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Influence of Inversion on Mg Mobility and Electrochemistry in Spinels

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Supporting Information

ABSTRACT: Magnesium oxide and sulfide spinels have recently attracted interest as cathode and electrolyte materials for energy-dense Mg batteries, but their observed electrochemical performance depends strongly on synthesis conditions. Using first-principles calculations and percolation theory, we explore the extent to which spinel inversion influences Mg^{2+} ionic mobility in $MgMn_2O_4$ as a prototypical cathode, and $MgIn_2S_4$ as a potential solid electrolyte. We find that spinel inversion and the resulting changes of the local cation ordering give rise to both increased and decreased Mg^{2+} migration barriers, along specific migration pathways, in the oxide as well as the sulfide. To quantify the impact of spinel inversion on macroscopic Mg^{2+} transport, we determine the percolation thresholds in both $MgMn_2O_4$ and $MgIn_2S_4$. Furthermore, we



analyze the impact of inversion on the electrochemical properties of the $MgMn_2O_4$ cathode via changes in the phase behavior, average Mg insertion voltages and extractable capacities, at varying degrees of inversion. Our results confirm that inversion is a major performance limiting factor of Mg spinels and that synthesis techniques or compositions that stabilize the well-ordered spinel structure are crucial for the success of Mg spinels in multivalent batteries.

1. INTRODUCTION

Multivalent (MV) batteries, such as those based on Mg^{2+} ,^{1,2} can potentially achieve high volumetric energy density via facile nondendritic stripping/deposition on an energy-dense metal anode.^{3–5} However, the development of viable MV technology is hindered by poor Mg diffusivity in oxide cathodes as well as poor Coulombic efficiencies in liquid electrolytes.^{2,5–7}

One pathway to improve Mg migration in solids is to utilize host structures where Mg occupies an unfavorable coordination environment.^{8–10} Spinels with composition AM_2X_4 (A = Mg, M = metal cations, X = O or S) are appealing structures in this regard because of their tetrahedrally coordinated Mg sites, rather than the preferred octahedral coordination of Mg. Theoretical calculations indeed predict reasonable Mg²⁺ migration barriers (~550–750 meV) in both oxide and sulfide spinels.^{11,12} Note that oxide spinels have long been used as cathodes and anodes in commercial Li-ion batteries.^{13–19}

Spinel- Mn_2O_4 is a particularly promising, energy-dense, MV cathode, as it is one of the few oxides^{20–25} to have shown electrochemically reversible Mg^{2+} intercalation.^{26,27} However, the cyclable Mg content, that is, the observed capacity, seems to depend strongly on the synthesis conditions.^{26–28} Several

studies on the MgMn₂O₄ structure^{29–33} have indicated that the spinel is prone to inversion, that is, Mg/Mn antisite disorder (see Section 2), where the degree of inversion can range from $20\%^{30}$ to $60\%^{.29}$ It has further been argued that the propensity of Mn³⁺ to disproportionate into Mn²⁺ and Mn⁴⁺ promotes spinel inversion and phase transformations.^{16,34} Since inversion directly affects the local cation arrangement, it may significantly impact the Mg²⁺ ionic mobility.^{35,36} For the rational design of improved Mg battery cathodes it is, therefore, crucial to understand how inversion in oxide spinels affects Mg²⁺ migration.

Inversion is not a phenomenon unique to oxides, and other chalcogenide spinels such as sulfides, which are also important cathode materials in MV technology,¹² are also known to exhibit inversion.^{37,38} A recent combined theoretical and experimental study has identified ternary sulfide and selenide spinels as promising Mg-ion conductors with potential applications as solid electrolytes in MV batteries.³⁸ Solid

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Figure 1. Schematic of a (a) normal and an (b) inverted spinel MgM_2X_4 (M = Mn, In and X = O, S). The blue and orange polyhedra correspond to the M (16*d*, *oct*) and Mg (8*a*, *tet*). The dashed rectangle indicates the vacant 16*c*, *oct* site and the dashed triangle the vacant 48*f tet* site. In panel b, green arrows display the exchange of Mg and M sites, leading to inversion in the spinel.

Table 1.	Notations	Used in	AM_2X_4	Structure	of Figure	1 ^a
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			sharing neighbors			
site	coordination	ion in normal spinel	face	edge	corner	no. sites
8 <i>a</i>	tet	A (Mg ²⁺)	16c	48 <i>f</i>	48f, 16d, 8b	8
16 <i>d</i>	oct	$M (Mn^{3+,4+}/In^{3+})$	8b, 48f	16c, 16d	8a, 48f	16
16c	oct	Vac	8a, 48f	16d, 16c	8b, 48f	16
48f	tet	Vac	16d, 16c	8a, 8b, 48f	8a, 8b, 16c, 16d	48
86	tet	Vac	16d	48 <i>f</i>	48f, 16c, 8a	8
a Vac indicates vacancy. No. sites is normalized against the conventional (cubic) cell of a normal spinel with 32 anions.						

electrolytes combine the advantage of improved safety with a high Mg transference number. Three promising compounds were reported, namely, $MgSc_2Se_4$, $MgSc_2S_4$, and $MgIn_2S_4$.³⁸ $MgIn_2S_4$ spinel had previously been reported,^{39,40} and the available literature as well as our own synthesis attempts (Figure S1 in Supporting Information, SI) indicate that the compound is prone to inversion, where the degree of inversion can be as high as ~85% (Table S2, SI).

In the present work, motivated by the importance of the spinel structure for MV battery technology, we explore the influence of spinel inversion on Mg mobility in ternary oxides and sulfides, using MgMn₂O₄ and MgIn₂S₄ as the prototype for each class of spinels. We consider all possible local cation environments that arise due to inversion and compute the activation barriers for Mg migration in each scenario using firstprinciples calculations. The high requirement for the ionic conductivity in solid electrolytes typically demands migration barriers to be <500 meV, as observed in solid Li-conductors,⁴¹ while cathodes can operate under lower ionic mobilities (barriers \sim 750 meV, see Section 4.1)² as the required length is less than for a conductor. Hence, we limit accessible Mg²⁺ migration paths to those with a barrier less than 500 and 750 meV for operation as a solid electrolyte and cathode, respectively. We will use MgMn₂O₄ as the prototype cathode for which we restrict barriers to 750 meV and MgIn₂S₄ as an example of an electrolyte (barriers <500 meV).

Our results indicate that inversion, in both solid electrolytes and cathodes, can simultaneously cause a decrease in activation barriers across certain migration trajectories while increasing the barriers across others, leading to a complex interplay of opening and closing of specific Mg migration pathways. To quantify the impact of these variations in the microscopic activation barriers on macroscopic Mg diffusion, we estimate the critical Mg concentrations (percolation thresholds) required to facilitate Mg²⁺ diffusion through the structure at different degrees of inversion. Note that Mg extraction from the cathode material creates Mg-vacancies that can affect the percolation properties. For example, vacancies can cause migration pathways that are inactive in the fully discharged composition to become accessible. Hence, for a cathode, we examine the variation of the percolation threshold with vacancy content in the spinel lattice. In electrolytes, the Mg concentration does not significantly vary and we do not consider the effect of Mg-vacancies in MgIn₂S₄. Our estimates indicate that stoichiometric MgMn₂O₄ and MgIn₂S₄ spinels remain percolating up to ~55-59% and 44% inversion, respectively. Finally, we discuss the impact of spinel inversion on Mg-electrochemistry in the Mn₂O₄ cathode by evaluating the 0 K phase diagram, average voltages, and the accessible Mg capacity at various degrees of inversion.

