A mechanism of defect-enhanced phase transformation kinetics in lithium iron phosphate olivine

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Antisite defects are a type of point defect ubiquitously present in intercalation compounds for energy storage applications. While they are often considered a deleterious feature, here we elucidate a mechanism of antisite defects enhancing lithium intercalation kinetics in LiFePO\textsubscript{4} by accelerating the FePO\textsubscript{4} \textrightharpoonup LiFePO\textsubscript{4} phase transformation. Although Fe\textsubscript{(Li)} antisites block Li movement along the [010] migration channels in LiFePO\textsubscript{4}, phase-field modeling reveals that their ability to enhance Li diffusion in other directions significantly increases the active surface area for Li intercalation in the surface-reaction-limited kinetic regime, which results in order-of-magnitude improvement in the phase transformation rate compared to defect-free particles. Antisite defects also promote a more uniform reaction flux on (010) surface and prevent the formation of current hotspots under galvanostatic (dis)charging conditions. We analyze the scaling relation between the phase boundary speed, Li diffusivity and particle dimensions and derive the criteria for the co-optimization of defect content and particle geometry. A surprising prediction is that (100)-oriented LiFePO\textsubscript{4} plates could potentially deliver better performance than (010)-oriented plates when the Li intercalation process is surface-reaction-limited. Our work suggests tailoring antisite defects as a general strategy to improve the rate performance of phase-changing battery compounds with strong diffusion anisotropy.

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INTRODUCTION

Point, line and/or planar defects are ubiquitously present in all materials and frequently have beneficial effects on material properties. The intentional introduction and control of defects plays a key role in the development of advanced materials with better performance and new functionality. Well-known examples include doping semiconductors to modify the band structure and using phase or grain boundaries to strengthen alloys. Like other materials, battery intercalation compounds contain various types of defects. “Defect engineering” is a promising strategy for this class of materials, which nonetheless has not yet been widely explored. In particular, recent studies find that antisite defects, which are common in battery compounds, can promote Li transport and enhance rate performance by opening up alternative diffusion channels with lowered migration energies in numerous lithium-ion battery electrode materials. Such phenomena have been reported in Li\textsubscript{1.217}Mn\textsubscript{0.467}Cr\textsubscript{0.3}O\textsubscript{2}, Li\textsubscript{1}Mn\textsubscript{1−x}Fe\textsubscript{x}PO\textsubscript{4} (ref. 10) and Li\textsubscript{4}Ti\textsubscript{5}O\textsubscript{12} (ref. 11) etc. In refs. 10-12, antisite defects are also reported to improve the stability and cyclability of cubic Li\textsubscript{4}Ti\textsubscript{5}O\textsubscript{12}, where the random mixing of Li and Ti on octahedral sites in the cubic phase electrochemically induced from amorphous TiO\textsubscript{2} enables reversible capacity that cannot be achieved otherwise. These examples demonstrate that the rational tailoring of antisite defects provides a potentially general approach to improving battery electrode properties.

Here we present a computational study that reveals a new mechanism of antisite defects enhancing the rate capability of intercalation compounds by accelerating surface-reaction-limited (SRL) phase transformation during battery charge/discharge. Antisite defects are generated when the sites of intercalating ions are occupied by other cations that are usually less mobile. A prominent effect of antisite defects in battery electrodes is to block the existing paths of intercalating ions and generate new migration pathways at the same time. For intercalation compounds with strong anisotropic transport properties, which are common among battery materials, such effect usually leads to the reduction in the diffusion anisotropy of intercalating ions. For example, lithium iron phosphate olivine (LiFePO\textsubscript{4}) is theoretically predicted to have predominantly one-dimensional (1D) Li diffusion in the [010] direction.\textsuperscript{7} However, experiments show that practically synthesized LiFePO\textsubscript{4} with just a few percent of Li–Fe antisite defects (Li/Fe) instead exhibit two-dimensional (2D) Li diffusivity.\textsuperscript{6,9} This discrepancy is explained by first-principles calculation\textsuperscript{10} that finds Li/Fe to impede Li movement along [010] migration channels but facilitate Li hopping between the channels via vacancies on Fe sites created by antisite defects. Similar observations of antisite defects reducing ion diffusion anisotropy are also reported for Li\textsubscript{1}Mn\textsubscript{1−x}Fe\textsubscript{x}O\textsubscript{2} (ref. 10), Li\textsubscript{4}Ti\textsubscript{5}O\textsubscript{12} (ref. 11), Na\textsubscript{2+x}Fe\textsubscript{2−x}S\textsubscript{2} (SO\textsubscript{4})\textsubscript{3} (ref. 11) and LiMnBO\textsubscript{3} (ref. 12).

