

Joint Charge Storage for High-Rate Aqueous Zinc-Manganese Dioxide Batteries

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Aqueous rechargeable zinc-manganese dioxide batteries show great promise for large-scale energy storage due to their use of environmentally friendly, abundant, and rechargeable Zn metal anodes and MnO2 cathodes. In the literature various intercalation and conversion reaction mechanisms in MnO₂ have been reported, but it is not clear how these mechanisms can be simultaneously manipulated to improve the charge storage and transport properties. A systematical study to understand the charge storage mechanisms in a layered δ -MnO₂ cathode is reported. An electrolytedependent reaction mechanism in δ -MnO₂ is identified. Nondiffusion controlled Zn²⁺ intercalation in bulky δ-MnO₂ and control of H⁺ conversion reaction pathways over a wide C-rate charge-discharge range facilitate high rate performance of the δ -MnO₂ cathode without sacrificing the energy density in optimal electrolytes. The Zn-δ-MnO₂ system delivers a discharge capacity of 136.9 mAh g⁻¹ at 20 C and capacity retention of 93% over 4000 cycles with this joint charge storage mechanism. This study opens a new gateway for the design of high-rate electrode materials by manipulating the effective redox reactions in electrode materials for rechargeable batteries.

With the rapid development of renewable energy such as wind and solar, the low cost and high power energy storage technologies become important for the reliability of the electric grid systems to accommodate peak-electricity integration.^[1-6] The electrochemical capacitors, basically with relatively low cost, can store energy in seconds with high power density but low energy

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density.[7-11] Therefore, it is highly desired to pursue alternative energy storage solution enabling both high energy densities and high power densities.[12,13] In recent years, the mild aqueous Zn-based batteries have attracted increasing interest because of the low cost and high capacity of Zn metal anode (819 mAh g⁻¹), suitable redox potential (-0.76 V vs standard hydrogen electrode), and nonflammability, nontoxicity, and high ionic conductivity of aqueous electrolytes.[14] A few cathode materials such as polymorphs of MnO₂,^[15–19] the vanadium-based oxides/ sulfides, [20–22] prussian blue analogs, [23,24] cation-deficient spinel $ZnMn_{2}O_{4}$, [25] NaV₂(PO₄)₃^[26] etc., have been investigated for Zn-based batteries. Among them, the MnO2 cathode has attracted the most interest due to its high capacity (308 mAh g⁻¹), low cost, and the diverse energy storage mechanisms.[16] Different from the widely investigated Li-ion and

Na-ion battery systems, the aqueous Zn-MnO₂ batteries exhibit complicated energy storage mechanisms. Zn²⁺ ion intercalation in $\alpha/\gamma/\delta\text{-MnO}_2$ forming spinel ZnMn₂O₄,^[27,28] Zn-buserite,^[29] layered and/or tunnel Zn_xMnO₂,^[28,30] H⁺ reaction in $\alpha\text{-MnO}_2$ nanofiber forming MnOOH,^[16] and Zn²⁺ and H⁺ cointercalation in akhtenskite-MnO₂ and polymer intercalated layered MnO₂ have been reported in literatures.^[15,17] The different observed storage mechanisms have been attributed to various crystal structure and particle size of cathodes of the starting materials, but have been a subject of significant discussion and debate.^[17] The lack of detailed understanding and control of the mechanisms have limited the development of more efficient and reliable electrode materials for practical applications.^[16]

Among various polymorphs of MnO_2 , the layer $\delta \cdot MnO_2$ polymorph exhibits a large interlayer spacing, which could be appealing for high-rate electrochemical performance. Here, we demonstrate that the charge storage mechanism in rechargeable aqueous $Zn\text{-}MnO_2$ batteries with layered $\delta \cdot MnO_2$ as cathode material can be systematically manipulated to produce optimum high rate electrochemical properties and long cycle life. Combining detailed crystal phase, morphology evolution, and electrochemical kinetic studies in $\delta \cdot MnO_2$ cathode, a joint nondiffusion Zn^{2+} ion intercalation in bulk $\delta \cdot MnO_2$ and H^+ conversion reaction were revealed (**Figure 1**). Such joint charge storage reaction in $\delta \cdot MnO_2$ is mainly associated with the

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Figure 1. Joint nondiffusion controlled Zn²⁺ intercalation and H⁺ conversion reaction mechanism in δ-MnO₂.

electrolyte media for redox reactions but varies with charge–discharge rates. In Zn(TFSI)₂-based electrolyte, nondiffusion controlled Zn²⁺ ion storage mechanism dominates the first step fast charge storage in bulk δ -MnO₂ without significant phase transition, while the diffusion controlled H⁺ conversion reaction dominates the following step reactions. This joint charge storage mechanism leads to the superior high-rate behavior of aqueous Zn-MnO₂ batteries without sacrificing much of the energy density, showing a high reversible capacity of 136.9 mAh g $^{-1}$ at 20 C and 93% capacity retention after 4000 cycles. This study opens a new approach of designing high-rate, high-energy electrode materials by manipulating the effective redox reactions in electrode materials.