While previous studies have analyzed the impact of inversion on structural, thermal, electronic, and magnetic properties, $^{30,42-45}$ the effect on Mg mobility in spinels has not yet



Figure 2. Local cation environments and various Mg hops considered in an inverted spinel structure. In all migration scenarios a Mg atom migrates from an occupied site (indicated by solid black circles) to an adjacent vacant site (dashed black rectangles), along the trajectory indicated by the arrows. Hops (a) 1 and (b) 2 occur with a *tet* \rightarrow *oct* \rightarrow *tet* topology, while hops (c) 3, (d) 4, and (e) 5 occur along an *oct* \rightarrow *tet* \rightarrow *oct* pathway. Blue and orange polyhedra correspond to Mg and M (M = Mn, In), while green polyhedra indicate mixed M/Mg occupancy. In the case of Hops 3, 4, and 5, the 8*a* sites corner-sharing with the intermediate 48*f* site are shown as gray polyhedra. The notation "edge" in panels c, d, and e corresponds to the 8*a* site that edge-shares with the 48*f*. Vac indicates vacancy.

been investigated. Understanding the influence of inversion on ion mobility will provide guidelines to tune the synthesis and electrochemical conditions of both cathodes and solid electrolytes, not only in MV systems but also in existing Li-ion architectures.⁴⁶ Finally, our results emphasize the importance of the topology of cation sites in setting the migration behavior within a general anion framework.⁴⁷

2. STRUCTURE

A spinel configuration is a specific ordering of cation sites (A and M in AM_2X_4) in a face-centered cubic (FCC) packing of anion sites (X), as shown in Figure 1. In a "normal" spinel, half of the octahedral (*oct*) sites, that is, 16*d*, are occupied by M atoms (Mn/In, blue octahedra in Figure 1), while 1/8 of the

tetrahedral (*tet*) sites (8*a*) are occupied by A (Mg, orange tetrahedra) cations.

Polyhedra in the spinel structure share faces, edges and corners, as summarized in Table 1. For example, the 8*a* sites that are occupied by A are face-sharing with vacant (Vac) 16*c* oct sites (dashed red square in Figure 1a), edge-sharing with vacant 48*f tet* (dashed red triangle) and corner-sharing with vacant *tet* (48*f*, 8*b*) and M-containing 16*d oct* sites.⁴⁸ Face-sharing polyhedra have the lowest cation—cation distance, leading to the highest level of electrostatic repulsion, followed by edge-sharing and subsequently corner-sharing polyhedra.⁴⁹ Indeed, the 16*c*, 48*f*, and 8*b* sites are vacant in spinel lattices (8*b* not shown in Figure 1) since they face-share with occupied 8*a* or 16*d* sites.

Inversion in a spinel structure refers to the collection of antisite defects in the 8*a* (A) and 16*d* (M) sublattices, as shown in Figure 1b. The degree of inversion, *i*, is defined as the fraction of 8*a* sites occupied by M cations, with a value of 0 (or 0%) and 1 (100%) indicating a normal and a fully inverted spinel, respectively. Thus, cations A and M are exchanged in inverted spinels (green arrows in Figure 1b), leading to a stoichiometry of $A_{1-i}M_i[A_{i/2}M_{1-(i/2)}]_2X_4$, compared to AM_2X_4 in normal spinels.

2.1. Possible Mg-Hops. Figure 2 and Table 2 summarize the possible local cation arrangements in a spinel structure that

Table 2. Summary of All Hops Considered for Evaluating Mg^{2+} Mobility in Inverted Spinels, Where M = Mn, In and Vac = Vacancy^a

hop	topology	intermediate site neighbor(s)	no. configurations
1	8a-16c-8a (tet-oct-tet)	16d (oct, M)	1
2	8a-16c-8a (tet-oct-tet)	16d (oct, Mg/M)	6
3	16d-48f-16d (oct-tet-oct)	8a (tet, Mg)	2
4	16d-48f-16d (oct-tet-oct)	8a (tet, M)	2
5	16d-48f-16d (oct-tet-oct)	8a (tet, Vac)	2

^{*a*}Neighbor column indicates the site that edge-shares with the intermediate site in the corresponding hop. The last column signifies the (maximum) number of configurations, along each migration trajectory, for which migration barriers have been calculated in this work. For example, along Hop 3, the corner-8*a* sites being cation-occupied and vacant are the two configurations considered.

can originate from inversion. The orange, blue, and green polyhedra in Figure 2 correspond to Mg, M, and mixed (Mg/M) occupation, respectively, with the arrows in each panel indicating the Mg migration trajectory. The dashed rectangles and triangles signify vacancies. Gray polyhedra correspond to 8a sites that are either cation occupied or vacant. While Figure 2a indicates the migration trajectory in a normal spinel, panels b, c, d, and e depict the possible Mg-hops that can occur in an inverted spinel. The subpanels in Figure 2b correspond to slices along perpendicular directions, that is, the 8a sites in the left subpanel of Figure 2b are perpendicular to the plane of the paper in the right subpanel.

In a normal spinel, the rate for Mg diffusion is determined by the hop between adjacent 8*a* tet sites face-sharing with a 16*c* octahedron, as shown in Figure 2a. Hence, the migration topology is tet-oct-tet, and referred to as "Hop 1" in our work. The intermediate 16*c* site in Hop 1 shares edges with six 16*d* oct sites ("ring" sites) that are occupied by M cations (2 out of 6 ring sites are shown in Figure 2a). It was recently proposed^{8,11,12} that the migration barrier in normal spinels, both oxides and sulfides, is predominantly set by the size of the shared triangular face (not shown in Figure 2a) between the 8*a* tet and 16*c* oct sites.

Along the tet-oct-tet migration pathway in inverted spinels (referred to as "Hop 2"), the 16d ring sites can be occupied by both M and Mg cations, as indicated by the six green polyhedra in the right subpanel of Figure 2b. To evaluate Mg²⁺ migration along Hop 2, we considered multiple configurations from 1 ring site occupied by Mg to all 6 ring sites being occupied by Mg. Since each ring site occupancy (e.g., 2/6 or 3/6 Mg) corresponds to a large number of possible cation decorations on the ring sites, we used the decoration that had the lowest electrostatic energy, as obtained by minimizing the Ewald energy of the unit cell⁵⁰ using classical charges in the spinel

framework. The specific cation arrangements used to evaluate the Mg migration barriers along Hop 2 are displayed in Figure S13.

As inversion leads to Mg²⁺ occupancy of 16d sites, Mghopping across 16d sites must also be considered. A 16d-16d hop can occur through two possible tetrahedral intermediate sites, the 8b and 48f. The 8b sites typically share all their triangular faces with occupied 16d sites and are therefore not open to Mg^{2+} migration due to high electrostatic repulsion, as shown by previous studies.^{8,36,47} However, the 48f sites share two triangular faces with vacant 16c sites, enabling them to act as viable intermediate sites for Mg²⁺ hopping. As such, we only consider the 16d-16d hop via the 48f as intermediate site, leading to a 16d-48f-16d topology (Figure 2c,d,e). The 48f shares one of its edges with an 8a tet site (Table 1), where the "edge-8a" can be occupied by Mg^{2+} ("Hop 3", Figure 2c), $M^{3+/4+}$ ("Hop 4", Figure 2d) or a vacancy ("Hop 5", Figure 2c). Additionally, across Hops 3, 4, and 5, we consider two scenarios where the 8*a* sites that share a corner with the 48*f* ("corner-8*a*", gray polyhedra in Figure 2) are either occupied by cations or left vacant.

