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Tuning Anionic Chemistry To Improve Kinetics of Mg Intercalation

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Supporting Information

ABSTRACT: Rechargeable Mg batteries (RMBs) hold great promise for high energy density in consumer electronics because of the high theoretical volumetric capacity and dendrite-free stripping/plating of Mg. Their development, however, is severely limited by the sluggish diffusion of Mg²⁺ in inorganic cathodes, mainly induced by strong interaction between Mg²⁺ and host anions. Herein, for the first time, we systematically investigate how anionic chemistry (O, S, and Se) affects Mg^{2+} migration in layered MX_2 (M = Ti and V; X = O, S, and Se), by combining theoretical density functional theory (DFT) calculations with electrochemical characterizations. At room temperature, TiSe₂ and VSe₂ achieve much better electrochemical performance than TiS₂ and VS₂, due to the



faster migration of Mg^{2+} in selenide than in sulfide and oxide, as demonstrated by electrochemical kinetic characterizations and DFT calculations. The improved kinetics of selenide can be attributed to three criteria: (i) larger diffusion channel, (ii) weaker interaction between Mg2+ and anion lattice, and (iii) higher electronic conductivity. The three criteria might not only be applicable to layered materials but also be generalizable to materials with other structures, like Chevrel phase, spinel, olivine, etc., which paves the way for the future design and modification of intercalation materials for RMBs.

■ INTRODUCTION

With the increasing demand for green energy, lithium-ion batteries (LIBs) have dominated electric vehicles and consumer electronics.¹⁻³ However, as the energy density of LIBs reaches the limitation of intercalation chemistry, rechargeable metal batteries are becoming promising candidates owing to the high-capacity metal anodes.⁴⁻⁶ Among them, rechargeable Mg batteries (RMBs) stand out, due to the merits of low potential $(-2.37 \text{ V vs } \text{Mg/Mg}^{2+})$, high volumetric energy density (3800 Wh/cm³), rich storage on earth, and, more importantly, highly reversible (close to 100% Coulombic efficiency (CE)) dendrite-free stripping/deposition of Mg in certain electrolytes.⁷⁻⁹ Therefore, RMBs are becoming one of the most promising candidates for application in consumer electronics, in which volumetric energy density and safety are the priority.

In recent years, significant advances have been achieved in RMB electrolytes.^{10–12} The development of RMBs, however, is severely limited by the lack of high-performance cathodes.^{13,14} Inspired by the great success in LIBs, intercalation materials, especially layered materials, are drawing the most attention as cathodes for RMBs because of their chemical stability, high redox potentials, and reversibility.¹⁵⁻¹⁸ However, layered cathodes for RMBs suffer sluggish migration of Mg²⁺ in solid hosts, due to the strong interaction between Mg²⁺ and anions

of host structure.¹⁹⁻²³ According to previous research, the effect of transition metals on the migration of Mg is very weak, $^{19,20,24}\ \mathrm{while}\ \mathrm{the}\ \mathrm{strong}\ \mathrm{interaction}\ \mathrm{between}\ \mathrm{anions}\ \mathrm{and}$ Mg^{2+} is the major contribution to the sluggish kinetics. ^{19,20,25-28} Recent findings demonstrate that oxide cathode materials exhibit extremely sluggish kinetics, 20,29,30 while soft anions (S and Se) have high Mg^{2+} mobility.^{19,31,32} The mechanism is not fully understood. A deep understanding of how anionic chemistry affects Mg²⁺ migration in layered materials would be of paramount significance for the future design of RMB cathode materials.