Layered δ -MnO₂ was synthesized for rechargeable aqueous Zn batteries using simple hydrothermal method according to previous report. The crystal structure of δ -MnO₂ is made of loosely bound layers of edge-shared MnO₆ located on the (001) plane (Figure 1). The scanning electron microscopy (SEM) image of the prepared δ -MnO₂ shows hierarchical nanoparticle morphologies with an average diameter of around 500 nm, which consists of small nanoflakes as shown in Figure 2a. High-resolution transmission electron microscopy (HRTEM) in Figure 2b reveals the lattice spacing of 0.7 nm for the (001) crystal plane of δ -MnO₂. The crystalline phase of as-prepared sample was further confirmed by X-ray diffraction (XRD), as shown in Figure 2c, which is in good agreement with the structure of layered birnessite δ -MnO₂ ([CPDS: 80-1098). [32]

We first analyzed the redox reactions in layered MnO_2 cathode by identifying the kinetic parameters of δ - MnO_2 . Figure 3a and Figure S1 of the Supporting Information show the cyclic voltammetry (CV) scanning curves of δ - MnO_2 from 0.1 to 10 mV s⁻¹ in 1 M $Zn(TFSI)_2$ with 0.1 M $Mn(TFSI)_2$

electrolyte. Pre-addition of manganese salt in electrolyte was used to suppress Mn²⁺ dissolution from MnO₂ electrodes.^[16] Two pairs of redox peaks denoted as peak 1, 2, 3, 4 were observed in δ-MnO₂ during charge and discharge, indicating two potential types of redox reactions. The current (i) of CV scanning is assumed to exhibit a dependency of the sweep rate (v): $i = av^b$, [33,34] where a and b are variable values, and the value b falls in a range of 0.5-1, depending on the charge storage kinetics in δ -MnO₂. The value b of 0.5 suggests that the charge storage in electrode material is controlled by solid-state diffusion of active ions, whereas value b of 1 indicates that the current is surface controlled electrochemical capacitive process in electrode material.[34] (Note that the zinc-based aqueous electrolytes show a high ionic conductivity of > 40 mS cm⁻¹ (Table S1, Supporting Information), indicating fast diffusion kinetics of active ions in the electrolyte. The diffusion of active ions in liquid electrolyte is therefore not considered as the ratelimiting step for the overall electrochemical redox reactions in solid δ -MnO₂ electrode. We plotted the dependence curves between $\log(i)$ with $\log(v)$ for the observed redox peaks, respectively (Figure 3b). The fitted curves for four peaks exhibit two different regions. In the scanning rate range of 0.1–1 mV s⁻¹, the fitting b value is approaching 1 (0.87, 0.81, 0.90, and 0.82 for peak 1 to 4, respectively), which indicates that the kinetics is fast and mainly nondiffusion controlled. In contrast, the value of b decreased to 0.76, 0.73, and 0.64 for peak 1, 3, and 4, respectively in the scanning rate range of 2-10 mV s⁻¹ (Note that the fitting curve for peak 2 is not shown because the peak shifted out of the voltage range). This indicates that the electrochemical charge storage in δ -MnO₂ cathode is contributed by both nondiffusion controlled and solid-state ion diffusion controlled reactions within a wide scanning rate range, where

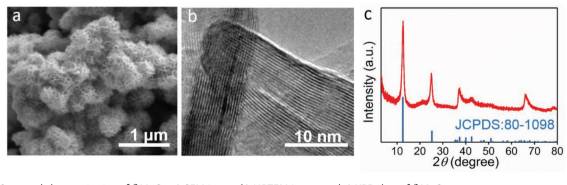


Figure 2. Structural characterization of δ -MnO₂. a) SEM image, b) HRTEM image, and c) XRD data of δ -MnO₂.

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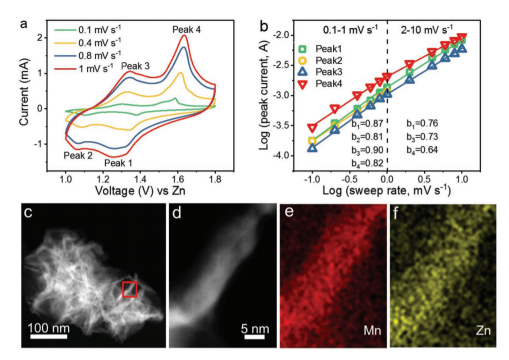


Figure 3. Kinetic analysis of the electrochemical reactions in δ -MnO₂. a) CV curves of Zn-MnO₂ batteries at different scan rates. b) b-value determination at each peak according to $i = a\nu^b$ (i is current; ν is sweep rate; a, b are variable values). Note the value of b_n (n = 1, 2, 3, 4) indicates the b value of peak n in the CV curves in (a) and Figure S1 (Supporting Information). c–f) STEM-HAADF images (c,d) and STEM-EDS mappings of the elemental distributions of e) Mn and f) Zn for the δ -MnO₂ electrode discharged to 1.43 V in the second cycle in the selected electrode area in Figure 3c. The red rectangular region in Figure 3c is magnified in Figure 3d. Electrolyte: 1 M Zn(TFSI)₂ electrolyte with 0.1 M Mn(TFSI)₂ additive in water.

the nondiffusion controlled charge storage contributes more effective capacity with a scanning rate lower than 1 mV s^{-1} .