In this work, we show that antisite defects in LiFePO\textsubscript{4} lead to an unexpected increase in the SRL phase transition rate. This is achieved by an increase in the surface reaction area for Li intercalation due to defect-enhanced Li diffusion along [100] even though antisites impede Li movement in the fast [010] diffusion direction. Analysis of the interplay between surface reaction and Li diffusion reveals that the inclusion of antisite defects in LiFePO\textsubscript{4} qualitatively changes the particle size dependence of phase transformation rate in the SRL regime. As a result, the rate performance of defect-containing LiFePO\textsubscript{4} is less sensitive to particle dimensions, which facilitates the use of larger particles to improve the packing density and reduce side reaction of the electrodes. Under galvanostatic conditions, the presence of antisites also reduces the risk of electrode damage from current hotspots by distributing the reaction flux more uniformly on particle surface. Due to the kinetic competition between surface reaction and bulk diffusion, an optimal defect concentration is
predicted to exist for a given particle geometry to maximize the Li (de)intercalation rate. Criteria for the co-design of defect content and particle morphology are proposed. Counterintuitively, we find that (100)-oriented LiFePO4 platelike particles may exhibit even better rate performance than (010)-oriented plate particles in the SRL kinetic regime at relatively low defect levels, whereas the latter is commonly viewed as the most desirable LiFePO4 particle morphology. While we demonstrate the possibility of using antisite defects to accelerate phase transitions in LiFePO4, it may have general applicability to other phase-changing battery materials that exhibit ion diffusion anisotropy.

RESULTS

Electrochemically driven first-order phase transformations in battery-electrode materials upon ion (de)intercalation are subject to the kinetic control of various rate-limiting steps. We previously show9 that the competition between Li diffusion and surface reaction can give rise to three distinct phase transformation modes in LiFePO4, i.e. bulk-diffusion-limited (BDL), SRL and an intermediate hybrid mode, in which phase boundary migration is BDL or SRL in different directions. As illustrated in Fig. 1, these transformation modes are associated with different phase growth morphologies and rates. They are not unique to LiFePO4 and can operate in other intercalation compounds. Phase transformation becomes SRL when Li insertion/extraction at the electrode particle surface is much more sluggish than its diffusion inside the particles. This could occur at small particle sizes, low applied over/under-potentials and/or low exchange current density for charge transfer, etc.9 In a pioneering study, Singh, Ceder and Bazant (SCB) investigated the SRL phase transformation kinetics using a depth-averaged model of LiFePO4.13 They show that a salient feature of the SRL mode is that phase boundary travels at a constant velocity perpendicular to the main Li intercalation direction (Fig. 1), which is in sharp contrast to the behavior of the BDL mode. Although the SCB theory assumes 1D Li transport in LiFePO4 in accordance with the DFT prediction,7 the characteristics of the SRL mode are independent of Li diffusion anisotropy and also apply to cases with higher dimensional Li diffusivity in the presence of antisite defects.9

Antisite defects accelerate SRL phase transformation

We study the SRL phase transformation process in defect-containing LiFePO4 particles using the mesoscale phase-field simulation method.14-16 which has been applied to the modeling of LiFePO4.17-21 An unexpected phenomenon that antisite defects can increase the phase boundary migration speed is observed from the simulations, in which the influence antisite defects exert on the discharge process is captured by their effect on the anisotropic Li diffusivity in LiFePO4. In ref. 15, Malik et al. evaluate from first-principles calculation the activation barrier required for Li$^+$ to cross over to a neighbor channel through an antisite in LiFePO4. The calculated Li migration barrier is employed to determine the defect concentration dependence of Li diffusivity along [010], [100] and [001] axes, $D_{[010]}$, $D_{[100]}$ and $D_{[001]}$ at 440 K. While $D_{[010]}$ at the room temperature (300 K) is also reported at several defect levels in ref. 15, the values of $D_{[100]}$ and $D_{[001]}$ at 300 K are not provided in that work. As they are required for our simulations, we calculated all the three diffusion coefficients at 300 K as a function of antisite concentration following the approach of Malik et al.10 (see Methods). As shown in Supplementary Fig. 1 in the Supplementary Information (SI), $D_{[100]}$ exhibits an approximately inverse relationship with $D_{[010]}$ at 300 K. Both of them approach $10^{-12}$ to $10^{-11}$ cm$^2$ s$^{-1}$ at high defect concentrations, consistent with our previous observation of 2D room-temperature Li diffusivity $D_{[010]}\approx D_{[100]}\approx 10^{-11}$ cm$^2$ s$^{-1}$ in defect-containing LiFePO4 microparticles.3 The calculated Li diffusivity values at 300 K are used in the phase-field simulations presented below.

