

Tuning Li-Ion Diffusion in α -LiMn_{1-x}Fe_xPO₄ Nanocrystals by Antisite Defects and Embedded β -Phase for Advanced Li-Ion Batteries

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Supporting Information



ABSTRACT: Olivine-structured LiMn_{1-x}Fe_xPO₄ has become a promising candidate for cathode materials owing to its higher working voltage of 4.1 V and thus larger energy density than that of LiFePO₄, which has been used for electric vehicles batteries with the advantage of high safety but disadvantage of low energy density due to its lower working voltage of 3.4 V. One drawback of LiMn_{1-x}Fe_xPO₄ electrode is its relatively low electronic and Li-ionic conductivity with Li-ion one-dimensional diffusion. Herein, olivine-structured α -LiMn_{0.5}Fe_{0.5}PO₄ nanocrystals were synthesized with optimized Li-ion diffusion channels in LiMn_{1-x}Fe_xPO₄ nanocrystals by inducing high concentrations of Fe²⁺-Li⁺ antisite defects, which showed impressive capacity improvements of approaching 162, 127, 73, and 55 mAh g⁻¹ at 0.1, 10, 50, and 100 C, respectively, and a long-term cycling stability of maintaining about 74% capacity after 1000 cycles at 10 C. By using high-resolution transmission electron microscopy imaging and joint refinement of hard X-ray and neutron powder diffraction patterns, we revealed that the extraordinary high-rate performance could be achieved by suppressing the formation of electrochemically inactive phase (β -LiMn_{1-x}Fe_xPO₄, which is first reported in this work) embedded in α -LiMn_{0.5}Fe_{0.5}PO₄. Because of the coherent orientation relationship between β - and α -phases, the β -phase embedded would impede the Li⁺ diffusion along the [100] and/or [001] directions that was activated by the high density of Fe²⁺-Li⁺ antisite (4.24%) in α -phase. Thus, by optimizing concentrations of Fe²⁺-Li⁺ antisite defects and suppressing β -phase-embedded olivine structure, Li-ion diffusion properties in LiMn_{1-x}Fe_xPO₄ nanocrystals can be tuned by generating new Li⁺ tunneling. These findings may provide insights into the design and generation of other advanced electrode materials with improved rate performance.

KEYWORDS: LiMn_{1-x}Fe_xPO₄, high-rate capabilities, $Fe^{2+}-Li^+$ antisite, β -LiMn_{1-x}Fe_xPO₄, lithium-ion battery

Lithium ion batteries (LIB) have become an effective energy storage system which have been widely used for transportation applications, including hybrid electric vehicles (HEV), plug-in hybrid electric vehicles (PHEV), and electric

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vehicles (EV).¹⁻³ However, the current commercial cathodes fall short of satisfying the needs for high energy density, acceptable rate capability, and excellent stability through rapid (dis)charge over thousands of cycles. For example, the wide application of LiNirCorMnrO2 is limited by its harsh synthesized conditions, security issues, and short working life,³⁻⁵ spinel LiMn₂O₄ suffers from severe capacity fade during electrochemical cycles,^{6,7} and olivine-structured LiFePO₄ also possesses low energy density due to its lower working voltage of 3.4 V.⁸ Recently, olivine-structured α -LiMn_{1-x}Fe_xPO₄ has become a promising candidate for cathode materials because of its higher working voltage of 4.1 V and thus larger energy density than that of LiFePO4 by doping manganese into LiFePO₄.^{9,10} One drawback of α -LiMn_{1-x}Fe_xPO₄ electrode is its relatively low electronic and Li-ionic conductivity,^{11,12} which has been effectively improved by several methods, such as carbon coating to enhance electronic conduction and nanosized active particles of electrodes with shortened Li-ionic diffusion length to enhance Li-ionic conduction.¹³⁻¹⁵ However, both experimental and theoretical studies claimed that the dynamic process of local Li-ionic diffusion could be tuned by structural defects, such as antisite occupancies and substitutions being common in the nanosized electrodes materials.^{16–19} Therefore, we believe that in order to ensure the efficient application of olivine-structured LiMn_{1-r}Fe_rPO₄ in LIB, electrochemical performance of the electrodes could be further enhanced while the fundamental knowledge of underlying Li-ionic transport processes versus defects is revealed.

