Sodium vanadate/PEDOT nanocables rich with oxygen vacancies for high energy conversion efficiency zinc ion batteries

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\textbf{Abstract}

Aqueous zinc-ion batteries (ZIBs) have received an increasing attention for large-scale energy storage due to its low cost and high safety. However, the sluggish kinetics stemming from the limited Zn\textsuperscript{2+} diffusion pathway and the strong electrostatic interaction between Zn\textsuperscript{2+} and the anion sublattice in the host crystal are great challenges for high-efficient Zn\textsuperscript{2+} storage. To circumvent these hurdles, bridge-oxygen sites oxygen vacancies are introduced in sodium vanadate Na\textsubscript{0.76}V\textsubscript{2}O\textsubscript{4} (NVO) through the in-situ polymerization of 3,4-ethylenedioxythiophene. In the resultant oxygen-vacancy rich NVO/poly(3,4-ethylenedioxythiophene) (PEDOT) nanocables (noted as V0\textsuperscript{2+}-PNVO), the interplanar space is enlarged and the electrostatic interaction is weakened by the introduction of “oxygen vacancies” for fast reversible Zn\textsuperscript{2+} diffusion and intercalation. The conductive PEDOT coating also provides desired electronic conductivity and improves the structural integrity of NVO. Consequently, V0\textsuperscript{2+}-PNVO based ZIBs exhibit an enhanced specific capacity (355 mA h g\textsuperscript{−1} at 50 mA g\textsuperscript{−1}), significantly enhanced energy conversion efficiency (>80%), and long lifespan (99% of the initial capacity remained after 2600 cycles).

1. Introduction

Aqueous rechargeable batteries have emerged as promising alternative technology for large-scale energy storage due to their inherent safety, affordable cost and feasible fabrication [1,2]. Zn\textsuperscript{2+} possess high theoretical capacity (820 mA h g\textsuperscript{−1}), low redox potential (~0.76 V vs standard hydrogen electrode) and excellent stability in water, leading to increasing research works on rechargeable aqueous zinc-ion batteries (ZIBs). [2,3] Until now, various cathode materials have been studied for high-performance aqueous ZIBs, such as manganese-based materials [4–6], vanadium-based composites [7,8], Prussian blue and its analogues [9], and transition-metal dichalcogenides [10,11]. Among them, vanadium-based materials can deliver high theoretical specific capacity (up to 588 mA h g\textsuperscript{−1}) and excellent electrochemical stability owing to their multiple valence states and open-framework structure [12,13]. Pure vanadium oxides, such as V\textsubscript{2}O\textsubscript{3} [14] and VO\textsubscript{2} [15], usually show poor rate performance and cycling stability because their moderate sluggish ionic diffusion kinetics would lead to structure degradation during the repeated intercalation/deintercalation process of Zn\textsuperscript{2+}. Introducing metal ions (Na\textsuperscript{+}, K\textsuperscript{+}, Mg\textsuperscript{2+}. Al\textsuperscript{3+}) [16–19] or/and structural water molecules [20] between [V-O] polyhedron layers to expand interlayer space is an effective strategy to enhance Zn\textsuperscript{2+} diffusion kinetics. Metal vanadate, such as sodium vanadate [7,21] and potassium vanadate [18] with a layer or tunnel structure have also received much attention as the inserted metal ions can improve the structural stability. It has been reported that 2D layered sodium vanadate, such as Na\textsubscript{0.76}V\textsubscript{2}O\textsubscript{4} and HNa\textsubscript{0.76}V\textsubscript{2}O\textsubscript{4}•4H\textsubscript{2}O, can deliver high ion diffusion coefficients (from 10\textsuperscript{−10} to 6×10\textsuperscript{−10} cm\textsuperscript{2} s\textsuperscript{−1}) and high capacities (281 and 304 mA h g\textsuperscript{−1} at 500 mA g\textsuperscript{−1}, respectively) [21], because a layer structure provides effective pathway for Zn\textsuperscript{2+} diffusion. Unfortunately, the structural destroy and capacity fading also accompanied, i.e., only 71% capacity retention after 2000 cycles was retained for Na\textsubscript{0.76}V\textsubscript{2}O\textsubscript{4} [21]. Sodium vanadate with a 3D rigid tunnel structure, such as Na\textsubscript{0.76}V\textsubscript{2}O\textsubscript{15} [21](notes as NVO), can effectively alleviate the structure collapse and exhibit reversible insertion/extraction of Zn\textsuperscript{2+}. However, the tunneled NVO shows an unsatisfactory capacity (135 mA h g\textsuperscript{−1} at 0.5 A g\textsuperscript{−1}) because the tunnel structure with smaller lattice space provide less effective diffusion paths for Zn\textsuperscript{2+} than the layer structure [21]. Thus, more efforts are necessary to achieve a better fundamental understanding and to modify the tunnel sodium vanadate as high-capacity and superior-stability cathode materials for ZIBs.