Figure 2 compares the discharge simulations for two LiFePO4 particles with different antisite contents at a constant underpotential $\Delta\phi = 35$ mV, which is below the underpotential required to form a metastable solid solution in LiFePO4. The particles are given (010)-oriented platelike shape in simulations to facilitate fast Li diffusion along the [010] direction and ensure that the phase transition is in the SRL regime. Such particle morphology is

![Image](https://example.com/image.png)

**Fig. 1** Schematics of three distinct kinetic modes of phase transformation in LiFePO4.
commonly obtained in hydrothermal synthesis. In the simulations, Li is inserted into the particle from (010) surface only, and zero-flux boundary condition is applied to (100) surfaces. The “defect-lean” particle contains 0.5% antisites and has a large \( D_{(010)} = 1.3 \times 10^{-10} \text{ cm}^2 \text{s}^{-1} \) but a much smaller \( D_{(100)} = 3.9 \times 10^{-14} \text{ cm}^2 \text{s}^{-1} \). The “defect-rich” particle contains 25% antisites and has a much reduced Li diffusion anisotropy with \( D_{(010)} = 5.4 \times 10^{-12} \text{ cm}^2 \text{s}^{-1} \) and \( D_{(100)} = 2.6 \times 10^{-12} \text{ cm}^2 \text{s}^{-1} \). Because phase boundaries are observed to be parallel to the [001] axis in platelike particles, \( D_{(010)} \) simulations were reduced to two dimensions to improve computation efficiency. Coherency stress that arises from the lattice mismatch between LiFePO 4 (LFP) and FePO 4 (FP) phases plays a critical role in inducing the metastable solid reaction region on the defect-rich particle. As such, more Li ions are inserted into the defect-rich particle per unit time, which results in a direct increase of the enhanced [001] diffusivity enabled by defects. It can be seen that the Li diffusion flux in the defect-rich particle is confined near phase boundary. However, significant Li diffusion flux along [100] exists within the entire defect-rich particle, which transports Li atoms inserted at surface locations distant from the phase boundary to the boundary to participate in phase transformation. This results in a larger surface reaction area. Therefore, a key insight we obtained is that an increased Li diffusivity along the phase boundary migration direction ([100] here) benefits the SRL phase transformation kinetics in LiFePO 4 by expanding the surface reaction region. This differs from the effect of faster Li diffusion on improving phase transformation kinetics in the BDL regime, in which a higher diffusivity enhances phase boundary velocity in the same direction. It represents a new mechanism of accelerating phase transitions through the interplay between Li diffusion and surface reaction, which has not been explored so far.

To further shed light on the effect of Li diffusion on SRL kinetics, especially the quantitative dependence of phase boundary velocity on [001] and [010] Li diffusivity, a series of particle discharge simulations under constant \( \Delta \phi = 35 \text{ mV} \) were carried out. Instead of obeying the relation given by Supplementary Fig. 1, the values of \( D_{(010)} \) and \( D_{(100)} \) are independently varied in simulations in order to study their respective effects on the phase transition kinetics. The [010] thickness of the particle is \( L_{(010)} = 50 \text{ nm} \), and its [100] dimension \( L_{(100)} \) is assumed to be much larger than the surface reaction region so that the phase boundary velocity \( V_{PB} \) is independent of \( L_{(100)} \). As shown in Fig. 4, varying \( D_{(010)} \) from 10^{-9} to 10^{-12} \text{ cm}^2 \text{s}^{-1} at a given \( D_{(100)} \) has little effect on the calculated phase boundary velocity \( V_{PB} \) and the surface reaction zone width \( W \), which is defined as the width of the region where \( j_s \) is larger than 35% of its peak value. This confirms that phase transition is indeed in the SRL regime and hence insensitive to \( D_{(010)} \). We note that \( D_{(010)} \) in defect-containing LiFePO 4 is predicted to increase beyond the bulk diffusivity value at very small [010] particle thickness when antisites can no longer block the 1D migration channels. The results above suggest that this particle size dependence of \( D_{(010)} \) does not have a large impact on the predicted acceleration of phase transition by antisites in the SRL regime. On the other hand, increasing \( D_{(100)} \) from 10^{-14} to 10^{-11} \text{ cm}^2 \text{s}^{-1} while keeping \( D_{(010)} \) constant causes both \( W \) and \( V_{PB} \) to increase by more than 20 times.

**Depth-averaged model**

We found that the 2D simulation results can be well approximated by a 1D depth-averaged model similar to the SCB theory, which is derived by assuming that Li concentration \( c \) is uniform along [010]...
the intrinsic diffuse interface width in the phase-V

Dependence of phase boundary velocity to enable much better rate capability in

Fig. 4  Dependence of phase boundary velocity $V_{PB}$ and surface reaction zone width $W$ on $D_{[100]}$ and $D_{[101]}$. Markers represent 2D phase-field simulation results. Solid and dashed lines are numerical and approximate analytical solutions to the depth-averaged model, respectively. Dashed arrow indicates the $V_{PB}$ value predicted by the SCB model, i.e. for the case of $D_{[100]} = 0$.

and a function of [100]-coordinate $x$ only. With the simplification, the governing equation of the phase-field model becomes

$$\frac{\partial c}{\partial t} = \frac{\partial}{\partial x} \left[ D_{[100]} V_{m} \varepsilon (1 - \varepsilon) \frac{\partial \mu_i}{\partial x} \right] + \frac{2J_i}{L_{[010]}}$$  \(1\)

where the surface reaction flux $J_i$ on one (010) facet is given by the Butler-Volmer equation (Methods). Equation 1 has the form of a reaction-diffusion equation. It reduces to the SCB model when the diffusion term disappears from Eq. 1 at $D_{[100]} = 0$. Phase boundary velocity $V_{PB}$ and surface reaction zone width $W$ predicted by this model are shown as solid lines in Fig. 4. The excellent agreement with the full simulations allows one to use the depth-averaged model to efficiently analyze the SRL kinetics in the presence of [100] Li diffusion.