2.2. Percolation Theory. While activation barriers for the various cation arrangements in Figure 2 determine the active Mg^{2+} migration hops (or channels) on the atomic scale, the macroscopic diffusion of Mg²⁺, which is essential for (dis)charge of cathodes or ionic conduction in solid electrolytes, depends on the existence of a percolating network of active migration channels. As the 8a-16c-8a channels form a percolating network throughout the spinel structure, stoichiometric normal spinels with Mg in 8a enable macroscopic diffusion of Mg²⁺ as long as the 8a-16c-8a hop is open, that is, the migration barrier for Hop 1 is below a threshold value. However, inversion leads to mixing of cation occupancies in both the 8a and 16d sites, potentially causing some 8a-16c-8achannels to close (due to higher Mg²⁺ migration barriers along Hop 2) while opening new channels typically closed in a normal spinel (e.g., Hops 3, 4, or 5). Hence, in addition to identifying facile microscopic hops, it is important to consider whether a percolating network of low-barrier migration channels exists. Analogous studies have been done on Li⁺ percolation in rocksalt lattices.³⁶

In percolation theory, the site percolation problem^{S1-54} identifies the critical concentration, $x = x_{crit}$ at which an infinite network of contiguous connected sites exists in an infinite lattice of randomly occupied sites. In terms of ionic diffusion, x_{crit} sets the "percolation threshold", above which percolating channels exist in a given structure and macroscopic ion diffusion is feasible. While percolation thresholds are accessible analytically for 2D lattices,⁵³ Monte Carlo (MC) simulations need to be used to estimate x_{crit} in 3D structures.

The existence of a percolating diffusion network in a structure at a certain x ($>x_{crit}$) does not imply that all ions in the structure can be (reversibly) extracted. Mg sites that are not part of a percolating network will form isolated clusters throughout the structure so that the amount of extractable ions is lower than the total concentration, that is, $x_{ext} < x$. The quantity x_{ext} can be assumed to correspond to the capacity of a cathode material. Numerically, x_{ext} is also estimated from MC simulations.³⁶

In summary, the two central quantities obtained from percolation MC simulations are the Mg concentration beyond which macroscopic diffusion is feasible (x_{crit}) and the fraction of extractable Mg ions in a percolating structure (x_{ext}) . To study

Mg diffusion in spinels, we modified the nearest neighbor model (normally considered in site percolation estimations) to include occupancies up to the third nearest neighbor (i.e., corner-sharing sites in Table 1). Two Mg sites in a given spinel arrangement are considered connected only if the migration channel linking them is open (i.e., the migration barrier is below an upper-limit). Thus, a percolating network of Mg sites is formed solely via open migration channels. Whether a channel is considered open will depend on the migration barrier for Mg hopping through it.

3. METHODS

The computational approaches to predict properties relevant to cathode materials have recently been reviewed by Urban et al.⁵⁷ Also, the ability of density functional theory (DFT)^{58,59} methods to predict materials with novel properties has been amply demonstrated.⁵⁰ As a result, all calculations in this work are done with DFT as implemented in the Vienna Ab Initio Simulation Package,^{61,62} and employing the Projector Augmented Wave theory.⁶³ An energy cutoff of 520 eV is used for describing the wave functions, which are sampled on a wellconverged k-point $(4 \times 4 \times 4)$ mesh. The electronic exchangecorrelation is described by the semilocal Perdew-Burke-Ernzerhof (PBE)⁶⁴ functional of the Generalized Gradient Approximation (GGA). Calculations on Mg₂Mn₂O₄ are always initialized with an ideal cubic structure while allowing for potential tetragonal distortions during the geometry relaxation as the spinel can be either cubic ($x_{Mg} \approx$ 0) or tetragonal $(x_{Mg} \approx 1)$ based on the concentration of Jahn–Teller active Mn^{3+} ions. The computed c/a ratio for the tetragonal-MgMn₂O₄ structure is in excellent agreement with experimental reports^{29,65} (see Section S12). For voltage and 0 K phase diagram calculations of Mg_xMn₂O₄, the PBESol exchange-correlation functional⁶⁶ is used to improve the description of the energetics,⁶⁷ while a Hubbard Ucorrection of 3.9 eV is added to remove spurious self-interaction of the Mn *d*-electrons.^{68–70}

The activation barrier calculations are performed with the Nudged Elastic Band (NEB) method.^{71,72} The barriers are calculated in a conventional spinel cell (32 anions), which ensures a minimum distance of ~8 Å between the elastic bands and reduces fictitious interactions with periodic images. We verified that migration barriers do not change appreciably (<3% deviation) when equivalent calculations are performed in larger supercells (see Figure S2). Seven images are introduced between the initial and final end points to capture the saddle point and the migration trajectory. All NEB results are based on the PBE functional, without Hubbard U.^{10,11} The migration barriers in spinel-MgIn₂S₄ are calculated with compensating electrons added as a background charge to ensure charge-neutrality of the structure at nonstoichiometric Mg concentrations.

As migration barriers are calculated in the conventional spinel cell, the degree of inversion (*i*) that can be modeled is constrained by the migration trajectory under consideration, in both the oxide and the sulfide. For example, along Hop 2 (Figure 2b), a 3/6 Mg ring site occupancy leads to 3 Mn/In atoms in the 8*a* sites, and consequently results in $i \approx 3/8 = 0.375$. Similarly, the barrier calculations along the 16d-48f-16d topology (Hops 3, 4, and 5), which require a minimum of 2 Mg atoms in the 16d sites (or 2 Mn/In sites in the 8*a*), correspond to $i \approx 0.25$.

Monte Carlo simulations are used to estimate the Mg percolation thresholds (x_{crit}) and the fraction of extractable Mg ions (x_{ext}). A 6 × 6 × 6 supercell of the primitive spinel structure is used, which corresponds to 1728 anion atoms (Figure S6 plots convergence behavior with supercell size). In MC simulations, a network of Mg sites is considered percolating when it spans the periodic boundaries of the simulation cell in one or more directions.⁷³ Inversion in the spinel is introduced during MC sweeps by labeling a number of random 8*a* and 16*d* sites, corresponding to the degree of inversion, as part of the "Mg sub-lattice". For example, the Mg sublattice in a normal spinel consists of all 8*a* sites. However, in an inverted spinel (with the degree

of inversion *i*) the Mg sublattice will be composed of (1 - i)% of all 8*a* sites and (i/2)% of all 16*d* sites.

To evaluate the M composition at which percolation occurs, a MC sweep is performed with the following steps:⁷³ (i) the supercell is initialized with M atoms in both M and Mg "sub-lattices", corresponding to a M_3X_4 (X = O, S) stoichiometry, (ii) M atoms on the Mg sublattice are randomly changed to Mg, (iii) after all Mg sublattice sites are changed (i.e., a stoichiometry of MgM2X4 is attained), M atoms on the M sublattice are randomly flipped to Mg. During an MC sweep, once a Mg atom replacement results in the formation of a percolating network, the current Mg concentration (x_{Mg}) is taken as an estimate of the percolation threshold (x_{crit}) , while for $x > x_{crit}$, the fraction of sites within the percolating network, x_{ext} is stored. The values of x_{crit} and x_{ext} are averaged over 2000 MC sweeps to guarantee well-converged estimates. The effect of vacancies on Mg percolation in the Mn-spinel is captured by initializing the Mg sublattice with varying vacancy concentrations, at a given degree of inversion, corresponding to a $Vac_zMn_{3-z}O_4$ stoichiometry ($z \le 1$). Whenever vacancies are initialized in a supercell, only the Mn atoms are changed to Mg during a MC sweep.