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https://doi.org/10.1016/j.enm.2021.05.003
Received 6 January 2021; Received in revised form 11 April 2021; Accepted 3 May 2021
Available online 9 May 2021
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To circumvent the challenges discussed above, vanadium-based materials are often composed with conductive materials, such as carbon materials [22–24] and conductive polymers [25,26] to improve the electrochemical kinetics and cycling capability of ZIBs with their high electronic conductivity. In addition, defect engineering, especially oxygen vacancies, are introduced as an effective strategy to modify the crystalline structure of materials and reduce the activation energy of Zn$^{2+}$ insertion and diffusion [15,23,27]. Thus, the reaction and transport kinetics are intrinsically improved, and Zn$^{2+}$ storage capability is significantly enhanced by oxygen vacancies [15,23].

In this work, oxygen vacancies in sodium vanadate, Na$_{0.76}$V$_6$O$_{15}$ nanobelts (NVO), were introduced through in-situ polymerization of poly(3,4-ethylenedioxythiophene) (PEDOT) coating and studied for ZIBs. The resultant oxygen-vacancy rich NVO/PEDOT nanocables (noted as Vo$^\ddagger$-PNVO) demonstrated a high capacity, much enhanced transport properties, and excellent cycling stability, attributing to: (1) enlarged interplanar space benefitting ion diffusion, (2) reduced polarization arising from the electrostatic interactions, (3) excellent electronic conductivity of PEDOT coating, and (4) enhanced mechanical integrity of NVO coated with conductive and stable PEDOT. Density functional theory (DFT) calculations are used to confirm the presence and site of oxygen vacancies.

2. Experimental section

2.1. Materials synthesis

2.1.1. Synthesis of Na$_{0.76}$V$_6$O$_{15}$ nanobelts (NVO)

0.5 g of commercial V$_2$O$_5$ powder was added into 50 mL of DI water and kept stirring at room temperature for 2 h to form solution A. Then, 1.0 g sodium dodecyl benzene sulfonate dispersed in 30 mL DI water by stirring was added in solution A. After stirring for another 2 h, the obtained orange suspension was transferred to 100 mL Teflon-lined stainless-steel autoclave and kept at 180 °C for 7 days. After solvent replacement with DI water and ethanol, and the sample was freeze dried in vacuum. Finally, the NVO was obtained after being heated in air at 450 °C for 3 h.

2.1.2. Synthesis of Na$_{0.76}$V$_6$O$_{15}$/PEDOT nanocables (Vo$^\ddagger$-PNVO)

The obtained NVO was put in a vacuum environment with 0.5 mL HCl at room temperature for 1 h. Then, the sample was moved to a vacuum reactor with 0.1 mL 3, 4-ethylenedioxythiophene (EDOT) monomers and kept at 80 °C for 6 h. With EDOT monomers polymerized at the surface of NVO, Vo$^\ddagger$-PNVO nanocables were obtained after further dried in air at 120 °C overnight.