To further quantify the capacity contribution, the correlation between the normalized capacity and $v^{-1/2}$ for δ -MnO₂ was established from sweep rate of 0.2 to 10 mV s⁻¹ in Figure S2a of the Supporting Information, where the nondiffusion controlled capacity remains constant, and capacity resulted by solid-state diffusion controlled process varies with $v^{-1/2}$.[10] In excellent agreement of the results in Figure 3b, two regions of capacity for δ -MnO₂ were observed. With ν < 1 mV s⁻¹, the normalized charge storage capacity decreases slightly with increasing the sweep rate, whereas the dependence of capacity on sweep rate becomes stronger at v > 1 mV s⁻¹. The critical sweep rate separating diffusion controlled capacity and nondiffusion controlled capacity is likely associated with the intrinsic charge transport properties in the electrode material and used electrolytes. We further quantified the nondiffusion controlled capacity in δ -MnO₂ in the inset of Figure S2a of the Supporting Information, with 76%, 82%, 86%, 85%, and 88% at the sweep rate of 0.2, 0.4, 0.6, 0.8, and 1 mV s^{-1} respectively. In addition, the redox peaks exhibit small voltage offset lower than 50 mV at sweep rate < 1 mV s⁻¹ (Figure S2b, Supporting Information), and this is similar to surface redox reactions with fast charge storage kinetics.^[33] The nondiffusion controlled capacity should be attributed to the "pseudocapacitance"-like charge storage in δ-MnO₂ cathode in aqueous electrolyte. The STEM-EDS mappings of discharged δ-MnO₂ cathode (1.43 V) in Figure 3c-f revealed the uniform distribution of elemental Zn with the same distribution of Mn in the nanoflakes, indicating the charge storage

of Zn^{2+} ion in the δ -MnO $_2$ cathode. Surprisingly, the nondiffusion controlled "pseudocapacitive"-like Zn^{2+} ion storage in δ -MnO $_2$ has a high specific capacity (238.8 mAh g $^{-1}$ at 0.2 C, 78% of theoretical capacity for $1e^-$ transfer). The high non-diffusion controlled capacity is in sharp contrast of the electrochemical capacitor that is commonly associated with the surface or near-surface redox reaction, [35] indicating Zn^{2+} insertion into the bulk structure of δ -MnO $_2$ instead of on the outer surface (Surface area: 34.08 m 2 g $^{-1}$; Figure S3, Supporting Information). The nondiffusion controlled Zn^{2+} intercalation could be related to the 2D crystal network of δ -MnO $_2$ that allows a fast transport pathway with little crystal phase change (To be discussed later). [33] Such phenomenon has also been observed for Li $^+$ intercalation pseudocapacitance in the bulk of 2D materials, such as T-Nb $_2$ O $_5$. [33]

In addition, the anion species of zinc salt or manganese salt strongly affect the ion association properties in solutions. We find that the charge storage mechanism in layer δ -MnO₂ varies with electrolyte media for the redox relations (Figure S4, Supporting Information). The quantification of charge storage kinetic in δ -MnO₂ in 1 m ZnSO₄ and 0.1 m MnSO₄ electrolyte was performed in Figure S5 (Supporting Information). Similar mixed charge storage were observed in δ -MnO₂. However, the separation of two different charge storage processes is much more obvious in ZnSO₄-based electrolyte. At sweep rate < 1 mV s⁻¹, the charge storage kinetics in δ -MnO₂ is dominated by nondiffusion controlled charge storage with b value close to 1 (capacity contribution: \approx 95%). While at sweep rate > 1 mV s⁻¹, the b approaches a value of 0.5 (Figure S5, Supporting Information).

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Furthermore, it was found that the joint charge storage mechanism in δ -MnO₂ is less associated with the morphology of δ -MnO₂ rather than the electrolyte. Similar charge storage mechanism was observed for the δ -MnO₂ with different particle size and surface area (Figure S6, Supporting Information). Zinc salts with a bulky anion such as TFSI- and TF-, seems to lead obviously to more nondiffusion controlled charge storage in δ-MnO₂ than that in the ZnSO₄-based electrolyte in the higher sweep rate range (Figures S7, S8, Supporting Information). δ-MnO₂ exhibits basically similar joint charge storage mechanism in electrolytes with different pH values as shown in Figure S9 (Supporting Information). Nevertheless, the pH value of electrolyte shows some modification on the joint charge storage mechanism in δ -MnO₂. Lower pH value seems to favor the nondiffusion controlled Zn^{2+} storage capacity in δ -MnO₂ under high sweep rate (Figure S9, Supporting Information). This could be because that the increasing of H+ concentration frees some Zn²⁺ cation from coordinating with OH⁻ in water solution and facilitates its intercalation.[36]