4. RESULTS

4.1. MgMn₂O₄. Figure 3 plots the ranges of Mg^{2+} migration barriers in $Mg_{x}Mn_{2}O_{4}$ (y-axis) for all hops of Figure 2 and



Figure 3. Ranges of Mg²⁺ migration barriers along the hops considered in spinel-Mg_xMn₂O₄₊ The dotted black line indicates the upper-limit of migration barriers (~750 meV) used to distinguish open and closed migration channels in percolation simulations. Solid red and dashed blue lines correspond to dilute Mg ($x_{Mg} \approx 0$) and dilute vacancy ($x_{Mg} \approx 1$) limits. Fractions along Hop 2 indicate the occupancy of Mg²⁺ in the 16*d* ring sites, while the legend "8*a* full (empty)" corresponds to cation-occupied (vacant) corner-8*a* sites along Hops 3–5. The barriers along Hop 1 are calculated at *i* ≈ 0, while Hops 3–5 have been done at *i* ≈ 0.25. Along Hop 2, *i* varies with Mg occupancy of the ring sites, ranging from *i* ≈ 0.125 at 1/6 Mg to *i* ≈ 0.75 at 6/6 Mg. The raw data from Nudged Elastic Band calculations are displayed in Figure S3 of the SI.

Table 2, while the raw data is included in Figure S3 of the SI. The migration barriers are calculated with respect to the absolute energies of the end points, nominally identical for a given Mg^{2+} hop. However, there are a few cases where the end point energies are different, since the local symmetry of the cation decoration is broken differently across the end points (e.g., 3/6 hop in Figure S3b). In such cases, the barrier is reported with respect to the end point with the lowest energy. The dotted black line in Figure 3 is the upper-limit of the Mg migration barrier, as required for reasonable battery perform-

ance,² and is used to determine the percolation thresholds (see Section 4.3). For a Mg₂Mn₂O₄ cathode particle of size ~ 100 nm being (dis)charged at a C/3 rate at 60 °C, the migration barrier upper-limit is ~750 meV (the upper-limit decreases to ~660 meV at 298 K).² Since full-cell Mg batteries so far have displayed superior performance at ~60 °C than at 25 °C,^{1,74} the value of ~750 meV has been used as the cutoff to differentiate "open" and "closed" Mg²⁺ migration channels. In terms of notations, the fractions used in Hop 2 (e.g., 1/6, 2/6, etc., yellow rectangle in Figure 3) correspond to the fraction of 16*d* ring sites (Figure 2b) that are occupied by Mg^{2+} . The terms "8a empty" and "8a full" along Hops 3, 4, and 5 in Figure 3 indicate that the corner-8a sites (Figure 2c,d,e) are vacant and occupied by cations, respectively. x_{Mg} in Figure 3 is the Mg concentration in the cell used for the barrier estimation, corresponding to the "dilute Mg" ($x_{Mg} \approx 0$, solid red lines) and "dilute vacancy" ($x_{Mg} \approx 1$, dashed blue lines) limits.

Mg migration barriers along Hop 1 (tet-oct-tet, normal spinel) at the dilute Mg and dilute vacancy limits are ~717 meV and ~475 meV, respectively (red rectangle in Figure 3), in good agreement with previous studies.^{8,11,75} Note that the dilute Mg (vacancy) limit for Hop 1 corresponds to the regime when no 8a sites, other than those required to model the hop, are occupied by Mg (vacancies). Since the migration barriers at both Mg concentration limits are below ~750 meV, Hop 1 is always open for Mg migration. Barriers along Hop 2 (yellow rectangle in Figure 3) decrease initially with Mg occupation of the 16d ring sites (\sim 393 meV at 2/6 vs 536 meV at 1/6) before increasing beyond 750 meV at 5/6 and 6/6 Mg. The nonmonotonic variation of the migration barriers along Hop 2 is due to the gradual destabilization of the 16c site. The increasing instability of the 16c also changes the migration energy profile (Figure S3b) from "valley"-like⁸ at 1/6 Mg to "plateau"-like at 5/6 Mg. Figure S14 shows the Mg migration barriers along Hop 2 when the ring sites are occupied by vacancies instead of Mg²⁺.

In the case of the *oct*-*tet*-*oct* Hops 3 and 4 (green and cyan rectangles in Figure 3), which, respectively, have tet Mg and Mn edge-sharing with the intermediate 48f site, the barriers vary drastically based on Mg content and occupancy of the corner-8*a* sites. For example, at (i) $x_{Mg} \approx 0$ and vacant corner-8*a*, the barrier along Hop 3 (~ 592 meV) is well below the upperbound of 750 meV, while the barrier is comparable along Hop 4 (~743 meV). At (ii) $x_{Mg} \approx 0$ and cation-occupied corner-8*a*, the barriers along Hops 3 and 4 increase significantly (~1388 meV and ~1418 meV) and surpass the upper-limit set for open channels. Eventually, at (iii) $x_{Mg} \approx 1$ (cation-occupied corner-8a), the barriers decrease to ~ 845 meV and ~ 784 meV along Hops 3 and 4, respectively. Note that the barriers along Hops 3 and 4 in Figure 3 are calculated at a degree of inversion, $i \approx$ 0.25. At a higher degree of inversion $(i \approx 1)$ and $x_{Mg} \approx 1$ (cation-occupied corner-8a), the barrier is ~ 1039 meV along Hop 4 (Figure S5). Hence, from the data of Figure 3, Hop 3 is considered closed for Mg migration whenever the corner-8a sites are cation-occupied, while Hop 4 is always considered a closed channel.

Mg migration barriers decrease significantly if the edge-8*a* is vacant (i.e., along Hop 5). For example, the migration barriers along Hop 5 (purple rectangle in Figure 3) are well below that of Hops 3 and 4 across the scenarios of (i) low Mg, vacant corner-8*a* (319 meV for Hop 5 vs 592 and 743 meV for Hops 3 and 4, respectively), (ii) low Mg, cation-occupied corner-8*a* (703 meV vs 1388 and 1418 meV), and (iii) high Mg, cation-

occupied corner-8a (570 meV vs 845 and 784 meV). Hence, Hop 5 is always open for Mg migration, since the barriers are below the upper limit of 750 meV.

In summary, the tet-oct-tet pathway (Hops 1 and 2) remains open for Mg migration in MgMn₂O₄ until a high degree of Mg occupation on the 16*d* ring sites (i.e., $\geq 5/6$ Mg) is present, which corresponds to high degrees of inversion (i > 0.625). The *oct-tet-oct* pathway is open only when the edge-8*a* is vacant (Hop 5) or when the corner-8*a* are vacant with Mg in the edge-8*a* (Hop 3).

4.2. $Mgln_2S_4$. Figure 4 plots the Mg^{2+} migration barriers in $Mgln_2S_4$ for the hops of Figure 2 (the raw data are shown in



Figure 4. Mg²⁺ migration barriers along each possible hop in spinel-MgIn₂S₄. The dotted black line indicates the upper-limit of migration barriers (~500 meV) used to distinguish open and closed migration channels in percolation simulations. Dashed blue lines indicate the dilute vacancy ($x_{Mg} \approx 1$)) limit. Fractions along Hop 2 indicate the occupancy of Mg²⁺ in the 16*d* ring sites, while the corner-8*a* sites are cation-occupied across Hops 3–5. The barrier along Hop 1 is calculated at *i* \approx 0, while Hops 3–5 have been done at *i* \approx 0.25. Along Hop 2, *i* varies with Mg occupancy of the ring sites, ranging from *i* \approx 0.125 at 1/6 Mg to *i* \approx 0.75 at 6/6 Mg. The raw data from Nudged Elastic Band calculations are displayed in Figure S4.