Scaling relation

The simulation results in Fig. 4 show that both $V_{PB}$ and $W$ hold the same parabolic relation with $D_{[100]}$, i.e. $V_{PB} \propto D_{[100]}^{1/2}$ and $W \propto D_{[100]}^{1/2}$ when $D_{[100]} > \sim 10^{-14} \text{ cm}^2 \text{ s}^{-1}$. Furthermore, $V_{PB}$ is proportional to $W$ over the entire range of $D_{[100]}$. When $D_{[100]} \rightarrow 0$, $W$ approaches the intrinsic diffuse interface width in the phase-field model, and $V_{PB}$ reaches the prediction of the SCB model. It shows that a defect-free LFP particle has a very low SRL phase boundary velocity ($<10^{-3} \text{ nm s}^{-1}$) because of a very narrow surface reaction zone (~2 nm). As a result, good rate performance can only be achieved when the FP $\rightarrow$ LFP first-order phase transition is bypassed. However, when an adequate amount of antisite defects are present in the particle, the surface reaction area can increase by orders of magnitude and results in a similar increase in the phase boundary velocity to enable much better rate capability in the absence of metastable solid solution.

A more transparent understanding of the scaling relation can be obtained from an approximate analytical solution to the depth-averaged model, which is based on the sharp-interface assumption (see derivation in Methods). The solution gives the following expressions of surface reaction zone width $W$ and phase boundary velocity $V_{PB}$:

$$W = \lambda_1 \sqrt{\frac{D_{[100]} L_{[010]}}{I_0}}$$  \(2\)

$$V_{PB} = \lambda_2 \sqrt{\frac{I_0 D_{[100]} L_{[010]}}{I_0}}$$  \(3\)

where the expressions of $\lambda_1$ and $\lambda_2$, which are functions of $\Delta \phi$, are given in Methods, and $I_0$ is the exchange current density. As shown in Fig. 4, Eqs. 2 and 3 show excellent agreement with the numerical solution when $W$ is significantly larger than the diffuse-interface width of the phase boundary. In addition to explaining the parabolic dependence of $W$ and $V_{PB}$ on $D_{[100]}$, Eq. 3 shows that $V_{PB}$ varies with $I_0$ and $L_{[010]}$ particle thickness $L_{[010]}$ as $I_0^{1/2}$ and $L_{[010]}^{1/2}$, respectively. Notably, these relations qualitatively differ from defect-free LFP particles with 1D Li diffusivity, in which $V_{PB}$ has a stronger dependence on $I_0$ and $L_{[010]}$, i.e. $V_{PB} \propto I_0$ and $V_{PB} \propto L_{[010]}$, according to the SCB theory. Such difference in the scaling behavior lies in that upon decreasing $I_0$ or increasing $L_{[010]}$, the surface reaction area $W$ increases in defect-containing particles (Eq. 2), which contributes to a less pronounced reduction of $V_{PB}$.

Particle size dependence of phase transformation time

The phase boundary speed predicted by Eq. 3 applies to the situation where the [100] particle size is larger than the surface reaction zone width $W$. When $L_{[100]} < W$, the entire (010) particle surface is active for Li intercalation during discharge. An interesting prediction thus arises: $V_{PB}$ should increase approximately linearly with $L_{[100]}$ in the SRL regime, i.e. the longer the travel distance of the phase boundary, the faster it moves. Accordingly, the particle transformation time, given by $t_{f} = L_{[100]} / V_{PB}$, should be insensitive to $L_{[100]}$. This prediction is confirmed by calculations shown in Fig. 5, which plots the average $V_{PB}$ and $t_{f}$ as a function of $L_{[100]}$ for Li intercalation into a defect-rich (25% antisites) particle at $\Delta \phi = 35 \text{ mV}$. It can be seen that $V_{PB}$ is proportional to $L_{[100]}$ and $t_{f}$ increases very slowly with $L_{[100]}$ up to $\sim 1 \mu \text{m}$, which makes $t_{f}$ only dependent on [010] particle size as $t_{f} \propto \sqrt{L_{[010]}}$. In contrast, $t_{f}$ is proportional to $L_{[100]}^{2} L_{[010]}$ in defect-free particles. Along with Eq. 3, this comparison shows that the inclusion of antisite defects results in qualitatively different particle size dependence of the SRL phase transition kinetics. The weaker dependence of $t_{f}$ on both $L_{[100]}$ and $L_{[010]}$ in defect-containing particles implies another benefit of antisite defects, i.e. they enable the rate performance of LiFePO$_4$ to degrade less severely with particle dimensions. This facilitates the use of larger particles in

Fig. 5  Effect of $L_{[100]}$ on the average phase boundary velocity $V_{PB}$ (red line) and total transformation time $t_{f}$ (black line), for Li intercalation into a particle with $D_{[100]} = 2.6 \times 10^{-12} \text{ cm}^2 \text{ s}^{-1}$ at $\Delta \phi = 35 \text{ mV}$. The [010] particle size is 50 nm.
applications, which can improve the packing density and reduce side reactions between particles and electrolyte in electrodes.