To further understand the different charge storage mechanism, the structural evolution of δ -MnO₂ was carefully investigated during cycling. **Figure 4**a shows the charge and discharge curve of δ -MnO₂ and the selected charge–discharge status for ex-situ XRD measurements in the second cycle at 0.2 C (equals to < 1 mV s⁻¹) in 1 m Zn(TFSI)₂ electrolyte. Note that the ex-situ XRD for δ -MnO₂ electrodes were tested after disassembling the cell without rising by water. Interestingly, no new XRD peak showed up beside the peaks from Zn(TFSI)₂ salts (Figure S10,

Supporting Information) in electrolyte before discharging to 1.43 V (Status I), indicating no visible crystal structure change of δ -MnO₂. The capacity before 1.43 V is, therefore, mainly produced by the nondiffusion controlled "pseudocapacitive"like Zn^{2+} intercalation in the bulk δ -MnO₂ without a phase change, [33] which is consistent with the more closer b value to 1 for the reduction peak 1 in Figure 3b. Along with the discharge process, a set of new peaks (marked by yellow solid line) appear from the discharge voltage of 1.22 V (Status II), and the intensity continuously increases until the end of discharge to 1 V (Status IV) along with the gradual reduction in the peak intensity of δ -MnO₂ phase (marked by gray dashed line). The set of newly appeared peaks (marked by yellow solid line) likely belongs to Zn(TFSI)₂[Zn(OH)₂]₃.xH₂O discharge product, which is an analogy of $ZnSO_4[Zn(OH)_2]_3.nH_2O$ (n = 3, 4 etc.) that was observed in Zn-MnO₂ batteries in ZnSO₄-based electrolytes (Figure S11, Supporting Information).^[16] The presence of $Zn(TFSI)_2[Zn(OH)_2]_3.xH_2O$ in the δ -MnO₂ cathode indicates the H⁺ conversion reaction with δ -MnO₂ to form MnOOH during further discharge process, which is evidenced by the XRD patterns of MnOOH as shown in Figure S12 (Supporting Information). Meanwhile, the OH- reaction with the Zn2+ ion dissolved from Zn anode and Zn(TFSI)2 aqueous electrolyte to form the accompanied discharge compound of $Zn(TFSI)_2[Zn(OH)_2]_3.xH_2O$ on δ -MnO₂ electrode along with the redox reaction of $MnO_2 + H \leftrightarrow MnOOH$, which is also evidenced by the SEM image of newly formed flakes in the δ -MnO₂ electrodes in discharged status (Figure S13a,

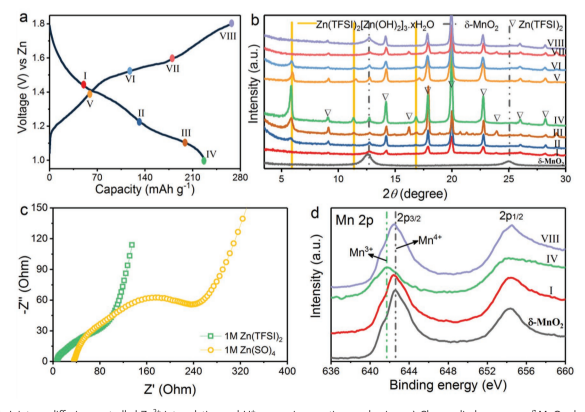


Figure 4. Joint nondiffusion controlled Zn^{2+} intercalation and H^+ conversion reaction mechanism. a) Charge–discharge curve δ -MnO₂ electrodes in the second cycle at 0.2 C. b) Ex-situ XRD patterns of δ -MnO₂ electrodes at selected states in (a). c) EIS analysis of Zn- δ -MnO₂ cells at discharged state. d) XPS spectra for Mn 2p at the selected states in (a). Electrolyte: 1 M Zn(TFSI)₂ electrolyte with 0.1 M Mn(TFSI)₂ additive in water.

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Supporting Information). We further compared the XRD patterns of δ-MnO₂ electrodes after rising with DI-water at the end discharge state. The Zn(TFSI)₂[Zn(OH)₂]₃.xH₂O peaks disappeared after complete rinse with water, indicating that formed $Zn(TFSI)_2[Zn(OH)_2]_3.xH_2O$ in δ -MnO₂ cathode is washable by water (Figure S11a, Supporting Information), which was confirmed by the disappearance of flake-like of $Zn(TFSI)_2[Zn(OH)_2]_3.xH_2O$ in the SEM image of δ -MnO₂ cathode at the discharge state of 1 V after DI-water rinse (Figure S13b, Supporting Information). In contrast, the $ZnSO_4[Zn(OH)_2]_3.nH_2O$ formed in δ -MnO₂ cathode in $ZnSO_4$ based electrolyte is not soluble in water and remains without change in XRD patterns before and after rinsing (Figure S11b, Supporting Information). Therefore, δ -MnO₂ electrode in Zn(TFSI)2-based electrolyte leads to both low ohm resistance and interfacial charge transfer -3 times lower than that in ZnSO₄-based electrolyte (Figure 4c), which would further benefit the rate capability of δ -MnO₂ cathode.