2.2. Material characterizations

X-ray diffraction (XRD) with Bruker D8 Discover X-ray diffractometer and an ImS 2-D detection system (50 kV, 1000 mA) was used to characterize the crystalline phase of materials. The composition of samples was identified with Fourier-transform infrared (FTIR) spectra using a Bruker-TENSOR27 FTIR spectrometer from 1800 to 400 cm$^{-1}$, Raman spectra performing on a Jobin-Yvon HR800 Raman microscope ($\lambda = 514$ nm) recorded from 100 to 1800 cm$^{-1}$, and Thermogravimetric and differential scanning calorimetry (TG-DSC) measurements on SDT Q600 from 25 to 700 °C (10 °C min$^{-1}$, flowing air). Electron spin resonance (ESR) measurements were carried out on Bruker BioSpin GmbH. X-ray photoelectron spectra (XPS) were obtained by using a Thermo ESCALAB 250XIIX spectrometer in the PHI-5000C ESCA system with Mg K$_\alpha$ radiation (hv = 1253.6 eV). The ratio of oxidation state was estimated from the corresponding peak area ratio. The morphologies and structures of materials were identified by field-emission scanning electron microscope (FE-SEM, S-4800) with an Energy-dispersive X-ray spectrometer (EDS), transmission electron microscopy and high-resolution transmission electron microscopy (TEM, HRTEM, JEOL-JEM2100) with Energy Dispersive X-ray spectrometry (EDS) element mapping analysis of High-Angle Anular Dark-Field Scanning Transmission Electron Microscopy (HADDF-STEM).

2.3. Electrochemical characterizations

Electrochemical performance of zinc ion batteries was tested using CR2032 coin-type cells fabricated in air, respectively. The corresponding electrolyte is 3 M zinc trifluoromethanesulfonate (98%, Zn(CF$_3$SO$_2$)$_2$) aqueous solution and 3 M ZnSO$_4$ aqueous solution for ZIBs, respectively. A glass fiber filter (Whatman, Grade GF/A) was chosen as the separator for ZIBs. Active material, conductive carbon and polyvinylidene fluoride (PVDF) were mixed in a mass ratio of 7: 2: 1 in N-methyl-2-pyrrolidione (NMP). Then, the obtained slurry was painted on the current collector (titanium foil) and heated at 100 °C in vacuum overnight. The mass loadings of the working electrodes are about 1.5 ~ 2 mg cm$^{-2}$. The electrochemical performance of the electrodes was tested by cyclic voltammetry (CV) and electrochemical impedance spectroscopy (EIS) measurements on CHI760e Shanghai Chenhua workstations. Galvanostatic charge/discharge (GCD) measurements were conducted with a multichannel battery testing system (LAND CT2001A). The working voltage of batteries was set from 0.3 to 1.5 V versus Zn$/\text{Zn}^{2+}$/Zn. Galvanostatic intermittent titration technique (GITT) was carried out at a current pulse of 50 mA g$^{-1}$ for 10 min and a following relaxation time for 30 min in each step. The internal reaction resistance (RR) was calculated using the IR drop with the applied current density and the resistances at different steps. EIS measurements were tested with a frequency range of 100 kHz to 0.1 Hz.

2.4. Density functional theory (DFT) calculations

DFT was used to identify the possible sites of oxygen vacancies in NVO by using Vienna ab initio simulation package (VASP) [28]. The exchange and correlation energy functional were treated by the Perdew–Burke–Ernzerh of the generalized gradient approximation (GGA) [29]. Interaction between ion and electrons was described with projector augmented wave pseudo potentials (PAW) [30,31] approach. The energy cutoff for the plane wave basis set was set as 500 eV, and the total energy was converged to 10$^{-5}$ eV. Effective U parameter of 3.1 eV was used to describe the electronic structure of V [32]. The lattice parameters of NaV$_6$O$_{15}$ were optimized to be 15.85 Å and 10.41 Å with $\alpha = 90°$, $\beta = 112°$, $\gamma = 90°$. K-points used for k-point sampling in irreducible region of the first Brillouin zone ensure the accuracy of the calculation within 1 meV.

3. Results and discussion

3.1. Composition and morphology of Vo$^\ddagger$-PNVO

Oxygen vacancies in Vo$^\ddagger$-PNVO nanocables were introduced with vaporized polymerization method [33]. In brief, with NVO as oxidant, vaporized EDOT in-situ oxidatively polymerized and chemically connected with NVO via the positive S cations in PEDOT and the negative O anions in NVO, forming a nanocable-structured Vo$^\ddagger$-PNVO. Meanwhile, oxygen vacancies and accompanied V$^{4+}$ were generated in Vo$^\ddagger$-PNVO in the non-oxygen atmosphere.