Herein, for the first time, we systematically investigated how different anionic chemistry (O, S, and Se) affects Mg²⁺ migration in layered MX₂ (M = Ti, V; X = O, S, Se), by combining theoretical calculations with electrochemical characterizations. Three kinetics criteria were proposed based on the correlation between anionic chemistry and intercalation kinetics in layered materials, which might be generalizable to other types of intercalation cathode materials. This work can provide the basis for electrode material design and pave the

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Figure 1. Discharge–charge curves of (a) $TiSe_2$ and (b) VSe_2 for the first three cycles. Cycling performance and Coulombic efficiency (CE) of (c) $TiSe_2$ and (d) VSe_2 with APC electrolyte and metal Mg anode between 0.5 and 2 V at a current density of 5 mA/g under room temperature.

way for exploring intercalation materials with fast kinetics for RMBs.

RESULTS AND DISCUSSION

Electrochemical Performance of MX₂. MX_2 (M = Ti, V; X = O, S, Se) with $P\overline{3}m1$ structure are selected as model cathodes to investigate the impact of anionic chemistry on Mg^{2+} migration kinetics, because they are representative and mostly investigated layered cathodes for RMBs.^{8,13,31,33} Density functional theory calculations demonstrate that TiS₂, TiSe₂, and VSe₂ are thermodynamically stable, while VS₂ is metastable exhibiting moderate energy above hull values of 20 meV/atom. Although common TiO₂ (anatase, brookite, and rutile)³⁴ and VO₂ (rutile)³⁵ are stable at room temperature (RT), layered VO₂ and TiO₂ with $P\overline{3}m1$ structure fall into the unstable range with large energy above hull (>85 meV/atom). Therefore, the discussion about the oxides only limits to calculation.

Based on the thermodynamic stability of MX₂ compounds, TiS₂, TiSe₂, and VSe₂ were synthesized by solid-state reaction with VS₂ by the hydrothermal method. As confirmed by X-ray diffraction (XRD) (Figure S1), all of the patterns of these compounds were indexed in the standard *P*3*m*1 space group, consistent with previous reports.^{36–39} As the particle sizes determine the diffusion length of Mg²⁺, we managed to synthesize the particles of these compounds with a similar size (1–20 μ m), which is also confirmed by scanning electron microscopy (SEM) (Figure S2). Thus, the difference in electrochemical performance between these isostructure materials can be only attributed to their difference in anionic chemistry.

Electrochemical performance of TiS_2 , $TiSe_2$, VS_2 , and VSe_2 cathodes was examined with excess Mg metal as the anode in all phenyl complex (APC) electrolyte between 0.5 and 2 V, using 2032 coin cells at room temperature. Figures 1a,b and S3a,b show the discharge/charge curve for the first three cycles at a current density of 5 mA/g. For sulfide, TiS_2 and VS_2 deliver the negligible capacities (less than 10 mA h/g) with low Coulombic efficiency (CE) (Figure S3a,b), indicating the high migration barriers and Mg²⁺ trapped in TiS_2 and VS_2 .²⁵ In contrast, much higher capacities with a low overpotential and

high CE are achieved for TiSe_2 and VSe_2 (Figure 1a,b). TiSe_2 delivers the first discharge capacity of 127 mA h/g (close to its theoretical capacity: 130 mA h/g) with two plateaus at 1.35–1.4 and 1.15–1.24 V, and 114 mA h/g can be recovered on charge with two plateaus at 1.73–1.75 and 1.57–1.58 V, corresponding to the CE of 89.7%. Similarly, VSe_2 (theoretical capacity: 128 mA h/g) can render a reversible capacity of 115 mA h/g after the first three cycles, with two plateaus on both discharge and charge, indicating similar intercalation and deintercalation processes between TiSe_2 and VSe_2 .