The charge process is highly reversible in the charge–discharge process. As charge voltage increased, the peak intensity of $Zn(TFSI)_2[Zn(OH)_2]_3.xH_2O$ gradually decreased (Status V, VI), and reversibly disappeared at the charge voltage of 1.6 V (Status VII). Between the charges voltages of 1.6 and 1.8 V, the XRD peaks did not change significantly for δ -MnO $_2$ phase except for the gradual intensity increase. This indicates that the capacity observed in this high charge voltage range is from the extraction of nondiffusion controlled Zn^{2+} ions storage in

the bulky structure of δ -MnO₂ phase with no obvious crystal structure change. No large flake-like Zn(TFSI)₂[Zn(OH)₂]₃.xH₂O was observed in δ -MnO₂ electrode charge to 1.8 V (Figure S13d, Supporting Information), confirming the reversible formation and decomposition of Zn(TFSI)₂[Zn(OH)₂]₃.xH₂O during cycling.

The X-ray photoelectron spectroscopy (XPS) analysis for Mn $2p_{3/2}$ peaks in δ -MnO $_2$ electrodes reveals the valance status change of Mn during charge and discharge in Figure 4d. The Mn2 $p_{3/2}$ peaks for δ -MnO $_2$ at discharge status I exhibit a dominated peak at 642.5 eV with a shoulder at 641.6 eV, which are ascribed to the existence of Mn⁴⁺ and Mn³⁺, respectively, in δ -MnO $_2$ electrode after initial Zn²⁺ intercalation. The Mn $2p_{3/2}$ peak significantly shifted to the lower binding energy with a dominated reduced peak of Mn³⁺ (641.6 eV) at the end of discharge (state IV);^[37] and it changed back to the original binding energy in the charge process (state VIII) along with the Mn oxidized to its initial state. The reversible valence status change for Mn 2p further confirms the reversible joint charge storage in δ -MnO $_2$ structure.

TEM images were further used to investigate the morphological and structural evolution of δ -MnO₂ electrode at lattice-scale during the charge–discharge process in 1 M Zn(TFSI)₂-based electrolyte in the first cycle. **Figure 5**a–c shows the TEM/HRTEM images of δ -MnO₂ electrode at the discharge state of 1 V. It was found that the pristine flake-like δ -MnO₂ particles were transformed into two kinds of crystal morphologies at

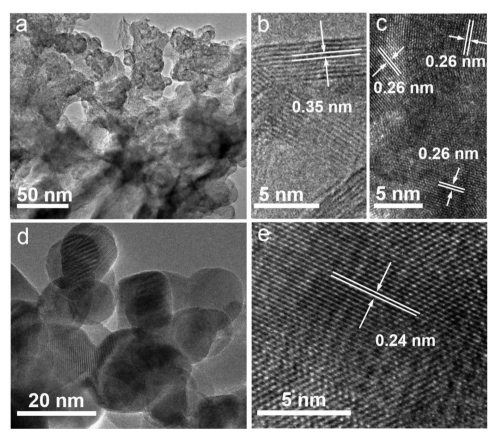


Figure 5. TEM images of δ-MnO₂ electrodes during the electrochemical process. a–c) MnO₂ electrodes discharged to 1 V. d, e) MnO₂ electrodes charged back to 1.8 V. (first cycle). Electrolyte: 1 $_{\rm M}$ Zn(TFSI)₂ electrolyte with 0.1 $_{\rm M}$ Mn(TFSI)₂ additive in water.



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higher magnification. The nanoflake like morphology with lattice distance of 0.35 nm corresponds to the (0 0 2) plane of δ-MnO₂ (Figure 5b). This well-preserved nanoflakes further confirm the nondiffusion controlled "pseudocapacitance"-like Zn²⁺ intercalation in bulk δ -MnO₂ without phase change, [33] which is consistent with ex-situ XRD results in Figure 4b. The other main morphology is the ≈ 5 nm nanoparticle aggregates with some disorder in crystals as demonstrated in the HRTEM of Figure 5c. The lattice distance of 0.26 nm in nanoparticles aggregates corresponds to the (0 4 0) planes of MnOOH phase, as evidenced by the XRD in Figure S12 (Supporting Information). The appearance of MnOOH confirms the proton reactions with δ -MnO₂. The two different morphologies after first discharge could modify the reaction kinetics for the Zn2+ intercalation and phase change conversion reaction with H+ in the following cycles due to the reduced diffusion length in nanoparticle for the reaction between MnO2 and H+ to form MnOOH. Furthermore, the morphology was converted into the crystal particle at the fully charged state of 1.8 V (Figure 5d). The HRTEM images indicate that the lattice distance of 0.24 nm is the (-1 1 1) plane of the original δ -MnO₂ electrode (Figure 5e), revealing the reversible crystal structural change of δ -MnO₂.