The XRD patterns of both Vo$^\ddagger$-PNVO and NVO demonstrate a pure Na$_{0.76}$V$_6$O$_{15}$ phase (JPSD No. 75-1653) in Fig. 1a. In this crystal structure, VO$_6$ octahedra connect the V$_6$O$_{15}$$^{-\delta}$ layers by corner-shared oxygen atoms and form a tunnel structure (insert in Fig. 1a). Three distinctive peaks at 2$\theta$ = 9.30°, 12.16° and 18.6° correspond to the (001), (200) and (202) planes of Na$_{0.76}$V$_6$O$_{15}$, respectively. The relatively reduced peak intensity of Vo$^\ddagger$-PNVO implies amorphous information which may be attributed to PEDOT and oxygen vacancies.

In the FTIR spectrum (Fig. 1b) of NVO, peaks at 493 and 553 cm$^{-1}$ originate from the stretching vibration of chain-coordinated V=O$_2$ bonds with
The peak at 767 cm\(^{-1}\) is related to the symmetric stretching vibration of bridge-coordinated V–O\(_6\) bonds, and those located at 995 and 958 cm\(^{-1}\) are attributed to the stretching vibration of terminal V=O bonds [13]. Absorption bands at around 1377 and 1631 cm\(^{-1}\) are caused by the vibration of Na–O bonds and O–H bonds from absorbed water, respectively [13,34]. In the spectrum of Vo'–PNVO, apart from the characteristic peaks of NVO, the stretching vibration of C–S bonds (690, 839, 933 cm\(^{-1}\)), C–O–C bonds (1207, 1140, 1087 cm\(^{-1}\)), C–C and C=C bonds (1397, 1522 cm\(^{-1}\)) are observed, suggesting the formation of PEDOT in Vo'–PNVO [35]. The peak of V–O\(_6\) bonds (767 cm\(^{-1}\)) vanishes, and only one peak for V=O bonds (984 cm\(^{-1}\)) is remained. Moreover, a blue shift in the peak of V–O\(_6\) bonds suggests the V–O\(_6\) bonds become stronger in Vo'–PNVO. [13] These changes should relate to the oxygen vacancies and the connections between PEDOT (positive S) and NVO (negative terminal O), respectively (further explorations in following part) [36]. The above analysis indicates that Vo'–PNVO remains its crystalline structure with tiny changes.

Raman spectra of Vo'–PNVO and NVO were collected as complementary fingerprints to the FTIR spectra (Fig. 1c). Peaks located at 122 and 152 cm\(^{-1}\) come from the bending vibration of O–V–O–V bonds, which are related to the compressive deformation of lattice in tunnel-structured NVO [13,37]. Comparatively, a red shift of O–V–O–V peak (3 cm\(^{-1}\)) in Vo'–PNVO suggests a weaker compression deformation [13]. As a result, a larger lattice space will be provided to facilitate the diffusion of Zn\(^{2+}\) in Vo'–PNVO, which can be confirmed by the following IFFT of HRTEM images. The bending vibration of V=O bonds in Vo'–PNVO becomes weak at 258 and 265 cm\(^{-1}\) due to the S–O connections with PEDOT [36], and the stretching signal of V–O\(_6\) bonds appears with a blue shift to 695 cm\(^{-1}\) [38], which is in accordance with FTIR analysis. The vibration of V–O\(_6\) bonds is slightly weakened at 440 and 516 cm\(^{-1}\) in Vo'–PNVO [23], while it disappears in FTIR analysis. This can be explained that V–O\(_6\) bonds are partially weakened by oxygen vacancies as oxygen vacancies are generated at the interface of NVO and PEDOT and inner part retains its original valence states, similar to our precious work Vo'–V2O5/polyaniline [33]. The formation of PEDOT is proved by the new bonds at 810, 1295, 1453, and 1560 cm\(^{-1}\), which are assigned to the C–S, C=C–C=, symmetric C=C=C=O and asymmetric C=C=C=O stretching modes, respectively. [25,39] Based on the FTIR and Raman spectra of Vo'–PNVO, PEDOT anchored NVO at the terminal O, and oxygen vacancies are most likely to form at O\(_6\) site (further confirmed by following theoretical calculations). In this case, there would be an enlarged space for Zn\(^{2+}\) diffusion (to be further discussed). The mass of PEDOT accounts for 17% in Vo'–PNVO, reflected by a mass loss and an exothermic peak at around 300 °C [40], as shown in TG-DSC results (Fig. S1).