In addition, two plateaus on the discharge and charge processes in TiSe₂ and VSe₂ indicate two steps for Mg²⁺ intercalation into interlayers of TiSe₂ and VSe₂, which are related to the different vacancy location of Mg²⁺ (tetrahedral or octahedral sites).^{38,40} The flat voltage plateaus indicate a two-phase region, induced by the formation of distortion in TiSe₂ and VSe₂ hosts.⁴¹ The distortion is a consequence of changing the ionic character of the bond via magnesium intercalation and electron donation to the host.^{42,43}

The phase change due to distortion in $TiSe_2$ and VSe_2 hosts was also investigated using cyclic voltammetry at a sweep rate of 0.2 mV/s between 0.5 and 2 V (Figure S4). Both $TiSe_2$ and VSe_2 exhibit two cathodic peaks during discharge but only one anodic peak during charge. Two cathodic peaks correspond to two voltage plateaus in discharge curves (Figure 1a,b), whereas two voltage plateaus merge into one peak during charge due to the closed voltages. At the same sweep rate, the current at cathodic and anodic peaks for selenide is higher than that for sulfide (Figure S4), indicating the fast intercalation reaction kinetics.⁴⁴

Cycling stability of TiS_2 , $TiSe_2$, VS_2 , and VSe_2 was evaluated in coin cells with APC electrolyte and an excess Mg negative electrode at 5 mA/g and room temperature (RT) (Figures 1c,d and S3c,d). TiS_2 and VS_2 deliver a low capacity of less than 10 mA h/g (Figure S3c,d). In contrast, $TiSe_2$ and VSe_2 maintain capacities of 80 and 86 mA h/g after 40 cycles, with the Coulombic efficiencies (CEs) above 99.3 and 99.5%, respectively (Figure 1c,d). The slight capacity decay may be associated with the minor dissolution of layered $TiSe_2$ and VSe_2 into electrolytes.



Figure 2. Ex situ XRD of (a) TiSe₂ and (c) VSe₂ for pristine (black), discharged to 0.5 V (red) and charged to 2 V (blue) for the first cycle, respectively. (004) peak shift of (b) TiSe₂ and (d) VSe₂ during discharging and charging processes, respectively.



Figure 3. Morphologies and energy-dispersive spectroscopy (EDS) elemental mapping of discharged $(a-d) TiS_2$, $(e-h) TiSe_2$, $(i-l) VS_2$, and $(m-p) VSe_2$. The scale bars in (a), (e), (i), and (m) represent 2 μ m. All of the samples are discharged to 0.5 V.

Mg²⁺ Diffusion-Controlled MX₂ Cathodes. The reversible intercalation of Mg²⁺ into selenide is confirmed by XRD (Figure 2) and energy-dispersive spectroscopy (EDS, Figure 3). The $P\overline{3}m1$ layered structure of TiSe₂ and VSe₂ maintains after the first discharge and charge processes (Figure 2a,c). During discharge, peaks of XRD shift to the lower angles, indicating that Mg²⁺ is intercalated into the layers of TiSe₂ and VSe₂ (Figure 2b,d). On charge, the peaks shift back to an initial angle but without overlapping the initial peaks, indicating the incomplete deintercalation of Mg^{2+} from $TiSe_2$ and VSe2 with some Mg2+ trapped. EDS was applied to quantify Mg²⁺ intercalation into sulfide and selenide (Figure 3). After discharging to 0.5 V, the amount of Mg^{2+} intercalated into selenide (0.51 Mg/TiSe₂ and 0.48 Mg/VSe₂) is much higher than the amount of Mg^{2+} intercalated into sulfide (0.03 Mg/TiS₂ and 0.05 Mg/VS₂), which is consistent with the discharge capacities.

The overpotential and quasi-equilibrium potential of TiS₂, TiSe₂, VS₂, and VSe₂ during discharge and charge were collected using galvanostatic intermittent titration technique (GITT) (Figures 4a,b and S5). The cells were allowed to relax at the open circuit for 5 h to reach quasi-equilibrium, after each 2 h discharging or charging at 5 mA/g in the first cycle (Figures 4a,b and S5). GITT can provide information on both thermodynamic hysteresis and kinetic polarization.^{45,46} The cumulative capacities of these four compounds are slightly higher than their capacities during the galvanostatic process due to the relaxation process in each discharge/charge process. Apparently, the overpotentials for TiS_2 and VS_2 (Figure S5) are much higher than those for TiSe₂ and VSe₂ (Figure 4a,b), respectively, especially during discharging process, indicating the faster kinetics for Mg²⁺ diffusion in selenide. The quasiequilibrium potentials for TiS₂ and VS₂ are higher than those for TiSe₂ and VSe₂, respectively, due to the stronger