Many studies have been dedicated to investigate the electrochemical performance of two polymorphs of LiFePO₄, that is, α - and β -LiFePO₄.^{20,21} Compared to the α -phase, the β phase has been proven to possess almost no electrochemical activation,^{21,22} owing to Li-ions kept in captivity to pass the high energy barrier for Li-ion transport through [PO₄] tetrahedra and [FeO₆] octahedra.²³ In our previous work, β -LiFePO₄ was actived by generating various defects to make new Li-ionic diffusion paths with the low energy barriers, however, electrochemical performance of the actived β -LiFePO₄ is still not as good as that of α -LiFePO₄.²⁰ Therefore, the appearance of β -phase usually leads to the performance degradation, and the α -phase has become the main candidate for the applications. In α -LiMn_{1-r}Fe_rPO₄ materials, the Li-ion transport properties can be altered by the Fe²⁺-Li⁺ antisite defects where Fe²⁺ ions reside inside the Li⁺ diffusion channels along the [010] direction. The formation of antisite defects has been reported by X-ray diffraction and transmission electron microscopy (TEM) measurements in α -LiMn_{1-r}Fe_rPO₄^{24,25} which would impede the Li⁺ transport along the diffusion channels.²⁶ On the other hand, it has been predicted by theoretical calculations that the anitsite defects with a high concentration (>2%) will allow the Li-ion diffusion crossover the (one-dimensiona) 1D diffusion channels and tend to shift the 1D to 2D and/or 3D diffusion.^{16,26-28} A number of experimental studies also demonstrated that electronic and Liionic conductivities are essentially two-dimensional in LiFePO₄ single crystals, owing to the presence of antisite defects.^{29,30} Moreover, it is believed that the formation of antisite defects would enhance the rate capabilities by changing the two-phase to single-phase solid solution reaction in olivine-type electrodes.^{5,2}

In this work, we induced high antisite defect concentrations of 5.27% and 4.24% into two kinds (named as 1# and 2# samples) of carbon-coated α -LiMn_{0.5}Fe_{0.5}PO₄ nanoparticles,

respectively, which would activate new Li⁺ tunneling in both of them. Crucial to this study is the suppression of forming electrochemically inactive β -LiMn_{1-x}Fe_xPO₄ nanodomains with size less than 10 nm inside the 100 nm sized α -LiMn_{1-x}Fe_xPO₄ nanocrystals in the 1# electrode but embedded in the 2# electrode, which blocks the Li-ion transport along the [100] and [001] directions, and thus causes the poor electrochemical performance in the 2# electrode. The specific capacities of the superior 1# LiMn_{0.5}Fe_{0.5}PO₄@C are 162, 127, 73, and 55 mAh g^{-1} at 0.1, 10, 50 and 100 C, respectively, which are higher than that of the 2# $LiMn_{0.5}Fe_{0.5}PO_4@C$ (over three times lower than that of 1# sample (15.8 mAh g^{-1}) at 100 C). The 1# electrode also displays excellent cycling lifetime, that is, sustaining about 74% of the total capacity during 1000 cycles at 10 C. Our findings revealed the importance of suppressing the β -phase formation to be embedded in α -phase that is responsible for the enhanced rate capabilities and extended cycling lifetime in olivine-type electrodes, which may shed light on the design of novel materials with desired performance.

