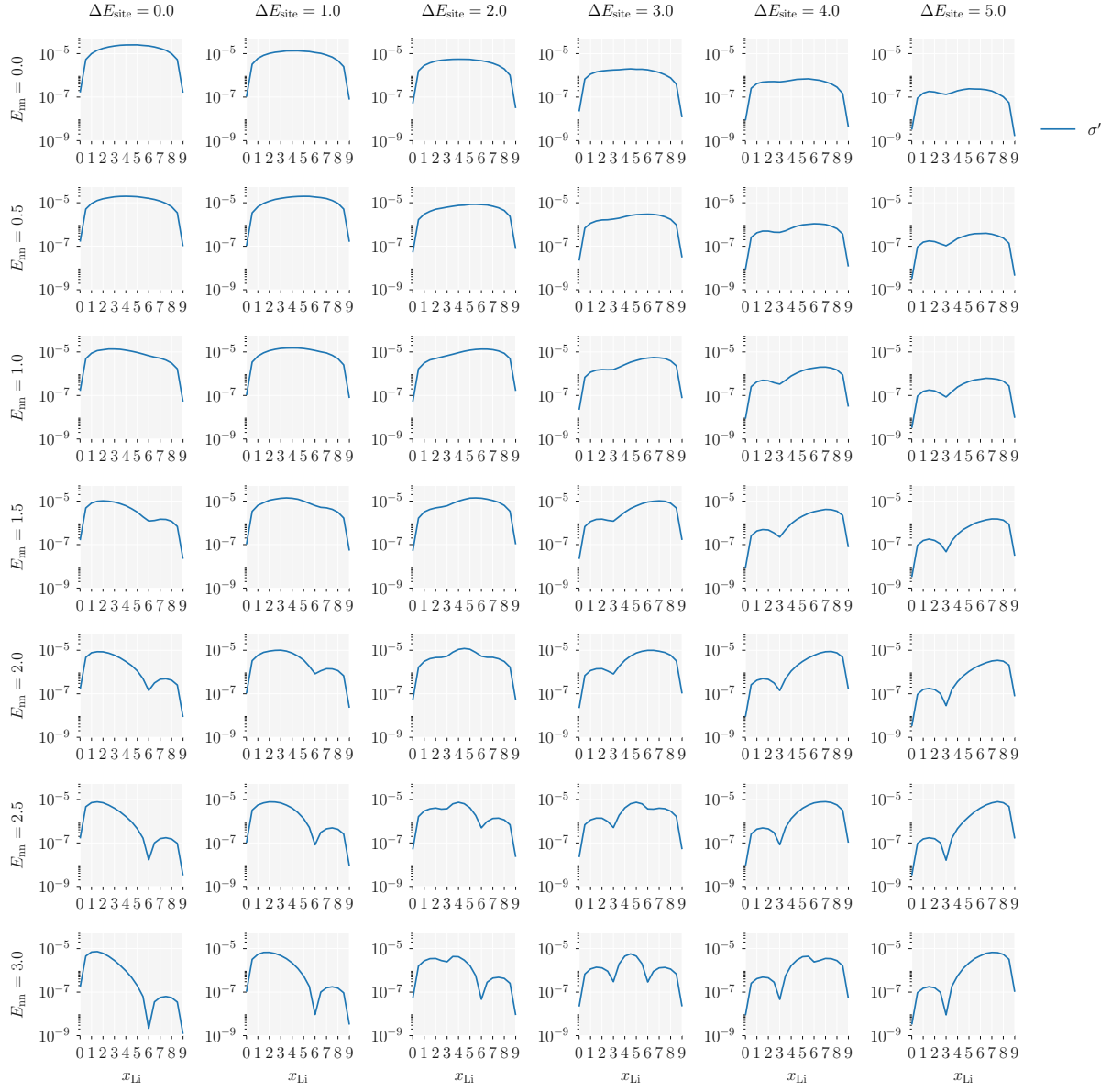


FIG. A4. Reduced ionic conductivity for the garnet lattice as a function of mobile-ion stoichiometry, x_{Li} , nearest-neighbour repulsion, E_{nn} , and site-occupation energy difference, ΔE_{site} . Each subplot shows the reduced ionic conductivity, σ' (Eqn. 13).

the thermodynamic factor, Θ , via $\tilde{D} = D_{\text{J}}\Theta$, where

$$\Theta = \frac{\partial \left(\frac{\mu}{kT} \right)}{\partial \ln x}. \quad (15)$$

⁶⁷ For a system with a single mobile species, the reduced conductivity is equal to the true ionic conductivity if $(VkT)/(q^2) = 1$.

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