Fig. 1d demonstrates the full survey XPS spectra of Vo’–PNVO and NVO. Compared with NVO, two new peaks of S 2 s and S 2 p are detected in Vo’–PNVO, which are the characteristic peaks of PEDOT. The S 2 p spectrum (Fig. 1e) can be deconvoluted into three peaks centered at 163.4, 164.6 and 168.3 eV, corresponding to S 2p3/2, S 2p1/2 and S O bonds, respectively [35,41]. The C 1 s spectrum of Vo’–PNVO in Fig. 1f is deconvoluted into three peaks of C=C=C=C, C=S, and C=O bonds in PEDOT [35,38,42], and the pure NVO without PEDOT only shows a weak signal of C 1 s spectrum (Fig. 1d). Fig. 1g displays asymmetric V 2p3/2
spectra of Vo⁺-PNVO and NVO. Calculated with the curve area ratio, the $V^{4+}/V^{4+}$ ratio of Vo⁺-PNVO (1.2) is lower than that of NVO (1.8), suggesting that a certain amount of $V^{4+}$ species are reduced to $V^{4+}$ when oxygen vacancies are generated in Vo⁺-PNVO. For O 1 s spectrum, NVO shows peaks of O–V and O–Na bonds at 529.9 and 531.4 eV, respectively (Fig. 1h) [23]. In contrast, the O 1 s spectrum of Vo⁺-PNVO is deconvoluted into four peaks at 529.6, 530.5, 531.5 and 532.8 eV, which correspond to the O–V bonds, oxygen vacancies, O–Na/O–S bonds and C–O bonds (thiophene ring) of PEDOT, respectively [23, 27, 43].

The bulk examination of oxygen vacancies was conducted by EPR measurements after above XPS surface characterizations. Oxygen vacancies as well as $V^{4+}$ can be identified by EPR due to unpaired electrons [15]. As shown in Fig. 1i, both NVO and Vo⁺-PNVO show characteristic EPR response at g = 1.999, and the peak intensity in Vo⁺-PNVO is slightly higher than that of NVO. This is because extra oxygen vacancies accompanied with $V^{4+}$ are generated in Vo⁺-PNVO besides the original $V^{4+}$ in NVO [15, 23, 38]. Considering the nanocable structure of Vo⁺-PNVO, the slight difference in bulk analysis (EPR) and significant change in surface analysis (XPS) confirm the surface distribution of oxygen vacancies in Vo⁺-PNVO. This surface distribution also agrees well with above XRD, FTIR and Raman analysis. Oxygen vacancies at the surface would facilitate the ion diffusion and electron transfer and suppress electrochemical polarization, improving the electrochemical rate performance and cycling stability of Vo⁺-PNVO.

The SEM images (Fig. 2a and b), TEM images (Fig. 2c and d) show a nanobelt structure for NVO, while Vo⁺-PNVO demonstrate a nanocable structure with a ~6 nm uniform coating. The uniform PEDOT coating is further confirmed by a homogeneous and broad distribution of S element, as shown in EDS elemental mappings of HADDF-STEM in Fig. 2i. The HRTEM image of NVO (Fig. 2e) and corresponding Inverse Fast Fourier Transform (IFFT) image (Fig. 2g) present perfect lattice fringes with an interplanar spacing of 3.46 Å, which is indexed to the (202) plane of NVO (JPD No. 75-1653). Comparatively, Vo⁺-PNVO show indistinct interface between PEDOT and NVO (Fig. 2f). This suggests the oxygen vacancies are induced in the near-surface layer of NVO, and it agrees well with the V–O₈ bond analysis based on FTIR and Raman spectra. Moreover, dislocations (red circles) are observed in Vo⁺-PNVO, and the corresponding interplanar spacing of (~406) plane is expanded from 1.66 Å to 1.98 Å (Fig. 2h). This is because oxygen vacancies cause the skewing of other atoms around and form dislocations in Vo⁺-PNVO, and the accompanied $V^{4+}$ with a bigger ionic radius (72 pm) also enlarge the interplanar distance than $V^{5+}$ (68 pm), which is consist with XPS and EPR results. As a result, ion diffusion would be enhanced and the electrostatic interactions would be weakened to improve the rate performance and cycling stability of Vo⁺-PNVO.