Figure S4). Since we consider MgIn₂S₄ as an ionic conductor, off-stoichiometric Mg concentrations are not of interest. Hence, all hops in Figure 4 are evaluated at the dilute vacancy limit ($x_{Mg} \approx 1$, dashed blue lines in Figure 4). The fractions used (1/6, 2/6, etc.) in Figure 4 are the number of 16*d* ring sites occupied by Mg²⁺ in Hop 2. Along Hops 3–5, we use cation-occupied corner-8*a* sites (i.e., "8*a* full" in Figures S4c, d, and e). The upper-limit of the Mg migration barrier for classifying open and closed migration channels (as indicated by the dotted black line in Figure 4) is set to ~500 meV, based on migration barriers of ~400–500 meV observed in fast Li-ion conductors, such as Garnets and Si-based thio-LISICONs.⁴¹

In the case of Hop 1, the barrier is ~447 meV, well below the upper limit of ~500 meV. Mg migration barriers along Hop 2 (yellow rectangle in Figure 4) follow trends similar to that of MgMn₂O₄ (Figure 3). For example, at low Mg occupation of the ring sites (1/6 or 2/6 Mg), the barrier is below the limits for percolating diffusion, before increasing beyond 500 meV at higher Mg content in the ring sites (>3/6 Mg). Also, the shape of the migration energy curve changes from a "valley" at 1/6 Mg (solid black line in Figure S4b) to a "plateau" beyond 2/6

Mg (solid red line in Figure S4b), indicating that the 16c site becomes progressively unstable with increasing Mg occupation of the ring 16d.

Along the 16d-48f-16d pathways (Hops 3, 4 and 5), the migration barriers are always higher than 500 meV, irrespective of the occupancy of the edge-8*a*. Indeed, the magnitude of the barriers are ~683 meV, ~531 meV, and ~504 meV for Mgoccupied, In-occupied and vacant edge-8*a*, respectively, indicating that the *oct-tet-oct* pathway will not be open for Mg²⁺ migration.

4.3. Percolation Thresholds. On the basis of the data of Figures 3 and 4, and the upper limits of Mg migration barriers set for $MgMn_2O_4$ (750 meV) and $MgIn_2S_4$ (500 meV), we compiled a list of conditions that enable the opening of the possible hops in Table 3. For example, Hop 1 (8*a*-8*a*) is open

 Table 3. Summary of Rules Used during Percolation

 Simulations with Conditions for an Open Channel^a

hop	topology	open under condition		
MgMn ₂ O ₄ – 750 meV				
1	8a-16c-8a	always open		
2	8a-16c-8a	max. 4/6 ring sites with Mg		
3	16 <i>d</i> -48 <i>f</i> -16 <i>d</i>	corner 8 <i>a</i> vacant		
4	16d-48f-16d	always closed		
5	16 <i>d</i> -48 <i>f</i> -16 <i>d</i>	always open		
$MgIn_2S_4 - 500 meV$				
1	8a-16c-8a	always open		
2	8a-16c-8a	max. 2/6 ring sites with Mg		
3	16 <i>d</i> -48 <i>f</i> -16 <i>d</i>	always closed		
4	16 <i>d</i> -48 <i>f</i> -16 <i>d</i>	always closed		
5	16 <i>d</i> -48 <i>f</i> -16 <i>d</i>	always closed		

^{*a*}Upper limit of migration barriers used to distinguish between open and closed channels is 750 and 500 meV for MgMn₂O₄ and MgIn₂S₄, respectively.

for all values of x_{Mg} and *i* for both Mg_xMn₂O₄ and MgIn₂S₄. Both the oxide and the sulfide spinel exhibit high barriers (>1 eV) for a 16*d*-8*a* hop (Figure S8), which would limit Mg transfer between an octahedral 16*d* site and an adjacent tetrahedral 8*a* site. Thus, in our percolation simulations, the 8*a*-8*a* (Hops 1 and 2) and the 16*d*-16*d* (Hops 3, 4, and 5) channels remain decoupled, and a percolating network consists solely of either 8*a*-8*a* or 16*d*-16*d* channels.

Figure 5a and b plot the percolation threshold (x_{crit}) black lines), at various degrees of inversion (*i*) in $Mn_{3-x}O_4$ and $In_{3-x}S_4$. The dashed yellow lines indicate the stoichiometric spinel, that is, M:X = 2:4. The blue (red) shaded region corresponds to Mg concentration ranges which do (do not) exhibit percolation. The *x*-axis in Figure 5 begins at a M_3X_4 (i.e., 50% M-excess or 100% Mg-deficient) configuration and spans concentrations up to $Mg_{1.5}M_{1.5}X_4$ (i.e., 25% M-deficient, 50% Mg-excess). Generally, percolation thresholds in the M-excess domain (i.e., $x_{crit} < 1$) are desirable as this implies that the stoichiometric spinel will possess percolating networks and will facilitate macroscopic Mg transport.

In the case of cathodes (Mn_2O_4) , Mg deintercalation from the framework creates vacancies, which can facilitate the formation of Mg percolating networks by opening certain migration channels (e.g., Hop 5 in MgMn_2O_4, Figure 3). Therefore, we explored the variation of the percolation threshold with vacancy concentration ("z" in Figure 5a) in the Mn-spinel. For the sake of simplicity, x in Figure 5 refers to



Figure 5. Critical concentration for Mg percolation (x_{crit}) in the (a) $(Mg_{x-z}Vac_z)Mn_{3-x}O_4$ and (b) $Mg_xIn_{3-x}S_4$ spinels are plotted as black lines at different degrees of spinel inversion *i*. The stoichiometric spinel concentration (M:X = 2:4) is indicated by the dashed yellow lines. Note that the zero on the *x*-axis corresponds to a stoichiometry of M_3X_4 (M = Mn/In and X = O/S). z indicates the vacancy concentration in the structure. The shaded red (blue) region in both panels indicates the Mg concentration range where macroscopic Mg diffusion is not possible (possible). The shaded gray region in panel a refers to the range of variation of the percolation threshold with vacancy content in the oxide cathode. The green circle, square and star in panel a correspond to sample scenarios discussed in the text.

the sum of Mg and vacancy concentrations. For example, x = 1 and z = 0.5 (green circle on the dashed black line) in Figure 5a indicates a composition of Mg_{0.5}Vac_{0.5}Mn₂O₄, while x = 0.6 and z = 0 (green square) corresponds to the M-excess spinel-Mg_{0.6}Vac_{0.5}Mn_{2.4}O₄.

The percolation threshold in the absence of vacancies (z = 0) is indicated by the solid black line in Figure 5a. When vacancies are introduced, the threshold decreases, as indicated by the x_{crit} at z = 0.4 (dotted black line) or 0.5 (dashed) consistently exhibiting lower values than x_{crit} at z = 0 in Figure 5a. For example, at x = 0.8 and i = 0.5 (indicated by the green star in Figure 5a), the spinel does not form a percolating network when there are no vacancies (z = 0, Mg_{0.8}Mn_{2.2}O₄), since x_{crit} ~0.88 > 0.8. However, the structure can percolate Mg when vacancies are introduced (z = 0.5, $Mg_{0.3}Vac_{0.5}Mn_{2.2}O_4$), as x_{crit} reduces to ~0.52 < 0.8. In a case such as this, the initial cathode structure may not be percolating, but introducing vacancies in the initial part of the charge can create a percolating zone on the cathode particle surface through which further Mg-removal can occur. However, upon discharge, the percolating structure could easily become nonpercolating if polarization increases the surface Mg concentration too rapidly.

At any degree of inversion, the magnitude of x_{crit} varies nonmonotonically and reduces only up to a vacancy content, z = 0.4 or 0.5 (see Figure S7a). Indeed, at x = 0.8 and i = 0.5(green star), an increase in z beyond 0.5 (such as z = 0.6, $Mg_{0.2}Vac_{0.6}Mn_{2.2}O_4$) causes the x_{crit} to increase to ~0.6, but the spinel continues to percolate. Thus, the shaded gray region in Figure 5a, which is bound by the z = 0.4, 0.5, and 0 lines, represents the extent of variation of x_{crit} with vacancy content in the cathode. Notably, the lowest value of x_{crit} is obtained at z =0.4 for $0 \le i \le 0.35$ and $0.595 \le i < 0.77$, and at z = 0.5 for 0.35 $\le i \le 0.595$, respectively.