Figure 4. Quasi-equilibrium voltage profile of (a) $TiSe_2$ and (b) VSe_2 obtained from GITT. The cells were allowed to relax for 5 h after every 2 h discharging or charging at 5 mA/g at RT. Representation of the transient voltage of galvanostatic pulse as a function of the square root of time from GITT for (c) $TiSe_2$ and (d) VSe_2 . (e) Nyquist plots for $TiSe_2$ cells at open-circuit voltage states obtained by electrochemical impedance spectroscopy (EIS) tests. The red dashed lines are the fitting curve by using the equivalent circuit, which consists of a resistor (R_1) paralleled with a constant phase element (CPE), and a CPE parallel with a resistor (R_2). (f) Plot of the imaginary resistance versus the inverse square root of angular frequency from EIS for $TiSe_2$. The red dashed line corresponds to the linear fitting with the slope marked out.

interaction between Mg^{2+} and sulfide host, which will be discussed below.

The chemical diffusion coefficient for Mg^{2+} in TiS_2 , $TiSe_2$, VS_2 , and VSe_2 at different charge/discharge states can be obtained from GITT using eq $I^{44,47}$

$$D = \frac{4}{\pi} \left(\frac{IV_{\rm M}}{Z_{\rm A}FS} \right)^2 \left[\frac{\mathrm{d}E(x)}{\mathrm{d}x} / \frac{\mathrm{d}E(t)}{\mathrm{d}\sqrt{t}} \right]^2 \tag{I}$$

where *I* is the applied constant current, $V_{\rm M}$ is the molar volume (112 cm³/mol for TiS₂, 205.79 cm³/mol for TiS₂, 115.07 cm³/mol for VS₂, and 208.86 cm³/mol for VS₂), assumed to be constant during the electrochemical process, $Z_{\rm A}$ is the charge number of electroactive species Mg²⁺, *F* is the Faraday constant (96 486 C/mol), *S* is the electrochemically active area between the electrode material and electrolyte from Brunauer–Emmett–Teller surface area (4585.32 cm²/g for TiS₂, 5706.68 cm²/g for TiSe₂, 16238.16 cm²/g for VS₂, and 4301.45 cm²/g for VSe₂), *E*(*x*) is the equilibrium voltage (i.e., open-circuit voltage) of the working electrode (vs reference electrode) at the composition of Mg_xTiS₂, and *E*(*t*) is the transient voltage response of the working electrode during the constant current interval.

The transient voltage versus the square root of the time of TiSe₂ and VSe₂ is plotted in Figure 4c,d, respectively. Plugging

the slope of the linear fitting into eq I, we can get the diffusion coefficient of Mg^{2+} in TiSe₂ and VSe₂ to be 9.65 × 10⁻²⁰ and 6.17 × 10⁻²⁰ cm²/s, respectively. In contrast, the diffusion coefficients in VS₂ and TiS₂ are 1.04 × 10⁻²³ and 4.20 × 10⁻²³ cm²/s, respectively. The diffusion coefficient of selenide is 3 orders of magnitude higher than that of sulfide, indicating the faster migration of Mg^{2+} in selenide.

Mg²⁺ diffusion coefficient obtained from GITT was also cross-checked using eq II from electrochemical impedance spectroscopy (EIS)

$$D = \frac{1}{2} \left[\left(\frac{V_{\rm M}}{AF\delta} \right) \left(\frac{dE}{dx} \right) \right]^2 \tag{II}$$

where δ is the slope of the Warburg straight line (Ω /s), *A* (cm²) is the effective contact area between the electrolyte and electrode materials, which can be obtained by multiplying *S* in eq I by the loading amount of active materials, and other symbols have the same meaning as they have in eq I.