Experimental Methods. Synthesis of LiFe_{0.5}Mn_{0.5}PO₄@C. Two kinds of LiMn_{0.5}Fe_{0.5}PO₄@C materials were prepared using the same starting materials but different methods, which were named as 1# LiMn_{0.5}Fe_{0.5}PO₄@C and 2# $LiMn_{0.5}Fe_{0.5}PO_4@C$, respectively. In the typical route, FeSO₄. 7H₂O (AR, 99%), MnSO₄ (AR, 99%), H₃PO₄ (AR, 85% solution), and LiOH·H₂O (AR, 85% solution) were used as starting materials in a molar ratio of 0.5:0.5:1.275:2.7, and ethylene glycol (EG) was applied as solvent. The synthesis process took place under nitrogen atmosphere. For the synthesis of both 1# and 2# LiMn_{0.5}Fe_{0.5}PO₄@C, 270 mmol LiOH·H₂O was dissolved in 111 mL of EG under ultrasonic apparatus to obtain LiOH solution. FeSO₄·7H₂O (50 mmol), 50 mmol MnSO₄, and 0.5 mmol ascorbic acid (as reducing agent to prevent the oxidation process of ferrous) were dissolved in 88 mL of EG to obtain FeSO4 and MnSO4 solution. Before EG was added, MnSO₄ was dissolved in 3 mL of deionized water due to its poor solubility in EG. H₃PO₄ (150 mmol) was dissolved in 22 mL of EG to obtain H_3PO_4 solution. Differently, for 2# LiMn_{0.5}Fe_{0.5}PO₄@C H₃PO₄ solution was slowly added into LiOH solution under magnetic stirring. Five minutes later, FeSO₄ and MnSO₄ solution was slowly added into the mixed solution. For 1# LiMn_{0.5}Fe_{0.5}PO₄@C, H₃PO₄ solution was slowly added into FeSO₄ and MnSO₄ solution under magnetic stirring. Five minutes later, LiOH solution was slowly added into the mixed solution. Then the same procedure was followed for both samples. After stirring for 10 min, the mixture was healed up to 180 °C for 5 h, and then cooled down to room temperature. The obtained precipitates were washed with deionized water and ethanol and dried in a vacuum drying oven at 70 °C for 6 h. Then the $LiMn_{0.5}Fe_{0.5}PO_4$ materials were collected. To achieve carbon coating, LiMn_{0.5}Fe_{0.5}PO₄ materials were mixed with 1.5 wt % of ascorbic acid and 18.5 wt % of glucose dissolved in deionized water and ethanol. After grinding for 30 min, the mixture was heated at 650 °C for 6 h in Ar atmosphere.

Characterization. The crystallographic structures of 1# and 2# LiMn_{0.5}Fe_{0.5}PO₄ were detected by using Bruker D8 Advance diffractometer with Cu K α (λ = 0.15418 nm) in the range of 10–120° at speed of 0.02° s⁻¹. The morphologies of the materials were measured on ZEISS Supra-55 scanning electron microscope. The neutron powder diffraction data were collected on the two materials using the BT-1 high-resolution powder diffractometer at the NIST Center for Neutron



Figure 1. Crystalline structures and electrochemical performance of 1# and 2# $\text{LiMn}_{0.5}\text{Fe}_{0.5}\text{PO}_4@C$ electrodes. (a) XRD patterns of 1# and 2# $\text{LiMn}_{0.5}\text{Fe}_{0.5}\text{PO}_4@C$ electrodes. (b,c) SEM images of 1# and 2# $\text{LiMn}_{0.5}\text{Fe}_{0.5}\text{PO}_4@C$, respectively. (d) Different rate performance of 1# and 2# $\text{LiMn}_{0.5}\text{Fe}_{0.5}\text{PO}_4@C$ electrodes at rates from 0.1 to 100 C, showing the increasing disparity between capacities at elevated rates Both electrodes were cycled between 2.0 and 4.5 V. (e) CV curves of 1# and 2# $\text{LiMn}_{0.5}\text{Fe}_{0.5}\text{PO}_4@C$ under the same scanning rate of 0.2 mV s⁻¹. The small and large half-widths of the CV peaks in 1# and 2# electrodes are marked by cyan and blue colors, respectively.