The structural and chemical evolution of δ -MnO₂ during charge and discharge process confirms the charge storage mechanisms in δ -MnO₂ and reveals the reaction sequence between nondiffusion controlled Zn^{2+} intercalation and H^{+} conversion reaction in δ -MnO₂. The reaction mechanism and charge storage sequence are mainly manipulated by the electrolyte media, and also depends on charge-discharge current rates. During the discharge process, nondiffusion controlled Zn^{2+} intercalation in bulk δ -MnO₂ first occurs and dominates the high voltage discharge process, and the H+ conversion reaction with MnO₂ dominantly takes place in the further discharge process under a medium charge rate as illustrated in Figure 1. Note that the specific charge storage process could vary under high C-rates. For the charge process, it is highly reversible with H+ conversion reaction at the early charge stage and nondiffusion controlled Zn²⁺ intercalation at the high voltage charge process. This joint charge storage in δ -MnO₂ with appropriate electrolyte media, such as Zn(TFSI)2-based electrolyte, facilitates fast charge storage kinetics and accounts for the observed high rate capability of δ -MnO₂ in the appropriate electrolyte as discussed in Figure 3.

The overall reactions for $Zn-\delta-MnO_2$ batteries in $Zn(TFSI)_2$ -based electrolyte can be the following processes during charge and discharge

Cathode:

$$MnO_2 + xZn^{2+} + 2xe^- \leftrightarrow Zn_xMnO_2$$
 (Nondiffusion – controlled reaction) (1)

$$H_2O \leftrightarrow H^+ + OH^-$$
 (2)

$$MnO_2 + H^+ + e^- \leftrightarrow MnOOH(Diffusion - controlled reaction)$$
 (3)

$$\frac{1}{2}Zn^{2^{+}} + OH^{-} + \frac{1}{6}Zn(TFSI)_{2} + \frac{\varkappa}{6}H_{2}O \leftrightarrow \frac{1}{6}Zn(TFSI)_{2}[Zn(OH)_{2}]_{3} \cdot \varkappa H_{2}O$$

Anode:

$$\left(\frac{1}{2} + x\right) Zn \leftrightarrow \left(\frac{1}{2} + x\right) Zn^{2+} + (1 + 2x)e^{-}$$
(5)

The electrochemical properties of layered δ -MnO₂ were then evaluated in coin cells using δ -MnO₂ as the cathode and Zn metal as the anode in different electrolytes. δ -MnO₂ cathode shows excellent rate performance, demonstrating high discharge capacities of 238, 220, 171, and 138 mAh g-1 at 0.2 C, 1 C, 5 C, and 20 C, respectively, in 1 M Zn(TFSI)2 with 0.1 M Mn(TFSI)₂ electrolyte (Figure 6a). (Note that the relatively low current rate, e.g. 0.2 C, will cause more of oxidation of preaddition of Mn2+ additive in electrolyte than the dissolution of Mn²⁺ from electrode during the slow charging process, leading to lower Coulombic efficiency at low current rate). The chargedischarge capacity exhibits less obvious decrease under 1 C rate, showing that the ion-diffusion is not a rate-limiting step for charge storage. In contrast, the rate capability of δ -MnO₂ cathode deteriorates a lot in ZnSO₄-based electrolyte with increased overpotential and reduced capacity utilization even though the same δ -MnO₂ cathode was used (Figure S14, Supporting Information). The improved performance of δ -MnO₂ could be likely ascribed to excellent kinetics of nondiffusion controlled Zn2+ ion intercalation in the bulk of the nanoflakelike δ -MnO₂ electrodes during charge and discharge processes in Zn(TFSI)2-based electrolyte in a wide C rate range. This is consistent with the kinetics of charge storage in layered δ -MnO₂

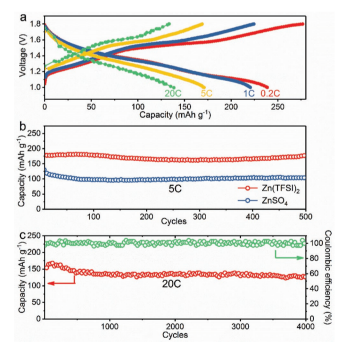


Figure 6. The electrochemical performance of Zn- δ -MnO $_2$ batteries. a) Rate performance of Zn- δ -MnO $_2$ battery in 1 M Zn(TFSI) $_2$ with 0.1 M Mn(TFSI) $_2$ additive electrolyte. b) Comparison of the cycling performance of δ -MnO $_2$ electrodes at 5 C in 1 M Zn(TFSI) $_2$ with 0.1 M Mn(TFSI) $_2$ additive and 1 M ZnSO $_4$ with 0.1 M MnSO $_4$ electrolyte, respectively. c) The long cycle performance of Zn- δ -MnO $_2$ battery at 20 C in 1 M Zn(TFSI) $_2$ with 0.1 M Mn(TFSI) $_2$ additive electrolyte.