The impedance data are fitted with the equivalent circuit shown in the inset of Figure 4e. The resistor R_s corresponds to the electrolyte resistance. The resistors R_1 and R_2 paralleled with the constant phase element (CPE) account for the surface reaction resistance of both TiSe₂ cathode and Mg anode. The ion diffusion in the host material is described as the Warburg element (Z_w) in low frequency. The simulated data from the



Figure 5. (a) Crystal structure of layered MX₂ (M = Ti, V; X = O, S, Se). Ti or V atoms (blue) are at the center of octahedra constructed of X atoms (green). Mg^{2+} migrates between the empty interlayer octahedral sites (orange) through the intermediate tetrahedral sites (olivine), which is marked in red dashed line. Calculated energy barrier for the migration of Mg^{2+} in layered structure along the minimum energy path as obtained by first-principles nudged elastic band (NEB) in calculations. (b) $VO_{2^{+}}$ (c) $VS_{2^{+}}$ and (d) $VSe_{2^{-}}$ within the dilute limit of cation insertion.

equivalent circuit well fit the impedance data for layered TiSe₂ at open-circuit voltage states (Figure 4e). Figure 4f shows the plot of the imaginary resistance determined by EIS as a function of the inverse square root of the angular frequency for TiSe₂. Plugging the slope of the linear fitting into eq II, the diffusion coefficient for TiSe₂ is 4.85×10^{-19} cm²/s. Applying the same method, the diffusion coefficients for TiS₂, VS₂, and VSe₂ are 3.10×10^{-22} , 3.20×10^{-22} , and 6.73×10^{-19} cm²/s, respectively. The Mg²⁺ diffusion coefficient for Ti or V sulfides and selenides obtained by the direct modeling of the EIS measurements is slightly higher than that determined by GITT, which has been reported in other cathodes.^{25,48} From both GITT and EIS measurements, the chemical diffusion coefficient for Mg²⁺ in Ti or V selenide are ~3 orders of magnitude higher than that in sulfide.

CV is an efficient tool to investigate the charge-storage kinetics of electrodes. In theory, the peak current (*i*) at a fixed potential obeys a power law relationship with the sweep rate $(v)^{49}$

 $i = av^b$

The value of *b* can be calculated from the slope of $\log(v) - \log(i)$ plots. A *b*-value of 0.5 indicates that the related electrochemical process is totally diffusion-controlled,⁵⁰ while b = 1 means a complete adsorption-controlled behavior, corresponding to a capacitor-like process.^{51,52} As shown in Figure S6a, all of the curves exhibit similar shapes for both cathodic and anodic processes at different sweep rates in the range of 0.1–1 mV/s. The log(*i*) versus log(*v*) plot for the anodic peak at ~1.8 V (versus Mg/Mg²⁺) shows that the *b*-values for TiSe₂ and VSe₂ are 0.58 and 0.57 (Figure S6b), respectively, indicating that the charge storage is mainly controlled by diffusion processes.

 Mg^{2+} Diffusion Mechanism in MX_2 . To reveal the mechanism behind the drastic difference among layered oxide, sulfide, and selenide in electrochemical performance and diffusion coefficient, nudged elastic band (NEB) calculations