Research. The high-energy X-ray (hard X-ray) wavelength is 0.011725 nm and the 2D detector is a PerkinElmer amorphous silicon flat panel detector. We took joint refinement method and used Fullprof Suite Toolbar to simulate the neutron powder diffraction data and hard X-rays data at the same time. The high-resolution transmission electron microscopy images and electron diffraction patterns of 1# and 2# LiMn_{0.5}Fe_{0.5}PO₄ were obtained by using a transmission electron microscopy (JEM-3200FS, JEOL) at an acceleration voltage of 300 kV. Fourier transform infrared spectrometer (FTIR) spectra were tested by a PerkinElmer FT-IR spectrometer.

Electrochemical Testing. A coin cell type CR2032 was used for electrochemical test. LiMn_{0.5}Fe_{0.5}PO₄@C, XC-72, and polyvinylidene fluoride (PVDF) were mixed with weight ratio of 5:3:2. The composite was dissolved in N-methylpyrrolidone (NMP) to form electrode slurry. The slurry was coated on aluminum and cut into sequin as working electrode. Lithium metal was used as the counter electrode, and 1 M $LiPF_6$ in a mixture of ethylene chloride (EC), diethyl chloride (DEC), and dimethyl chloride (DMC) was used as an electrolyte with a volume ratio of 1:1:1. All the cells were assembled in a glovebox with water/oxygen content lower than 1 ppm and tested at room temperature. We also packaged the coin cell with conductive black and reduced the content to be 10%, and the percentages of the final samples and PVDF in the working electrodes were 80% and 10%, respectively. The rate performance of XC-72 was also measured between 2.0 and 4.5 V, and the working electrodes were composed of 80%XC-72 and 20% PVDF. The electrochemical data were collected between 2.0

and 4.5 V at different C-rates. The cyclic voltammetry (CV) results were recorded by a CHI electrochemical workstation (CHI604E).

Result and Discussion. The 1# and 2# LiMn_{0.5}Fe_{0.5}PO₄@ C samples, which are synthsized by two different methods (see the Experimental Methods for detailed information), possess different morphologies and electrochemical performance, especially different capacities at high current densities. X-ray diffraction (XRD) patterns of 1# and 2# LiMn_{0.5}Fe_{0.5}PO₄@C are shown in Figure 1a, demonstrating the formation of olivinestructured α -LiMn_{0.5}Fe_{0.5}PO₄ in both of these two samples. The particle size of α -LiMn_{0.5}Fe_{0.5}PO₄ in sample 2# is only a little larger than that in sample 1#, as illustrated by scanning electron microscopy (SEM) and transmission electron microscopy (TEM) images in Figure 1b,c and S1. We also performed Maud refinement on the XRD patterns of the two materials, which shows that the averaged particle sizes of 1# and 2# are 53 and 61 nm, respectively (Figure S2). Figure S3 shows that the specific capacity of 1# LiMn_{0.5}Fe_{0.5}PO₄@C (162 mAh g^{-1}) is a little larger than that of 2# LiMn_{0.5}Fe_{0.5}PO₄@C (155 mAh g^{-1}), when the electrodes were cycled between 2.0 and 4.5 V (vs Li/Li⁺) at 0.1 C. However, such difference between capacities becomes more and more significant when the cycling rate is increased up to 100 C (Figure 1d and S4). It is worth to note that the capacity of 1# $LiMn_{0.5}Fe_{0.5}PO_4@C$ (54.6 mAh g^{-1}) is over three times larger than that of 2# sample (15.8 mAh g^{-1}) at 100 C. Such large disparity between capacities at high rates can still be detected, after using different amount of conductive carbon as shown in Figure S5. The





Figure 2. High concentrations of $Fe^{2+}-Li^+$ antisite defects in 1# and 2# $LiMn_{0.5}Fe_{0.5}PO_4@C$ electrodes (inserted in the plots), as determined by joint Rietveld refinement of (a) neutron powder and (b) hard X-ray diffraction patterns, with black crosses for the experimental data points, red lines for fitted profiles, blue lines for difference, and green bars for Bragg positions.