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as discussed in Figure 3 and the low charge transfer impedance in Figure 4c. The manipulation of redox reactions in δ -MnO₂ cathode via reaction media provides a new gateway to design high rate and high energy batteries systems beyond the modification of electrode material itself. In addition, the Zn-MnO2 battery also demonstrates higher cycling stability with higher capacity in Zn(TFSI)₂-based electrolyte than that in ZnSO₄ electrolyte at 5 C (Figure 6b). Excellent long-cycle stability at high operation rate of 20 C was shown, with a discharge capacity of 136.9 mAh g⁻¹ and high-capacity retention of 93% over 4000 cycles (Figure 6c). The combination of superior high-rate performance, long cycle life, environmentally friendly and low cost of electrode materials in Zn-MnO2 batteries enables an excellent alternative solution for energy storage. In addition, the Zn anode also exhibits good stability in the Zn(TFSI)2-based electrolyte with a stabilized Coulombic efficiency of >99% over thousand cycles under shallow plating/stripping as shown in Figure S15 (Supporting Information). However, approaches to improve the stability and reversibility of Zn anode under higher utilization is still critical for the practical application.

In summary, we demonstrate a new charge storage mechanism in δ-MnO₂ for designing high-rate performance Zn-MnO₂ batteries. A joint nondiffusion controlled Zn²⁺ intercalation in bulk δ-MnO₂ and H⁺ conversion reaction pathway was identified in δ -MnO₂ cathode during charge and discharge, in which the charge storage mechanisms could be tailored through electrolytes, especially via the anion of zinc salt. Fast reaction kinetics not limited by ion diffusion was realized for Zn-δ-MnO₂ batteries in Zn(TFSI)₂-based mild aqueous electrolyte in a wide C-rate range. This mixed energy storage mechanism contributes to superior high rate electrochemical performance of 136.9 mAh g⁻¹ discharge capacity at 20 C and 93% capacity retention after 4000 cycles. This study opens a new gateway to the design of high-rate electrode materials by manipulating the effective redox reactions in electrode materials for rechargeable batteries.

Experimental Section

Material Synthesis: δ -MnO $_2$ was synthesized by hydrothermal method. KMnO $_4$ and MnSO $_4$ with a molar ratio of 6:1 were put into a beaker and stirred to make the reactant totally dissolved. The solution was put into a Teflon contained autoclave and heated at 160 °C for 12h. The solution was then cooled to room temperature. δ -MnO $_2$ powder was obtained through the centrifuge, washed with deionized water three times, and dried at 60 °C overnight.

Characterizations: XRD measurements were performed using a Rigaku Miniflex II diffractometer with Cu K α radiation ($\lambda=1.5406$ Å). SEM was obtained on Helium ion microscope. Transition electron microscopy (TEM) images were conducted on FEI Titan at 300 kV. XPS measurements were performed using a Physical Electronics Quantera Scanning X-ray Microprobe. The specific surface area of δ -MnO $_2$ powder was determined by nitrogen adsorption curve, which was conducted on the surface area analyzer (Quantachrome, USA) and estimated by BET (Brunauer–Emmett–Teller) method.

Electrochemical Measurements: δ-MnO₂ was mixed with acetylene black and PVDF with a weight ratio of 6:3:1 in NMP solvent. The slurry was spread on carbon paper current collector and heated at 80 °C under vacuum condition overnight. The electrode was punched into 0.5 inch diameter disc and mass loading was around 1 mg cm⁻². 2023 coin cell was assembled to evaluate the electrochemical performance of

 $\delta\text{-MnO}_2$ with Zn foil (Alfa Aesar) as counter electrode and glass fiber as the separator. The galvanostatic charge/discharge was carried out in the voltage range of 1–1.8 V on a LANHER battery tester (Wuhan). Cyclic voltammetry (CV) measurements were conducted using CHI660E. Electrochemical impedance spectra (EIS) were performed on Bio-Logic Instruments (VSP) using a three-electrode cell configuration with δ -MnO $_2$ as the working electrode and Zn metal as both reference and counter electrodes.

Supporting Information

Supporting Information is available from the Wiley Online Library or from the author.

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Conflict of Interest

The authors declare no conflict of interest.

Keywords

Aqueous $Zn\text{-}MnO_2$ batteries, high-rate batteries, joint charge storage, battery reaction mechanisms