The stoichiometric {Mg/Vac}Mn₂O₄ spinel at i = 0 (dashed yellow line in Figure 5a), permits macroscopic Mg diffusion, since the percolation threshold ($x_{crit} \approx 0.44$ for z = 0-0.4) is in the Mn-excess domain (i.e., $x_{crit} < 1$). When vacancies are absent in the stoichiometric spinel (z = 0), which corresponds to the discharged MgMn₂O₄ composition, the structure percolates Mg up to $i \approx 0.55$. Upon charging, the presence of vacancies (z = 0.5) enables Mg percolation within Mg_{0.5}Vac_{0.5}Mn₂O₄ up to $i \approx 0.59$. At higher degrees of inversion (0.59 < i < 0.77), the oxide spinel requires Mn-deficient concentrations (i.e., x > 1) to facilitate Mg percolation, as illustrated by $x_{crit} \approx 1.05-1.13$ (z = 0.5-0) at i = 0.6. At i > 0.77, the oxide does not form a percolating Mg network at any level of Mn-deficiency (for $z \le 1$) in the lattice.

In stoichiometric ionic conductors, such as $MgIn_2S_4$, the vacancy concentration is low and therefore vacancies are not expected to play a major role in macroscopic Mg transport. Specifically in MgIn₂S₄, vacancies do not open additional migration channels, as indicated by the closed Hop 5 in Figure 4. Indeed, the percolation threshold in the In-spinel does not change up to a vacancy content, z = 0.2 in the structure (see Figure S7b). At z = 0, the x_{crit} in $In_{3-x}S_4$ (solid black line in Figure 5b) increases continuously with increase in inversion, with $x_{crit} \approx 0.435$, and 0.74 at i = 0, and 0.4, respectively. Thus, at low *i*, stoichiometric MgIn₂S₄ should exhibit significant ionic conductivity. However, at higher degrees of inversion (i > i)0.44), the sulfide spinel does not form percolating networks at any Mg-concentration, owing to the absence of open 16d-16dchannels in combination with the 8a-8a channels being closed beyond 2/6 Mg ring site occupancy (Table 3).

In general, mobility requirements in an ionic conductor are more stringent than in a cathode, consistent with the stricter cutoff of 500 meV we applied to the migration barriers in $MgIn_2S_4$.^{41,76} Indeed, a sulfide spinel Mg-cathode (such as $Mg_xTi_2S_4^{-74}$) exhibiting similar activation barriers with inversion as $MgIn_2S_4$ will not suffer from any percolation bottlenecks, since the barriers across all cation arrangements are well below the milder 750 meV cutoff set for cathodes (Figure 4).

4.4. Impact of Inversion on Cathode Electrochemistry. Under ideal conditions, the structure of an ionic conductor (such as $MgIn_2S_4$) should not undergo significant changes during operation. Thus, the extent of inversion should, in principle, be measured using characterization experiments postsynthesis (the calculated formation energies of various inverted configurations in spinel-MgIn₂S₄ are plotted in Figure S11). However, in a cathode material such as Mg_xMn₂O₄, which can generate mobile Mn²⁺ ions (Figure S9) through disproportionation of Mn³⁺, the degree of inversion (*i*) can change during electrochemical cycling.^{16,34} Consequently, structural changes in a cathode during cycling should manifest themselves as changes in the voltage profile and observed capacity, which can be benchmarked with theoretical predictions.^{2,21} To evaluate the effect of inversion on the voltage profile of Mg_xMn₂O₄, we calculated the phase diagram and energy of the intercalation system at 0 K as a function of Mg content under various degrees of inversion.^{16,17,21,77}

To evaluate the ground state hull of the $Mg_xMn_2O_4$ system, we enumerated over 400 Mg-vacancy configurations, at different Mg concentrations ($x_{Mg} = 0, 0.25, 0.5, 0.75$ and 1) and different degrees of inversion (i = 0, 0.25, 0.5, 0.75 and 1). Figure 6a displays structures with formation energies (*y*-axis) below 200 meV/Mn₂O₄ at different Mg concentrations (xaxis), and the formation energies of all the Mg-vacancy configurations considered are plotted in Figure S10 of the SI. Notably, formation energies in Figure 6a have been referenced to the noninverted (i = 0), empty Mn₂O₄ and magnesiated (MgMn₂O₄) spinel configurations. For each configuration, the degree of inversion is indicated by the corresponding symbol used, ranging from i = 0 (black circles) to i = 1 (red stars).

Overall, the Mg_xMn₂O₄ system is phase separating at 0 K across noninverted (i = 0) MgMn₂O₄ and Mn₂O₄ domains, since the ground state hull of the system (dashed black line in Figure 6a) only exhibits two configurations (i.e., MgMn₂O₄ and Mn_2O_4). Some solubility at low Mg content may be possible given the low positive mixing energy at $x_{Mg} = 0.25$ for the noninverted spinel ($E_{\rm formation} \approx 14 \text{ meV}/\text{Mn}_2 \breve{O}_4$). At higher Mg content, the formation energies are very high for the noninverted spinel (Figure S10), making a solid solution behavior very unlikely. Inversion becomes likely to occur at intermediate Mg compositions, as the low positive formation energies are on the scale of the configurational entropy. For example, $E_{\text{formation}} \approx 11 \text{ meV}/\text{Mn}_2\text{O}_4$ at i = 0.25 and $x_{\text{Mg}} = 0.5$ (green square at x = 0.5 in Figure 6a). Hence, inversion at intermediate states of magnesiation is likely. While Mg by definition has to be mobile in Mn_2O_4 to operate as a cathode, Mn mobility, which is required for spinel inversion to occur, depends strongly on its valence state.^{16,34} Typically, Mn³⁺ can be mobile through a temporary disproportionation mechanism, generating mobile Mn²⁺ (Figure S9).^{16,34}

Figure 6b plots the average voltages as a function of x_{Mg} at different *i* by taking the lowest $E_{formation}$ configuration at each *i* and x_{Mg} .⁷⁷ The average voltage for Mg insertion in the noninverted (*i* = 0) configuration is ~2.84 V (dashed black line in Figure 6b), in agreement with previous theoretical estimates.^{11,78} Inversion does increase the average insertion voltage (averaged over $x_{Mg} = 0$ to 1) marginally compared to the normal spinel, with specific values of ~2.92, 2.99, 2.97, and 2.99 V at *i* = 0.25, 0.5, 0.75, and 1, respectively. Notably, the phase behavior of the Mg_xMn₂O₄ system under inversion will be different compared to the normal spinel due to the formation of metastable inverted states at intermediate Mg compositions.

The extractable Mg content (x_{ext} , see Section 2.2), obtained as a function of inversion from our Monte Carlo simulations, indicates the extractable capacity of a cathode particle, and is shown in Figure 6c for stoichiometric MgMn₂O₄. The *y*-axis



Figure 6. (a) Ground state hull (or 0 K phase diagram) of the $Mg_xMn_2O_4$ system, with the zero of the formation energy referenced to the noninverted (i = 0) magnesiated ($MgMn_2O_4$) and empty (Mn_2O_4) spinel configurations. (b) Average voltage curves under i in $Mg_xMn_2O_4$, obtained using the lowest formation energy structures at each i across Mg concentrations. (c) Percentage of the theoretical capacity that can be reversibly extracted is plotted as a function of inversion in stoichiometric MgMn_2O_4.

indicates the % of the cathode's theoretical capacity (~270 mAh/g for MgMn₂O₄), that can be cycled reversibly. At low degrees of inversion, the extractable capacity in the stoichiometric spinel decreases roughly linearly with the degree of inversion, reaching ~41% (~110 mAh/g) at *i* = 0.4. The extractable Mg content decreases more rapidly from *i* = 0.4 to *i* = 0.5, before stabilizing around ~15% (~40 mAh/g) between *i* = 0.5 and 0.6. Eventually, none of the Mg becomes extractable beyond *i* = 0.61, reflecting the trends in the percolation thresholds ($x_{crit} \approx 0.59$ at stoichiometric MgMn₂O₄, Figure 5a) at high degrees of inversion. Note that, the overall amount of cyclable Mg (shown in Figure 6c) and by the phase behavior

as a function of x_{Mg} . For example, if the Mg removal occurs via a two-phase reaction (as is the case for the noninverted spinel), then the presence of a nonpercolating layer on the surface may prevent extraction of Mg from the bulk, even if percolation conditions are still favorable in the bulk material.