Galvanostatic cycling behavior

Besides constant underpotential condition, we also studied the difference between defect-rich and defect-lean particles under galvanostatic discharge or constant current conditions. Figure 6a compares the distributions of Li intercalation flux \( j_i \) on the (010) surface of particles with different defect contents when galvanostatically discharged to 50% state of charge at 0.005 C (nC = fully discharged in 1/n hours). While the total amount of Li intercalated into the particles per unit time is the same, the peak value of \( j_i \) decreases sharply with \( D_{[100]} \) because faster \([100]\) diffusion causes the reaction flux to be more evenly distributed on (010) surface. A 123-fold reduction in the peak flux occurs when \( D_{[100]} \) increases from 0 cm² s⁻¹ (0% antisites) to 2.6 × 10⁻¹² cm² s⁻¹ (25% antisites). Meanwhile, increasing \( D_{[100]} \) also significantly decreases the underpotential required from 30 mV to 0.3 mV as shown in Fig. 6b. Therefore, antisite defects are beneficial under galvanostatic cycling conditions by reducing polarization and mitigating degradation caused by current hotspots and electrochemical shock.²⁸⁻³⁰

Optimization of defect concentration and particle morphology

After demonstrating the benefits of antisite defects on phase transformation kinetics in the SRL regime, we ask the question whether there exists an optimal defect concentration and how it depends on the LiFePO₄ particle geometry. We still consider the plate-like particle morphology here for its practical relevance. The SRL transformation rate is maximized when the entire plate surface is active for Li intercalation during (dis)charge, which requires \( L_{[100]} < W \). Using Eq. 2, this leads a criterion on \([100]\) Li diffusivity

\[
D_{[100]} > \frac{L_{[100]}^2}{2} \frac{L_{[010]}}{\Delta \phi} \frac{V_{PB}}{W} \tag{4}
\]

On the other hand, Li bulk diffusion along the [010] plate thickness direction should be sufficiently facile so that it does not limit the transformation kinetics. This condition can be described by the inequality \( L_{[010]}^2 / D_{[010]} < W / V_{PB} \), where \( L_{[010]}^2 / D_{[010]} \) is the characteristic Li diffusion time along the [010] axis and \( W / V_{PB} \) is the time a [010] channel stays active for Li intercalation. Applying Eqs. 2 and 3, we derive a criterion on [010] Li diffusivity

\[
D_{[010]} > \frac{L_{[010]}^2}{2} \frac{L_{[010]}}{\Delta \phi} \frac{V_{PB}}{W} \tag{5}
\]

Equations 4 and 5 provide the guidance on tuning the defect concentration for given particle sizes, or conversely, the particle geometry for a given defect content.

Because increasing antisite defect concentration has opposite effects on \( D_{[100]} \) and \( D_{[010]} \), Eqs. 4 and 5 may not always be satisfied simultaneously and an optimal defect level may exist. We numerically examine the defect concentration dependence of SRL transformation rate in 2D phase-field simulations, in which a (010)-oriented LiFePO₄ particle of \( L_{[100]} \times L_{[010]} = 400 \text{ nm} \times 50 \text{ nm} \) is discharged at \( \Delta \phi = 35 \text{ mV} \). Figure 7 shows that the lithium intercalation time keeps decreasing with antisite concentration up

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**Fig. 6** Effect of \( D_{[100]} \) on SRL phase transformation behavior in a particle of size \( L_{[100]} \times L_{[010]} = 400 \text{ nm} \times 50 \text{ nm} \) galvanostatically discharged at 0.005 C. a Li intercalation flux distribution \( j_i \) in logarithmic scale at 50% state of charge in defect-rich, defect-lean and defect-free particles, in which \( D_{[100]} = 2.6 \times 10^{-12}, 3.9 \times 10^{-14} \) and \( 0 \text{ cm}^2 \text{s}^{-1} \), respectively. b Dependence of the underpotential applied to the particle on \( D_{[100]} \).