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- [1] B. Dunn, H. Kamath, J.-M. Tarascon, Science 2011, 334, 928.
- [2] S. Chu, A. Majumdar, Nature 2012, 488, 294.
- [3] J. Liu, J. G. Zhang, Z. Yang, J. P. Lemmon, C. Imhoff, G. L. Graff, L. Li, J. Hu, C. Wang, J. Xiao, Adv. Funct. Mater. 2013, 23, 929.
- [4] D. Larcher, J.-M. Tarascon, Nat. Chem. 2015, 7, 19.
- [5] S. Chu, Y. Cui, N. Liu, Nat. Mater. 2017, 16, 16.
- [6] J. Rugolo, M. J. Aziz, Energy Environ. Sci. 2012, 5, 7151.
- [7] W. Wei, X. Cui, W. Chen, D. G. Ivey, Chem. Soc. Rev. 2011, 40, 1697.
- [8] L. L. Zhang, X. Zhao, Chem. Soc. Rev. 2009, 38, 2520.
- [9] P. Simon, Y. Gogotsi, Nat. Mater. 2008, 7, 845.
- [10] V. Augustyn, P. Simon, B. Dunn, Energy Environ. Sci. 2014, 7, 1597.
- [11] S.-L. Kuo, N.-L. Wu, J. Electrochem. Soc. 2006, 153, A1317.
- [12] Z. Fan, J. Yan, T. Wei, L. Zhi, G. Ning, T. Li, F. Wei, Adv. Funct. Mater. 2011, 21, 2366.
- [13] Z. S. Wu, K. Parvez, X. Feng, K. Müllen, Nat. Commun. 2013, 4, 2487
- [14] M. Song, H. Tan, D. Chao, H. J. Fan, Adv. Funct. Mater. 2018, 28, 1802564.



www.advmat.de

- [15] J. Huang, Z. Wang, M. Hou, X. Dong, Y. Liu, Y. Wang, Y. Xia, Nat. Commun. 2018, 9, 2906.
- [16] H. Pan, Y. Shao, P. Yan, Y. Cheng, K. S. Han, Z. Nie, C. Wang, J. Yang, X. Li, P. Bhattacharya, *Nat. Energy* **2016**, *1*, 16039.
- [17] W. Sun, F. Wang, S. Hou, C. Yang, X. Fan, Z. Ma, T. Gao, F. Han, R. Hu, M. Zhu, J. Am. Chem. Soc. 2017, 139, 9775.
- [18] N. Zhang, F. Cheng, J. Liu, L. Wang, X. Long, X. Liu, F. Li, J. Chen, Nat. Commun. 2017, 8, 405.
- [19] S.-D. Han, S. Kim, D. Li, V. Petkov, H. D. Yoo, P. J. Phillips, H. Wang, J. J. Kim, K. L. More, B. Key, *Chem. Mater.* **2017**, *29*, 4874
- [20] J. Ding, Z. Du, L. Gu, B. Li, L. Wang, S. Wang, Y. Gong, S. Yang, Adv. Mater. 2018, 30, 1800762.
- [21] D. Kundu, B. D. Adams, V. Duffort, S. H. Vajargah, L. F. Nazar, *Nat. Energy* 2016, 1, 16119.
- [22] F. Wan, L. Zhang, X. Dai, X. Wang, Z. Niu, J. Chen, Nat. Commun. 2018, 9, 1656.
- [23] R. Trócoli, F. La Mantia, ChemSusChem 2015, 8, 481.
- [24] L. Zhang, L. Chen, X. Zhou, Z. Liu, Adv. Energy Mater. 2015, 5, 1400930.
- [25] N. Zhang, F. Cheng, Y. Liu, Q. Zhao, K. Lei, C. Chen, X. Liu, J. Chen, J. Am. Chem. Soc. 2016, 138, 12894.

- [26] G. Li, Z. Yang, Y. Jiang, C. Jin, W. Huang, X. Ding, Y. Huang, Nano Energy 2016, 25, 211.
- [27] C. Xu, B. Li, H. Du, F. Kang, Angew. Chem. 2012, 124, 957.
- [28] M. H. Alfaruqi, J. Gim, S. Kim, J. Song, D. T. Pham, J. Jo, Z. Xiu, V. Mathew, J. Kim, *Electrochem. Commun.* 2015, 60, 121.
- [29] B. Lee, H. R. Lee, H. Kim, K. Y. Chung, B. W. Cho, S. H. Oh, Chem. Commun. 2015, 51, 9265.
- [30] M. H. Alfaruqi, V. Mathew, J. Gim, S. Kim, J. Song, J. P. Baboo, S. H. Choi, J. Kim, *Chem. Mater.* 2015, 27, 3609.
- [31] G. G. Yadav, J. W. Gallaway, D. E. Turney, M. Nyce, J. Huang, X. Wei, S. Banerjee, *Nat. Commun.* 2017, 8, 14424.
- [32] G. Du, J. Wang, Z. Guo, Z. Chen, H. Liu, Mater. Lett. 2011, 65, 1319.
- [33] V. Augustyn, J. Come, M. A. Lowe, J. W. Kim, P.-L. Taberna, S. H. Tolbert, H. D. Abruña, P. Simon, B. Dunn, Nat. Mater. 2013, 12, 518.
- [34] T. C. Liu, W. Pell, B. Conway, S. Roberson, J. Electrochem. Soc. 1998, 145, 1882.
- [35] Y. Liu, F. Zhou, V. Ozolins, J. Phys. Chem. C 2012, 116, 1450.
- [36] A. K. Katz, J. P. Glusker, S. A. Beebe, C. W. Bock, J. Am. Chem. Soc. 1996, 118, 5752.
- [37] F. Zeng, Y. Pan, Y. Yang, Q. Li, G. Li, Z. Hou, G. Gu, Electrochim. Acta 2016, 196, 587.