5. DISCUSSION

In this work, we have used DFT-based NEB calculations to assess the changes in the activation barrier for Mg^{2+} migration arising from inversion in both oxide (MgMn₂O₄) and sulfide (MgIn₂S₄) structures. From our results (Figures 3 and 4), we can conclude that inversion has a significant impact on both oxides and sulfides, by opening and closing specific migration trajectories. To extrapolate the impact of the various Mg^{2+} migration barriers on macroscopic Mg diffusion, we estimated the percolation thresholds under different degrees of spinel inversion. Furthermore, we analyzed the impact of spinel inversion on cathode properties of $Mg_xMn_2O_4$ by evaluating the average voltages and practical capacities at different degrees of inversion.

5.1. Factors Influencing Barriers in MgMn₂O₄. Trends from activation barriers of Figure 3 suggest that Mg migration along the 8a-16c-8a pathways (Hops 1 and 2) can improve significantly with Mg occupation of the 16*d* ring sites (up to 4/6 Mg), at low degrees of inversion. Additionally, the 16d-48f-16d channels open for Mg migration whenever the edge-8*a* is vacant. However, high degrees of inversion detrimentally affect Mg²⁺ motion, due to the closing of both 16d-16d (corner- and edge-8*a* become occupied by the metal cation) and 8a-8a channels (high migration barriers at high Mg in the ring sites). Although we have specifically considered the case of spinel-Mg_xMn₂O₄, similar trends can be expected for other oxide spinels, given the similarity in Mg migration barriers along Hop 1 with different 3*d*-metals.¹¹

Previous studies have used electrostatic considerations to partially explain trends in Li⁺ activation barriers in a Mn₂O₄ spinel.⁴⁶ Indeed, the reduction in Mg migration barriers along Hops 1, 3, 4, and 5 (Figure 3) with increasing Mg concentration can be attributed to lower electrostatic repulsions at the corresponding intermediate sites caused by the reduction of Mn⁴⁺ to Mn³⁺. For example, the barrier reduces from 717 to 475 meV along Hop 1 and 1388 to 845 meV along Hop 3, as x_{Mg} increases from ~0 to ~1. However, Mg²⁺ activation barriers generally depend on steric and bonding constraints in addition to electrostatics, which are often difficult to deconvolute over a range of NEB calculations. For example, the Mg²⁺ activation barriers across Hop 2 (yellow bar in Figure 3) at low Mg occupation in the ring sites (1/6, 2/6) are lower than Hop 1 (red bar, Figure 3), which may be attributed to reduced electrostatic repulsion on the intermediate 16c (due to Mg²⁺ replacing higher valent Mn in the ring sites). However, barriers along Hop 2 increase beyond Hop 1 and eventually beyond the limit of \sim 750 meV at higher Mg in the ring sites (5/6, 6/6), despite lower electrostatic repulsion. Thus, the high Mg content in the ring sites decreases the stability of the intermediate 16c. One possible reason for the instability of the 16c site could arise from charge-deficient oxygen atoms being shared with adjacent, Mg²⁺-occupied (instead of Mn^{3+/4+}) 16d sites. Indeed, the instability of the 16c (e.g., in the case of 6/6 Mg in Hop 2) is quantified by longer (DFT-based) ≈ 2.3 Å Mg–O bonds, compared to ~ 2.08 Å in 16d with Mg (along the same hop) and ~2.13 Å in rocksalt MgO.

For the 16d-48f-16d hops in Figure 3 (Hops 3-5), electrostatic effects are more dominant than for the tet-oct-tet hops (Hops 1, 2), primarily due to the intermediate 48f edgesharing with an 8a. Indeed, the cation centers in edge-sharing tetrahedra are closer (~ 2.15 Å experimentally between 48f and 8a in an ideal LiMn₂O₄-spinel⁷⁹) than in edge-sharing octahedra (~2.88 Å between 16c and 16d). Consequently, the Mg barriers are consistently lower with a vacant edge-8a (Hop 5, Figure 3) compared to Mg/Mn-filled edge-8a (Hops 3, 4 in Figure 3). Also, Mg²⁺ activation barriers (at $x_{Mg} \approx 0$) increase significantly when the corner-8a sites are cationoccupied rather than vacant (Figure 3). A closer look at the cation-cation distances across corner-sharing 48f and 8a (~2.88 Å in ideal $LiMn_2O_4$) reveals that the corner-sharing tetrahedra within a spinel framework may experience electrostatic repulsion as high as edge-shared octahedra (i.e., 16c and 16d). Thus, the combination of cation–cation repulsion arising from both edge- and corner-8a sites results in the high barriers along Hops 3 and 4.

5.2. Barriers in Sulfides versus Oxides. Activation barriers calculated in $MgIn_2S_4$ (Figure 4) exhibit similar trends to $MgMn_2O_4$ (Figure 3), resulting from analogous trends in electrostatics, steric and bonding environments. However, the absolute changes in barriers in the sulfide are remarkably lower than the oxide. For example, the absolute difference between the lowest and the highest Mg migration barriers of MgMn₂O₄ (at $x_{Mg} \approx 1$) across Hops 1 through 5 is ~662 meV (1055–393 meV), while this is a much lower ~236 meV (683-447 meV) for MgIn₂S₄. Similarly, the barriers along the 16d-48f-16dtrajectory are far less sensitive to the edge-8a occupancy in the sulfide (504-673 meV) than in the oxide (570-845 meV at $x_{Mg} \approx 1$). Surprisingly, the migration barrier with an edge-8*a* occupied by Mg²⁺ is higher (~683 meV) than when the edge-8a is occupied by In^{3+} (~531 meV), suggesting that the In-S bonding environment screens the higher In³⁺ charge better than the Mg–S bonds screen Mg²⁺.

Lower activation barriers for Mg in sulfides have been reported before,^{11,12,38} which have been assigned to robust electrostatic screening, high polarizability, higher degree of covalency and large volume per anion of S²⁻ compared to $O^{2-2,76}$ For example, a Mg_xTi₂S₄⁷⁴ cathode will not suffer from any percolation bottlenecks, if the barriers across all cation arrangements are similar to the calculated values in MgIn₂S₄ (i.e., < 750 meV, Figure 4). But a more stringent upper-bound of ~500 meV on the barrier in a solid-state conductor^{41,76} indicates that inversion can significantly affect a sulfide ionic conductor by closing all 16d-16d channels and several 8a-8achannels with high Mg in the 16d ring (Figure 4). Since ionic mobility is expected to improve with larger anions and higher covalency (such as Se^{2-} compared to S^{2-} and O^{2-}), inversion is expected to affect Mg-mobility to a lesser extent in Mgcontaining Se-spinels, such as MgSc₂Se₄, compared to oxides and sulfides.