were performed to compute the diffusion activation barriers along the migration path for layered MX_2 (M = Ti, V; X = O, S, Se) in the limit of dilute Mg^{2+} insertion. Layered MX_2 $(P\overline{3}m1)$ consists of stacking sequences of the MX₂ slabs, each of which is composed of stacking of close-packed twodimensional triangular lattices of X (Figure 5a). The M atoms occupy octahedrally (tetrahedrally for oxides) coordinated interstitial sites between the X planes. Mg²⁺ diffuses between the stable octahedral sites (tetrahedral sites for oxides) through intermediate tetrahedral sites (octahedral sites for oxides), where the energy exhibits a local minimum before moving to the adjacent octahedral sites (Figures 5a and S7). The migration for Mg^{2+} in layered VO₂ and VS₂ exhibits the activation barriers of 1032 and 593 meV (Figure 5b,c), respectively, which are too high for Mg²⁺ diffusion at room temperature (RT),²⁰ whereas VSe₂ displays the lowest activation barrier of 346 meV (Figure 5d), affording the Mg²⁺ migration at RT. The different profile of oxides from sulfides and selenides is related to the unstable layered structure, while the positions of energetic minima located at \sim 0.2 and \sim 0.8 of the diffusion path are due to some distortion of the layered structure during Mg^{2+} diffusion. Similarly, the migration energy barrier for TiSe₂ is also lower than that for TiS_2 (Figure S8). In addition to oxide, sulfide, and selenide, we also extend our experiments and calculations to telluride (Figure S9). Layered $TiTe_2$ is prepared with the solid-state method (Figure S9a) and delivers a capacity of \sim 86 mA h/g (close to its theoretical capacity of 88 mA h/g) during the first three cycles with lower overpotential than TiSe₂ (Figure S9b). In addition, the energy barrier for the migration of Mg^{2+} in TiTe₂ is calculated to be lower than that of TiSe₂ (Figure S9c), indicating the better diffusion kinetics of Mg²⁺ in TiTe₂.

The trend for decreasing migration barriers in the series of "oxide–sulfide–selenide" compounds with the same structure mainly results from the different anionic chemistry in electronegativity (O > S > Se) and ionic radii (O²⁻ (1.32 Å) $< S^{2-}$ (1.82 Å) $< Se^{2-}$ (1.93 Å)).⁵³ In MX₂ (M = Ti, V; X = O,



Figure 6. Charge rehybridization upon the diffusion of Mg^{2+} between layers of (a) TiO₂, (b) TiS₂, (c) TiSe₂, (d) VO₂, (e) VS₂, and (f) VSe₂ with layer spacing (unit/Å). Areas of charge accumulation are shown in yellow, while depletion is shown in blue. Oxygen atoms are shown as small red spheres, sulfur as small yellow spheres, selenium as small kelly spheres, Ti as large sapphire spheres, V as big green spheres, and Mg at the center as an orange sphere. The isovalue number used for displaying the differential charge density is 0.003.

S, Se), Mg²⁺ diffuses between layers from the stable octahedral sites, crossing through a three-coordinated oxygen (sulfur or selenium) face into the intermediate tetrahedral site and finally to the next stable octahedral site through a symmetric path. The size of Mg²⁺ diffusion channel is dictated by the space between layers. From TiO₂ (VO₂) to TiSe₂ (VSe₂), the spacing between layers increases from 4.99 (4.98) to 6.29 Å (6.37 Å) (Figure 6), indicating the increasing size of diffusion channel, affording faster Mg²⁺ mobility in selenide than that in sulfide and oxide.

The activation energy of Mg^{2+} motion depends on, in particular, the deformability of the electron clouds of the anionic framework.²⁵ The high deformability will easily accommodate the charge brought by intercalated Mg²⁺, which contributes to Mg²⁺ mobility. When Mg²⁺ is intercalated, electrons accompanying Mg2+ are donated to the host. The increased negative charge around the Ti or V ions, in turn, raises the energy of d states relative to that of the X p states, which will increase the ionic character of the M-X bonds (M = Ti, V; X = O, S, Se) due to a reduction in the hybridization between M d states and X p states. Figure 6 shows the charge rehybridization upon the diffusion of Mg²⁺ between layers of MX₂. These charge difference plots were obtained by subtracting the charge densities of both MX₂ structures and an isolated Mg atom from the charge density of Mg_vMX₂. The charge difference plots therefore show how charge redistributes upon the addition of Mg to MX₂. Yellow color signifies an accumulation, while blue signifies a depletion of electron density. As Mg²⁺ migrates through the layers of MX₂, it will drag the electron cloud residing on the coordinating X anions with it. The surrounding M and X ions will therefore be required to rehybridize as the cation moves. While TiX₂ (or VX₂) hosts consist of the same $P\overline{3}m1$ structure, the different degree of rehybridization results from the different anionic chemistry (O, S, and Se). The higher the degree of rehybridization, the lower the deformability.²⁵ Obviously, the charge rehybridization for layered TiX₂ (or VX_2) decreases in the series of "O-S-Se", indicating the increasing deformability in this series, which accounts for the lowest diffusion barriers in TiSe₂ and VSe₂.