Based on above experimental analysis, the site of oxygen vacancies in NVO was further identified by DFT calculations. Different from orthorhombic V₂O₅ which consists of VO₃ pentahedrons, tunneled NVO consists of VO₆ octahedrons. As shown in Fig. 3a and b, VO₆ octahedrons connect each other through V–O₈ bonds and link layers with V–O₈ bonds. According to the crystalline symmetry, there are three possible sites in NVO: terminal O₈ (Site 1), O₈ (Site 2) and O₈ (Site 3). The calculations suggest that the free energy of oxygen vacancy on Site 2 (~317.585 eV) is smaller than that on Site 1 (~317.268 eV) and Site 3 (~317.259 eV), as shown in Fig. 3b. This result suggests that oxygen vacancies are easier to form at O₈ site, in agreement with the FTIR and Raman results. The V–O₈ bonds between layers are broken by oxygen vacancies, which can increase interplanar distance and reduce electrostatic interactions between Zn²⁺ and NVO, rendering fast Zn²⁺ diffusion and reversible Zn²⁺ intercalation in Vo⁺-PNVO.
3.2. Electrochemical characteristics and properties of Vo⁻-PNVO

The electrochemical characteristics and properties of Vo⁻-PNVO and NVO as cathode materials for ZIBs were evaluated by using coin cells with 3 M Zn(CF₃SO₂)₂ aqueous solution as electrolyte. In the cyclic voltammetry (CV) curves of NVO at 0.1 mV s⁻¹ (Fig S2a), there are three pairs of redox peaks centered at 1.1/1.28 V, 0.52/0.85 and 0.42/0.75 in the first cycle, which correspond to \( \text{V}^{5+}/\text{V}^{4+} \) and \( \text{V}^{4+}/\text{V}^{3+} \), respectively [19,37,38]. These redox peaks move to 1.1/1.33, 0.50/0.88 and 0.41/0.79 V in the subsequent 3rd and 4th cycles, respectively. The gradually enlarged voltage gap in redox pairs and reduced CV curve area suggest serious polarization and capacity decay in NVO. This is related to irreversible reactions, namely “dead Zn²⁺”, happened in the Zn²⁺ insertion/extraction processes, similar in Na₀.₃₃V₂O₅ [44]. In contrast, the 1st and 2nd CV curves of Vo⁻-PNVO are greatly different from the subsequent curves. Specifically, there are three pairs of redox peaks at 0.72/0.83, 0.56/0.70 and 0.34/0.56 V at beginning (Fig. 4a), indicating a multiple-step Zn²⁺ intercalation/de-intercalation reaction mechanism [44]. The almost vanished redox pair at around 1.10/1.28 V suggests that the redox reaction of \( \text{V}^{5+}/\text{V}^{4+} \) is weakened due to the introduction of oxygen vacancies in the shallow layer of Vo⁻-PNVO, which corresponds to the emergence of new redox pair at 0.72/0.83 V [15]. When vanadium ion has an oxygen octahedron coordination, an oxygen vacancy would render a change in electronic structure, causing the further splitting or deviation of orbital energy and leading to a change of charge-discharge voltage [45]. In the 3rd and 4th cycles, only two obvious pairs of redox peaks at 0.72/0.87 and 0.46/0.56 V are remained without deviation. This phenomenon indicates that the redox reactions in Vo⁻-PNVO become reversible from the 3rd cycle. As further compared in Table 1, the smaller voltage difference in the redox pairs of Vo⁻-PNVO suggests lower polarization and faster electrochemical kinetics than NVO. Oxygen vacancies can reduce the polarization with positive charge by balancing the strain/stress from Zn²⁺ and Vo⁻-PNVO. As the V⁴⁺ with partially filled 3d orbitals can capture and transfer electrons in redox reactions, the Zn-ion storage reaction kinetics would be catalyzed [17]. Thus, the number of d orbitals located at the conduction band bottom and the valence band top increases due to the introduced oxygen vacancies and V⁴⁺, enhancing the electrical conductivity and catalyzing the electrochemical reactions in Vo⁻-PNVO.