5.3. Percolation under Inversion. Estimations of percolation thresholds (x_{crit}) in the Mg_xMn_{3-x}O₄ system (Figure 5a) indicate that spinel inversion should not detrimentally affect macroscopic Mg²⁺ diffusion across the structure up to a fairly high degree of inversion, $i \approx 0.55-0.59$. However, Mg-excess concentrations are required to ensure percolating networks form at i = 0.6-0.7, while the spinel completely ceases to percolate Mg beyond i = 0.77 (Figure 5a). Given the preponderance of conversion reactions under Mg-excess concentrations in the oxide spinel, specifically the

decomposition of $Mg_xMn_{3-x}O_4$ (x > 1) into MgO and MnO,² it is of paramount importance that the chemically synthesizable, stoichiometric {Mg/Vac}Mn₂O₄ remains percolating. Efforts should be made to reduce or precisely control the amount of inversion (i.e., *i* < 0.6), by carefully tuning synthesis temperature and cooling rate^{29,30} during MgMn₂O₄ synthesis.

Higher Mg conductivity, as is required for a solid state electrolyte, demands a lower cutoff for the migration barrier along a pathway. In the case of MgIn₂S₄, where we used a 500 meV cutoff, MC simulations indicate that the stoichiometric spinel should remain percolating up to $i \approx 0.44$. However, high degrees of inversion ($i \approx 0.85$) can be observed during MgIn₂S₄ synthesis (Figure S1). As a result, strategies to limit inversion (i.e., i < 0.44) in sulfide spinel ionic conductors, such as chemical doping and careful calibration of synthesis conditions, need to be sought.

5.4. Voltages and Capacities. Inversion can also significantly impact electrochemical properties, such as phase behavior, average voltages, and extractable capacities in an oxide-spinel cathode (Figure 6). For example, the average voltage for Mg intercalation, across $x_{Mg} = 0-1$ in the Mn₂O₄spinel, is higher in an inverted spinel compared to a normal spinel (Figure 6b). Mg intercalation experiments in spinel- Mn_2O_4 have reported a marginally higher average voltage (~2.9 $(V)^{26}$ than predicted for the normal spinel (~ 2.84 V), with extraction voltages as high as \sim 3.5 V during the charging cycle, which might be an indication of the spinel inverting during electrochemistry. Also, the calculated 0 K phase diagram of the $Mg-Mn_2O_4$ system (Figure 6a) suggests that the tendency to invert is the highest at an intermediate Mg concentration, as indicated by low $E_{\text{formation}}$ (<50 meV/Mn₂O₄) configurations with i = 0.25 at $x_{Mg} = 0.5$. Hence, the degree of inversion in the Mn-spinel can indeed change dynamically during electrochemical Mg cycling, especially due to the presence of mobile Mn²⁺ ions (Figure S9). As reported by Ling et al.,⁸⁰ the mobility of Mn²⁺ within the spinel can also depend on the local arrangement of Mg^{2+} ions. Thus, from the data in Figure 6a, we expect the degree of inversion to vary largely between 0 and 0.25 during Mg-cycling. Also, previous Mg-cycling experiments in spinel-Mn₂O₄ have reported solvent cointercalation based phase transformations,^{28,81} which can be aided by the presence of mobile Mn²⁺ ions.

Additionally, the first Mg-site that will be (de)intercalated in the spinel will depend on the degree of inversion on the surface of the cathode particle. For example, if the degree of inversion is ~0 on the surface, then Mg²⁺ ions present in the 8*a* sites will be deintercalated first from magnesiated-MgMn₂O₄. Similarly, in a partially inverted surface of a discharged cathode, the Mg²⁺ ions in 16*d* sites that are connected via Hop 3 channels will be extracted as well as those in 8*a* sites connected via Hop 1 and open Hop 2 channels. In the case of a partially inverted surface in a charged-Mn₂O₄ cathode, the Mg²⁺ ions are more likely to first insert into 16*d* channels connected via Hop 5, since Hop 5 exhibits lower Mg migration barriers compared to Hop 1 (Figure 2) at $x_{Mg} \approx 0$.

Since the percolation threshold in the oxide cathode can change with the vacancy concentration during Mg (de)intercalation (Figure 5a), a dynamic change in the degree of inversion during Mg-cycling can cause polarization within the cathode particle. For example, if *i* changes from 0.55 to 0.59 while charging the MgMn₂O₄ cathode, during the following discharge the spinel is percolating only up to Mg_{0.5}Vac_{0.5}Mn₂O₄ (z = 0.5 in Figure 5a) at *i* = 0.59. For further discharge into the structure, that is, from Mg_{0.5}Vac_{0.5}Mn₂O₄ to MgMn₂O₄, a reduction in *i* to 0.55 is necessary, which can lead to hysteresis in the voltages during the charge and discharge cycles. Importantly, the extractable Mg content in stoichiometric MgMn₂O₄ decreases continuously with inversion, reaching values of ~63% (171 mAh/g) and ~17% (46 mAh/g) at *i* = 0.25 and 0.5 (Figure 6c), respectively. Thus, strategies to minimize changes in *i*, during Mg²⁺ cycling, such as cation-doping of Mn to prevent Mn²⁺ generation, should be employed to ensure reversible Mg (de)intercalation.

6. CONCLUSION

Spinels are promising materials in the development of multivalent battery electrodes and solid electrolytes but are prone to antisite disorder in the form of spinel inversion. With the example of two prototypical oxide and sulfide spinels, $MgMn_2O_4$ (cathode) and $MgIn_2S_4$ (solid electrolyte), we demonstrated that inversion can significantly impact both Mgion mobility and electrochemical properties. Using firstprinciples calculations, we analyzed the migration barrier for Mg²⁺ hopping in different local cation arrangements and found that inversion can both open and close select migration pathways on the atomic scale. To quantify the influence of local barrier changes on the macroscopic transport of Mg²⁺ ions, we determined the minimal M-deficiency x in $Mg_xM_{3-x}X_4$ required for percolation. Using a cutoff of 750 and 500 meV for cathodes and solid electrolytes, respectively, we found that the stoichiometric MgMn₂O₄ and MgIn₂S₄ compositions are Mg percolating up to ~55-59% and 44% inversion. Since the degree of inversion in the spinels considered in this work may vary between 20% and 85% depending on the method of preparation,^{29,30,38} a careful calibration of the synthesis conditions is essential to ensure sufficient Mg transport and to reduce the resultant impedance. In addition, spinel inversion can affect the electrochemical properties of cathode materials by changing the phase behavior, average voltage, and extractable capacities. Specifically, we find that the degree of inversion can change dynamically during electrochemical Mg cycling, as indicated by the 0 K phase diagram of the $Mg_{x}Mn_{2}O_{4}$ system and the activation barriers for Mn^{2+} hopping. Notably, even low degrees of inversion (i < 0.4)can detrimentally reduce the extractable capacity in stoichiometric MgMn₂O₄, with an estimated 15% decrease in capacity with every 10% increase in inversion. Thus, spinel inversion can hinder the electrochemical performance of both cathodes and solid electrolytes in MV systems and synthesis efforts must always be made to stabilize the normal spinel structure.

Given that the Mg^{2+} migration barriers over a range of oxide¹¹ and sulfide spinels¹² show similar trends, we expect similar behavior upon inversion in other spinel materials. Finally, the framework developed in this work, particularly the data reported on percolation thresholds and extractable Mg, is readily transferable to other spinels that have potential applications in Li-ion, Na-ion, Ca/Zn-multivalent, and other battery fields.

ASSOCIATED CONTENT

Supporting Information

The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/acs.chemma-ter.7b02820.

X-ray characterization data for $MgIn_2S_4$ and several nudged elastic band calculations in both $MgMn_2O_4$ and $MgIn_2S_4$; formation energies of all Mg-vacancy configurations considered in the $Mg_xMn_2O_4$ system (PDF)

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Notes

The authors declare no competing financial interest.

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