The electronic conductivity of the host is another important parameter that determines the mobility of Mg^{2+} . It is proposed that intercalation diffusion rates are dependent on the ease with which the electron cloud in the host can travel through the lattice to accompany the moving cation, which has been demonstrated by Mg^{2+} -intercalated Mo_6S_8 .⁵⁴ With the metallic

electronic structure, the ionic charges of Mo₆S₈ anions are shielded from the highly charged Mg ion by the itinerant electrons of the framework, resulting in the easy migration of ${\rm Mg}^{2+}$ in the lattice. However, when the charge brought by ${\rm Mg}^{2+}$ satisfies the electronic configuration of S atoms and induces a metal-to-semiconductor transition of Mg₂Mo₆S₈, the mobility of Mg^{2+} within $Mg_2Mo_6S_8$ becomes poor. Density of states (DOS), as a major factor in the determination of the electronic conductivity of inorganic materials, was calculated by first-principle calculation (Figure S11). A wide energy gap is observed in the total DOS of layered TiO₂, indicating the nature of semiconductor (Figure S11a). While for TiSe₂ and VSe₂ (Figure S10c,f), the electronic states around the Fermi level (E_f) consist of both 3d orbital of Ti (V) and 4p orbital of Se, indicating that TiSe₂ and VSe₂ are metals with the Fermi energy line entered into the conduction band. In addition, the electronic conductivities of TiS₂, TiSe₂, VS₂, and VSe₂ were experimentally measured to be 1.04×10^5 , 7.84×10^6 , 4.57×10^6 10^6 , and 9.60×10^6 S/m, respectively, which is consistent with the reported electronic conductivity order of selenide > sulfide > oxide.⁵⁵

Based on the discussion above, the layered selenide displays the lowest Mg^{2+} migration barriers and the best electrochemical performance among layered MX_2 (M = Ti, V; X = O, S, Se), which can be summarized into three main factors:^{19,20,54,59} (i) larger sizes of diffusion channel, due to the longer interlayer distance and Mg–Se bond; (ii) less strong interaction between Mg^{2+} and hosts, which can be demonstrated by less charge rehybridization upon Mg^{2+} diffusion between layers; (iii) higher electronic conductivity so that the electron cloud in the host can travel more easily through the lattice to accompany the moving Mg^{2+} .

The three criteria for fast Mg^{2+} migration kinetics developed from layered cathodes with a trigonal structure are also applicable to materials with monoclinic ($P2\overline{1}m$) and triclinic structures ($P\overline{1}$) (Figure S12). NbSe₃ shows much faster kinetics for Mg^{2+} intercalation and higher magnesiation level with a higher CE (Figure S13) than NbS₃ (Figure S14). More broadly, the reported electrochemical performance of cathodes with Chevrel phase,^{25,27} spinel,^{19,60} and olivine^{61,62} structures also supports the three criteria. The high Mg^{2+} mobility can be achieved only when all three criteria are satisfied at the same time. For example, Chevrel phase (Mo_6S_8 or Mo_6Se_8) with large diffusion channel, weak interaction between Mg^{2+} and host lattice, and metallic electronic structure has been proven to be the most successful cathode for RMBs.^{63,64} Sulfide spinel compounds have experimentally demonstrated to achieve high electrochemical performance,^{19,60} while the corresponding oxide spinel compounds with smaller diffusion channel, stronger interaction between Mg²⁺ with host anions, and lower electronic conductivity have not been demonstrated with any electrochemical activity for Mg²⁺ storage.²⁰ Olivine MgFeSiO₄ is calculated with a lower migration barrier of 600 meV⁶⁵ and experimentally achieves high electrochemical performance,⁶² whereas olivine Mg_{0.5}FePO₄ with a barrier of 1025 meV hardly intercalates any Mg²⁺,⁶⁶ due to the stronger interaction between Mg²⁺ with PO₄⁴⁻ than that with SiO₄⁴⁻.