Figure 3. Microstructures of 2# LiFe_{0.5}Mn_{0.5}PO₄@C. (a) A TEM image showing a single 2# LiFe_{0.5}Mn_{0.5}PO₄@C particle projected along the [010] direction, as indicated by the electron diffraction pattern in (b). (c,d) HRTEM images obtained from the local regions as marked by black boxes c and d in (a), respectively. (e,g) Structural models of α - and β -phases, respectively, and the unit cells of these two phases are marked by black lines, corresponding to the white boxes in (c,d). O, P, Fe/Mn, and Li ions are denoted as red, gray, brown, and green spheres, respectively. (f,h) FFT patterns corresponding to the HRTEM images in (c,d), respectively.

different performance between these two samples is also evidenced by the CV curves in Figure 1e. The CV curves of both electrodes possess two redox peaks at \sim 3.5 and \sim 4.1 V

versus Li/Li⁺, corresponding to the Fe³⁺/Fe²⁺ and Mn³⁺/Mn²⁺, respectively. However, when the scan speed increases from 20 to 100 mV s⁻¹, the redox peak of Mn³⁺/Mn²⁺ becomes absent,

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Figure 4. Origin of the different high-rate performance in $\text{LiFe}_{0.5}\text{Mn}_{0.5}\text{PO}_4$ electrodes. (a) Capacity differences between 1# and 2# $\text{LiFe}_{0.5}\text{Mn}_{0.5}\text{PO}_4@C$ at a high current density. The total capacity of 1# sample was set to be 1. (b–d) Schematic illustration of Li⁺ transport in $\text{LiFe}_{0.5}\text{Mn}_{0.5}\text{PO}_4@C$ with different concentrations of $\text{Fe}^{2+}-\text{Li}^+$ antisite defects. The data inserted in (b–d) exhibit the different antisite concentrations. The red dotted box and the red baffle in (c) represent the inactive Li⁺ and β -phase, respectively.

which is caused by the polarization effect (Figure S6). The halfwidths of the CV peaks of 2# LiMn_{0.5}Fe_{0.5}PO₄@C are wider than that of 1# LiMn_{0.5}Fe_{0.5}PO₄@C. The voltage differences between the Fe^{3+}/Fe^{2+} and Mn^{3+}/Mn^{2+} redox peaks of 2# $LiMn_{0.5}Fe_{0.5}PO_4@C$ are bigger than that of 1# LiMn_{0.5}Fe_{0.5}PO₄@C, which have been marked by blue and cyan colors in Figure 1e, respectively. All of these features indicate that the transmittability of Li-ions in sample 1# is higher than that in sample 2#. Furthermore, both of the two materials demonstrate a long-term cycle stability at 10 C, because about 74% and 49% of total capacity were maintained after 1000 cycles at 10 C in 1# and 2# LiMn_{0.5}Fe_{0.5}PO₄@C (Figure S7), respectively. Therefore, our 1# LiMn_{0.5}Fe_{0.5}PO₄@ C nanoparticles display superior high-rate capability and longterm cycling stability, which have not been demonstrated in other LiMn_{0.5}Fe_{0.5}PO₄-based materials reported in previous studies (Table S1).^{13,14,31-34}