**Fig. 7** Optimization of defect concentration and particle morphology for SRL phase transformation kinetics. The total transformation time \( t_f \) is plotted as a function of \( D_{[100]} \) or \( D_{[010]} \) for (010)-oriented \( L_{[100]} \times L_{[010]} = 400 \text{ nm} \times 50 \text{ nm} \) (solid line) and (100)-oriented \( L_{[100]} \times L_{[010]} = 50 \times 400 \text{ nm} \) (dashed line) platelet particles discharged at \( \Delta \phi = 35 \text{ mV} \).
ments and modeling studies establish the formation known for its exceptional rate capability. Recent experiments on LiFePO₄ surface in contact with electrolyte is conservatively estimated to be ~10⁻¹² cm² s⁻¹, which leads to an effective in-plane diffusivity of ~10⁻¹⁴ cm² s⁻¹ in 150nm-thick, (010)-oriented LiFePO₄ plates. Compared to D_{[100]} = 10⁻¹² cm² s⁻¹ in defect-rich particles, this represents a relatively small contribution to [100] Li transport although its importance will certainly increase with decreasing particle size, which warrants further study. Conversely, like Li surface diffusion antisite defects can also promote phase separation at high underpotentials where a metastable solid solution can form (Δϕ > 45 mV in our model). In a phase-field modeling study, Dargaville and Farrell show that increasing [100] Li diffusivity induces phase separation in LiFePO₄ under high discharge currents, which otherwise favor the solid-solution intercalation behavior. Antisite defects may have a beneficial effect on SRL intercalation kinetics even in this regime. For instance, Supplementary Fig. 4a compares the lithiation rate of defect-rich (25% antisites) vs defect-less (0.5% antisites) particles under Δϕ = 50 mV. It shows that the former is fully lithiated at ~8000 s, when the latter only reaches ~60% lithiation. The reason for such difference lies in that defect-enhanced [100] Li diffusion causes phase separation to initiate in LiFePO₄ metastable solid solution at an earlier time of ~5700 s (see Supplementary Fig. 4b), after which the Li chemical potential at the particle surface is reduced to the LFP/FP two-phase equilibrium level. As shown in Supplementary Fig. 4b, this in turn increases the surface reaction overpotential in the Butler–Volmer kinetics to generate a larger Li intercalation flux. A detailed analysis of the interplay between antisites, phase separation and Li intercalation kinetics will be presented elsewhere.

Finally, we discuss potential experimental strategy to examine the predicted beneficial effects of antisite defects in LiFePO₄. We suggest that our predictions could be tested experimentally by annealing hydrothermally synthesized LiFePO₄ plate-like particles at different temperatures to vary the defect level. In general, increasing the annealing temperature reduces the amount of defects in electrode materials. For instance, this approach has been demonstrated by Chen and Graetz, who report that the FeLi antisite concentration in LiFePO₄ prepared by a hydrothermal method decreases from 8% to 0% when the post-synthesis annealing temperature is increased from 440 °C to 500 °C. In ref. 2, Kim et al. synthesized Li₄MnP₂O₈ with ~0%, 20% and 30% Li/Mn antisites by calcinating samples at 700 °C, 650 °C and 600 °C, respectively, which are tested to confirm the effect of antisite defects on promoting Li diffusion in this material.

DISCUSSION

In summary, we reveal that the inclusion of antisite defects in LiFePO₄ particles could result in orders-of-magnitude improvement in SRL phase transformation kinetics by increasing the active surface area for Li intercalation. Such phenomenon originates from the experimentally confirmed effect of antisite defects on reducing the Li diffusion anisotropy and is expected to be applicable to other phase-changing intercalation compounds with anisotropic ion transport properties. We numerically and analytically study the interplay between Li diffusion and surface reaction in the SRL regime. The results show that antisite defects qualitatively change the scaling dependence of the phase transformation rate on Li diffusivity, exchange current density and particle size. Upon potentiostatic discharge, the Li intercalation rate deteriorates more slowly with increasing particle dimensions in defect-rich particles, which facilitates the use of larger electrode particles without severely compromising rate to increase the surface reaction area and accelerate SRL phase transformation in the two-phase coexistence regime, where phase separation is fully developed. This phenomenon can be analyzed within the same theoretical framework of this work by considering the contribution of surface diffusion to D_{[100]} in Eq. 1. Li diffusivity on (010) LiFePO₄ surface in contact with electrolyte is conservatively estimated to be ~10⁻¹² cm² s⁻¹, which leads to an effective in-plane diffusivity of ~10⁻¹⁴ cm² s⁻¹ in 150nm-thick, (010)-oriented LiFePO₄ plates. Compared to D_{[100]} = 10⁻¹² cm² s⁻¹ in defect-rich particles, this represents a relatively small contribution to [100] Li transport although its importance will certainly increase with decreasing particle size, which warrants further study. Conversely, like Li surface diffusion antisite defects can also promote phase separation at high underpotentials where a metastable solid solution can form (Δϕ > 45 mV in our model). In a phase-field modeling study, Dargaville and Farrell show that increasing [100] Li diffusivity induces phase separation in LiFePO₄ under high discharge currents, which otherwise favor the solid-solution intercalation behavior. Antisite defects may have a beneficial effect on SRL intercalation kinetics even in this regime. For instance, Supplementary Fig. 4a compares the lithiation rate of defect-rich (25% antisites) vs defect-less (0.5% antisites) particles under Δϕ = 50 mV. It shows that the former is fully lithiated at ~8000 s, when the latter only reaches ~60% lithiation. The reason for such difference lies in that defect-enhanced [100] Li diffusion causes phase separation to initiate in LiFePO₄ metastable solid solution at an earlier time of ~5700 s (see Supplementary Fig. 4b), after which the Li chemical potential at the particle surface is reduced to the LFP/FP two-phase equilibrium level. As shown in Supplementary Fig. 4b, this in turn increases the surface reaction overpotential in the Butler–Volmer kinetics to generate a larger Li intercalation flux. A detailed analysis of the interplay between antisites, phase separation and Li intercalation kinetics will be presented elsewhere.