We want to emphasize that synergy among these three criteria is necessary to achieve high Mg^{2+} mobility, i.e., Mg^{2+} mobility would be too low to achieve high magnesiation level if any of these three criteria is not well satisfied. Besides the just mentioned oxide spinel compounds and olivine $Mg_{0.5}FePO_4$, layered MoS_2 with a large interlayer spacing of 6.2 Å and moderate interaction between Mg^{2+} and host anions delivers very low magnesiation degree because of its semiconductor electronic structure.^{67,68} In other words, these three factors can explain most of the experimental findings in intercalation materials for RMBs. Accordingly, the effect of anionic chemistry on the kinetics of Mg^{2+} migration in intercalation compounds might apply to materials not only with a trigonal structure but also with monoclinic and triclinic structures, as well as reported Chevrel phase, spinel, and olivine.

In addition to different structures, the above discussion shows that the three criteria could be extended from chalcogen anions (oxide, sulfide, and selenide) to other anions, like halogen anions (fluoride, chloride, and bromide), hydroxyl, and polyanion. For example, polyanionic compounds might provide good mobility because of a more covalent bonding framework with weaker electrostatic interactions between Mg²⁺ and the host.⁶⁹ Also, some Prussian blue analogues have been demonstrated to intercalate Mg²⁺ reversibly⁷⁰ and benefit from large voids in the open framework structure and weak interaction of Mg²⁺ with the framework constituents.

The three criteria can also provide the following material design principle or modification measures to potentially improve the magnesiation kinetics of intercalation compounds: (i) replacing anion species with less electronegative ones and (ii) incorporating monovalent anions to reduce the electrostatic interaction between intercalant and host and improve the intercalated Mg^{2+} mobility. For example, theoretical calculations predict that partially substituting the O atom with F in one corner of "transition-metal-oxygen" octahedron increases the Mg ion mobility for both VPO₄F and FeSO₄F.^{13,71} In addition, (iii) combining mixed transition-metal ions in oxides with an open-tunnel structure might contribute to charge redistribution and electronic conductivity,⁷² which helps in the Mg²⁺ intercalation.

CONCLUSIONS

In this paper, we systematically studied how anionic chemistry affects Mg intercalation kinetics using layered MX₂ (M = Ti, V; X = O, S, Se) structure as an example system. The mechanism for higher electrochemical performance and lower migration barriers of selenide than sulfide and oxide was revealed by the correlation between anionic chemistry and intercalation kinetics. Three kinetics criteria are proposed to contribute to the better kinetics of layered selenide: (i) larger sizes of diffusion channel; (ii) less strong interaction between Mg²⁺ and host anions; and (iii) higher electronic conductivity. The

criteria derived from layered materials might be generalized to other intercalation compounds, including Chevrel phase, spinel, olivine, etc. This systematic study will provide direction and standard for selecting and designing high-performance cathodes with fast kinetics for RMBs.

ASSOCIATED CONTENT

S Supporting Information

The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/acs.chemma-ter.8b05218.

Experimental details, XRD and SEM of pristine MX_{2} , electrochemical characterizations, calculated energy barriers for the migration of Mg in TiS₂ and TiSe₂, DOS of MX_{2} , and electrochemical performance of NbSe₃ and NbS₃ (PDF)

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Notes

The authors declare no competing financial interest.

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