In order to understand the origin of the enhanced electrochemical performance in sample 1#, the concentrations of antisite defects in both samples were accurately determined by jointly refining and simulating the neutron powder and hard X-ray diffraction patterns. Such analysis method allows us to take the advantages of both techniques at the same time, that is, recognition on Fe/Mn by neutron diffraction and identifying precise lattice parameters by hard X-ray diffraction.^{35,36} The refinement results are shown in Figure 2 and Tables S2 and S3. The concentrations of Fe²⁺-Li⁺ antisite defects are 5.27% and 4.24% in 1# and 2# LiMn_{0.5}Fe_{0.5}PO₄@C, respectively, which are higher than the value of <2% in LiMn_{0.5}Fe_{0.5}PO₄@C (si higher than that of 2# LiMn_{0.5}Fe_{0.5}PO₄@C (Figure S8). In

order to verify whether the Mn^{2+} -Li⁺ antisite defects are formed in both samples, we fixed the occupation ratio of Fe and refined the sites of Mn and Li ions. The occupation ratios of Li, Fe, and Mn ions at 4c sites (the center of FeO₆ or MnO₆ octahedra) are 3.88%, 46.08% and 50.04% in sample 2, respectively, which indicates that the Mn²⁺-Li⁺ antisite defect is nonexistent. A similar refinement result was also obtained in 1# LiMn_{0.5}Fe_{0.5}PO₄@C. These results are similar as that reported in LiFe_{1-x}Mn_xPO₄ samples, in which the Mn ions at 4c sites can be barely exchanged with Li ions, owing to the larger ionic radius of Mn²⁺ than that of Fe²⁺ and Li^{+.37}

After identifying the structural differences through neutron and XRD diffractions at the electrode scale, the crystalline structures in the localized areas of these two materials were measured by using high-resolution transmission electron microscopy (HRTEM) imaging. A single nanoparticle of 2# $LiMn_{0.5}Fe_{0.5}PO_4$ (@C sample, as shown in Figure 3a, was used to obtain the electron diffraction pattern (Figure 3b) in which the zone axis can be indexed to be the [010] direction of α -LiMn_{0.5}Fe_{0.5}PO₄. However, the HRTEM images obtained from the local areas c and d in Figure 3a revealed the existence of different structures in α -LiMn_{0.5}Fe_{0.5}PO₄ (Figure 3c,d). In contrast to the olivine structure of α -LiMn_{0.5}Fe_{0.5}PO₄ in region c, as evidenced by the HRTEM image in Figure 3c and the corresponding fast Fourier transform (FFT) pattern in Figure 3f, the HRTEM image (Figure 3d) and corresponding FFT pattern (Figure 3h) display a different structure of β - $LiMn_{1-x}Fe_xPO_4$ in the nearby local region d with a size of about 8 nm. According to the FFT patterns in Figure 3f,h, the crystalline orientation relationship between α -LiMn_{0.5}Fe_{0.5}PO₄ and β -LiMn_{1-x}Fe_xPO₄ can be determined in the following: $[010]_{\alpha}//[011]_{\beta}$, $(100)_{\alpha}//(100)_{\beta}$, $(001)_{\alpha}//(011)_{\beta}$. The lattice

spacing of $(400)_{\alpha}$ and $(002)_{\alpha}$ planes match very well with that of $(200)_{\beta}$ and $(0-22)_{\beta}$, respectively, thus the α - and β -phase cannot be distinguished in the electron diffraction pattern of the whole particle in Figure 3b. It is worth to note that the theoretical misfit along the [100] direction between α - and β -LiFePO₄ phases is about 7.4%.²¹ However, substitution of Fe for Mn can alter the lattice spacing of both phases,^{37,38} which may minimize the disparity between the lattice spacing, and then allow the perfect match to occur. In addition, no peaks of β -phase were observed in the XRD pattern in Figure 1a, which may result from the inadequate amount of β -phase. Similar HRTEM measurements were also performed on single nanoparticles of 1# LiMn_{0.5}Fe_{0.5}PO₄@C sample, but no β phase was found in the local regions of the nanoparticles, as shown by the typical HRTEM images in Figure S9.