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performance. Antisites also induce a more uniform distribution of the reaction flux on particle surface under galvanostatic discharge conditions, which reduces the risk of electrode degradation due to electrochemical shock. We show that the optimal defect concentration that maximizes Li intercalation rate is dependent on particle geometry, and (100)-oriented plate-like LiFePO₄ potentially offers better rate performance than (010)-oriented plates during SRL (dis)charge process. Our work highlights the promise and opportunities of improving battery electrode compounds through intentional defect manipulation guided by a mechanistic understanding.

METHODS

Antisite concentration dependence of Li diffusivity

We follow the approach described in ref. 15 to determine the Li diffusivity as a function of antisite concentration in LiFePO₄ based on Li migration barriers calculated from the DFT calculations. In short, a 1D random walk model is set up to simulate the hopping of a Li ion between two Fe₁ᵦ antisites within a [010] channel, which are separated by a distance d that depends on the antisite concentration p as d = (1 − p⁻¹)λ₀/2 where λ₀ is the lattice constant of LFP along [010]. The Li ion makes fast random jumps within the channel with an activation barrier of 270 meV taken from ref. 7. Whenever Li hops to a site next to an Fe₁ᵦ, it may cross over to the nearest migration channel to circumvent the obstruction of the antisite with an activation energy of 491 meV as calculated by Malik et al. 10. The frequencies of the two types of jumps are given by \( v = \nu \exp(-E_c/k_BT) \), where \( v = 10^{\nu s^{-1}} \) is the attempted frequency and \( E_c \) is the activation energy. With \( \nu \) as input, 500 kinetic Monte Carlo simulations are run to estimate the average time \( t \) it takes for the Li ion to escape the channel blocked by the two antisites. The Li diffusivity is calculated as \( D = L^2/(2\langle t \rangle) \) where \( L \) is the channel length and \( \langle t \rangle \) is the time spent in the channel. Our work verifies this by reproducing the Li diffusivity values at 440 K and 500 K at 300 K reported in ref. 11 and then used the method to determine Li diffusion for different defect concentrations at 300 K, which is shown in Supplementary Fig. 1.

Phase-field model

We use a previously reported phase-field model 25 to simulate phase transformation in LiFePO₄ upon Li intercalation, which is briefly described here. The site occupancy fraction of Li in LiFePO₄, \( c(t) \), serves as the field variable to distinguish between LFP (c = 1) and FP (c = 0) phases in the model. Li diffusion and the LFP ↔ FP phase transformation are described by the Cahn–Hilliard equation for \( c(t) \) 46,47:

\[
\frac{dc}{dt} = -\nabla \cdot J = -\nabla \left[ \frac{D_v}{RT} (c^2 - c) \right] (1)
\]

where the chemical potential \( \mu_c \) is given by

\[
\mu_c = \frac{\partial \mu_{\text{chem}}}{\partial c} - \kappa \nabla^2 c = \mu_0 + \frac{1}{RT} \left[ \nabla \ln \left( \frac{c}{1-c} \right) + \Omega(1-2c) \right] - \kappa \nabla^2 c \quad (2)
\]

In Eqs. 6 and 7, \( D_v \) is the diffusion coefficient tensor, \( V_m = 43.8 \text{ cm}^3 \text{ mol}^{-1} \) is the molar volume of LiFePO₄ 49. \( R \) is the gas constant and \( \mu_0 \) is the equilibrium Li chemical potential at LFP/FP two-phase coexistence. A regular solution model is used to describe the homogeneous chemical free energy density \( \mu_{\text{chem}} \), of LiFePO₄ with a \( \Omega = 12 \text{ kmol}^{-1} \). The gradient coefficient \( k \) is given a value of 1.68 \text{ x } 10^{12} \text{ J cm}^{-3}, which produces a phase boundary energy of 0.072 J m⁻² that averages the (100), (010) and (001) interface energies obtained from first-principles calculations 50.

As the boundary condition, the Li intercalation flux at particle surface is described by the Butler–Volmer equation:

\[
\dot{j}_s = \frac{i_0}{F} \left[ \exp \left( \frac{\alpha V_m \eta}{RT} \right) - \exp \left( \frac{-\left(1-\alpha \right) V_m \eta}{RT} \right) \right] \quad (3)
\]

where \( \eta = \mu_{\text{Li}} - \mu_{\text{Li}} \) is the surface reaction overpotential with \( \mu_{\text{Li}} \) and \( \mu_{\text{Li}} \) being Li chemical potentials in the surrounding electrolyte and particle surface, respectively. The underpotential \( \Delta \phi \) is related to \( \mu_{\text{Li}} \) as \( \Delta \phi = F \left( \mu_{\text{Li}} - \mu_{\text{Li}} \right) / V_m \). We set \( i_0 = 0.01 \text{ A m}^{-2} \) (ref. 32) and \( \alpha = 0.5 \) in Eq. 8. Supplementary Table 1 in SI summarizes the parameters used in the model and the sources of their values.