The results from multiple characterizations at both the electrode and single-particle scales imply that in both of the highly defective 1# and 2# α -LiMn_{0.5}Fe_{0.5}PO₄ materials, the β - $LiMn_{1-x}Fe_xPO_4$ phase (<10 nm nanodomains) was only embedded in α -LiMn_{0.5}Fe_{0.5}PO₄ phase (about 100 nm nanocrystals) in sample 2#, which possesses worse electrochemical performance than that of sample 1#. The different electrochemical performance of the two materials at high current density were briefly shown in Figure 4a. We proposed one possible mechanism of the Li-ion transport, phase morphology, and their correlation to the different electrochemical performance in these two α -LiMn_{0.5}Fe_{0.5}PO₄ samples at the atomic scale. In a defect-free α -LiMn_{0.5}Fe_{0.5}PO₄ nanoparticle, the Li-ion migrates preferentially along 1D channels, that is, the [010] direction (Figure 4b). However, the concentrations of $\mathrm{Fe}^{2+}\mathrm{-Li}^{\!+}$ antisite defects are high enough in both samples 1 and 2, thus they may generate new tunneling of Li⁺ that are different from the 1D channels, allowing the Liion diffusion along [100] and/or [001] directions with considerable Li-ion diffusivity.^{16,29,30} The transport network made by the new tunneling may be associated with the solidsolution transformation with facile Li-ion migration at elevated rates, which leads to the superior high-rate capabilities in sample 1 (Figure 4d). We hypothesize that a similar electrochemical performance should have been detected in sample 2#, owing the similar level of defect concentrations in these two samples. However, according to the HRTEM results in Figure 3, some β -LiMn_{1-x}Fe_xPO₄ nanodomains were formed in the matrix of 2# α -LiMn_{0.5}Fe_{0.5}PO₄ with the planes of $(100)_{\beta}$ and $(011)_{\beta}$ connecting very well with $(100)_{\alpha}$ and $(001)_{\alpha}$ respectively. Owing to the highly poor Li-ion diffusivity in β phase, the formation of β -phase tends to block parts of Li-ion diffusion paths so as to slow down the Li-ion transport along the [100] and [001] directions in a whole particle of $2\# \alpha$ - $LiMn_{0.5}Fe_{0.5}PO_4$ (Figure 4c), as been evidenced by the large half-widths of CV peaks of sample 2 in Figure 1e. In addition, Li-ions diffuse inefficiently along [010] direction that are blocked by the antisite defects, which also leads to substantially low ionic transport, and then the degradation of high-rate performance in sample 2#.

Conclusions. In summary, we synthesized and compared two kinds of olivine-structured α -LiMn_{0.5}Fe_{0.5}PO₄ electrodes with different electrochemical performance and phase morphologies. Both of these two electrodes contain high concentrations of Fe²⁺-Li⁺ antisite defects. These defects may be responsible for the activation of facile Li-ion transport via new tunneling at high rates. Crucially, by using a suitable synthesis route, β -LiMn_{1-x}Fe_xPO₄ phase with poor Li-ion

diffusivity was only suppressed in local regions of a single nanoparticle of the 1# electrode. Because of the coherent orientation relationship between α - and β -phases, the β -phase would effectively impede the Li-ion migration along the [100] and [001] directions. Therefore, high-rate capabilities were significantly enhanced in the 1# electrode, which cannot be achieved in the 2# electrode prepared by a different synthesis process. Together with the excellent high-rate capability, the 1# electrode shows superior long-term cycling stability compared to other LiMn_{0.5}Fe_{0.5}PO₄-based electrodes. Our findings demonstrate the importance of controlling the formation of electrochemically inactive β -phase in olivine-based electrodes. Both the outstanding performance and the atomistic mechanisms of it may pave the way toward clarifying the requisite synthesis strategies for novel electrodes with optimized electrochemical performance.

ASSOCIATED CONTENT

S Supporting Information

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Materials characterization, electrochemical performance, and crystallographic details of 1# and 2# LiMn_{0.5}Fe_{0.5}PO₄@C (PDF)

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Notes

The authors declare no competing financial interest.

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