1D depth-averaged model

The model is derived in a similar way as in ref. 13. Integrating Eq. 6 along the [010] axis, one has:

\[
\frac{dc}{dt} = \frac{1}{L_{010}} \int_0^{L_{010}} \frac{D_v}{RT} (c^2 - c) \frac{\partial \mu_c}{\partial c} dy + \frac{2j_s}{L_{010}} \quad (4)
\]

where \( c(x,t) = \int_0^{L_{010}} c(x,y,t) dy / L_{010} \) is the average Li concentration in the [010] direction. With the assumption of facile Li transport along [010], so that \( c(t) \) is uniform along [010], i.e. \( c(x,y,t) = c(x,t) \), Eq. 7 becomes:

\[
\mu_c(x,t) = \frac{\partial \mu_{\text{chem}}(c(x,t))}{\partial c} - \kappa \frac{\partial^2 c}{\partial x^2} \quad (5)
\]

Accordingly, the 2D Cahn–Hilliard equation (Eq. 6) is reduced to the depth-averaged equation (Eq. 1), in which the overbar on \( c \) is dropped.

Analytical solution to the depth-averaged model

An analytical expression of the traveling wave solution to the depth-averaged model can be derived with a few approximations. Applying the ansatz \( c(x,t) = c(x-v_Ft) \) to Eq. 1, one obtains:

\[
\frac{dc}{dt} = \frac{D_v}{RT} \left( c^2 - c \right) + \frac{2j_s}{L_{010}} \quad (6)
\]

Letting \( x = 0 \) be the phase boundary location, Eq. 12 is completed by the following boundary conditions:

\[
\begin{align*}
\dot{c}(x=0) & = c_{\text{FP}} - c_{\text{LFP}} \quad (7) \\
\dot{c}(x=\infty) & = c_{\text{FP}} - c_{\text{LFP}} \quad (8) \\
\end{align*}
\]

Function \( L \) is defined as \( L(c,j_s) = \int_{c_{\text{LFP}}}^{c_{\text{FP}}} \left( \frac{1}{2} c_j^2 + \phi(c) \right) dc \) and \( j_s = j_s / \phi(c) \) is the dimensionless reaction flux. Inverting Eq. 14, the solution is explicitly expressed as:

\[
\begin{align*}
\dot{c} & = -2 \sqrt{-\frac{\phi(c)}{2 \Phi(c)}} x < 0 \\
\dot{c} & = 2 \sqrt{-\frac{\phi(c)}{2 \Phi(c)}} x > 0
\end{align*}
\]

Supplementary Fig. 5 shows that the Li concentration profile predicted by Eq. 15 agrees very well with the numerical solution.

As mentioned above, the surface reaction zone is defined as the region in which \( j_s(j_s) > \phi(c_{\text{MAX}}) \), where \( \phi(c_{\text{MAX}}) = j_s(c(0) = 0) \) is the peak reaction flux and \( \alpha \) is given a somewhat arbitrary value of 0.35. Using Eq. 15, the surface reaction zone width \( W \) is given by:

\[
W = \left[ K_{\text{FP}} \left( j_s^{-1}(\phi_{\text{MAX}}) \right) + K_{\text{LFP}} \left( j_s^{-1}(\phi_{\text{MAX}}) \right) \right] \sqrt{\frac{FD_v}{L_{010}} 0 L_{010}} \quad (16)
\]

Calculating the phase boundary velocity \( V_m \) from mass conservation, i.e.:

\[
V_m = 2 \int_{-\infty}^{\infty} \dot{c}(x) dx / (\Delta c L_{010})
\]

one obtains:

\[
V_m = \frac{1}{\Delta c} \int_{-\infty}^{\infty} \dot{c}(x) dx = \frac{1}{\Delta c} \int_{-\infty}^{\infty} \left( \int_{-\infty}^{\infty} \left( j_s(K_{\text{LFP}}(y)) + j_s(K_{\text{FP}}(y)) \right) dy \right) dy \equiv \frac{1}{\Delta c} \int_{-\infty}^{\infty} \dot{c}(x) dx
\]

Figure 4 shows that \( V_m \) and \( V_m \) predicted by Eqs. 16 and 17 have very good agreement with the numerical solutions except at very small \( D_{1000} \).
where $W$ is comparable to the diffuse phase boundary width and the sharp-interface assumption is not valid.

**DATA AVAILABILITY**

The data that support the findings of this study are available from the corresponding author upon request.

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**AUTHOR CONTRIBUTIONS**

M.T. and K.Y. conceived the project. K.Y. and L.H. performed the phase-field simulations. K.Y. and M.T. carried out the theoretical analysis and the kinetic Monte Carlo simulations. M.T., L.H. and K.Y. wrote the paper.
COMPETING INTERESTS
The authors declare no competing interests.

ADDITIONAL INFORMATION
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