In the Galvanostatic charge-discharge (GCD) curves of the Vo⁻-PNVO (Fig. S2b), all the charge/discharge voltage plateaus connect and form a long and tilting platform. The initial discharge specific capacity of Vo⁻-PNVO is 355 mA h g⁻¹ at 0.05 A g⁻¹, and the specific capacity remains 295, 290, 285 mA h g⁻¹ in the following three cycles, in accordance with the CV curves. The detailed storage mechanism will be explored further explored later. The values of Vo⁻-PNVO are higher than that of NVO (315, 277, 230, 213 mA h g⁻¹ at 0.05 A g⁻¹). Moreover, Vo⁻-PNVO exhibits superior rate capability with capacities of 256, 234, 228, 222, 221, 205 and 165 mA h g⁻¹ at 0.1, 0.2, 0.4, 0.8, 1, 2 and 4 A g⁻¹, respectively, and the capacity almost recovers when the rate switches back to 0.05 A g⁻¹ (Fig. 4b). Compared with NVO, the improved capacity and higher rate performance of Vo⁻-PNVO are closely related to PEDOT and oxygen vacancies. Note that it has been reported that pure PEDOT plays a relatively small role in capacity contribution (24 mA h g⁻¹ at 0.2 A g⁻¹, and even smaller) and poor cycling performance [38,46]. The main role of PEDOT is to create oxygen vacancies and to simultaneously form a conductive coating to improve the overall conductivity and protect the core [25]. Thus, oxygen vacancies should mainly contribute to the high capacity. The rate performance of Vo⁻-PNVO in regards of fast charge transfer kinetics will be discussed next.

The high rate performance of Vo⁻-PNVO is reflected by energy efficiency, which is important for practical applications. In Fig. 4c, the smaller hysteresis area of the GCD curve suggests Vo⁻-PNVO a higher

![Figure 3](image-url) Fig. 3. (a) The crystal structure of NVO. (b) The possible sites for oxygen vacancies in NVO, and (c) corresponding calculated free energy. The smallest free energy on Site 2 suggests O₅ site is the location of oxygen vacancies in Vo⁻-PNVO.

**Table 1**

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<tr>
<th>Sample</th>
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<th>Difference (V)</th>
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<td>0.46/0.56</td>
<td>0.12</td>
</tr>
<tr>
<td>PNVO</td>
<td>0.72/0.87</td>
<td>0.15</td>
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<tr>
<td>NVO</td>
<td>0.41/0.79</td>
<td>0.38</td>
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<tr>
<td></td>
<td>0.50/0.88</td>
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<td>1.10/1.33</td>
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</table>
energy efficiency and less energy loss than NVO. Calculated by the ratio of discharge and charge energy densities, the energy efficiency of Vo–PNVO is higher than that of NVO in different current densities (Fig. 4c). The enhanced energy efficiency of Vo–PNVO is ascribed to highly reversible Zn$^{2+}$ intercalation/de-intercalation due to the introduction of PEDOT and oxygen vacancies. In addition, the Vo–PNVO-based ZIBs exhibit an energy-density/power-density of 228 W h kg$^{-1}$/40 W kg$^{-1}$ and 105 W h kg$^{-1}$/3160 W kg$^{-1}$ at 0.05 and 4.0 A g$^{-1}$, respectively, higher than NVO and the recently reported vanadium-, manganese- and Prussian blue-based ZIBs, as shown in Ragone plot in Fig. 4d.

Vo–PNVO also deliver excellent cycling performance. As shown in Fig. 4e, 106% capacity retention (275 mAh g$^{-1}$) and approximately 100% coulombic efficiency were obtained in Vo–PNVO after 500 cycles at a small current density of 0.1 A g$^{-1}$, while the capacity of NVO decays rapidly in 100 cycles (70% capacity retention). The well overlap of the 1st and 100th GCD curves of Vo–PNVO confirms the excellent cycling stability and high reversibility of Vo–PNVO (Fig. 4f). The nanocable structure of Vo–PNVO still remains well after 500 cycles, while the NVO structure was destroyed as small parts in 100 cycles, confirmed by the SEM and TEM images in Fig. S3a–d. Further tested at a large current density of 4 A g$^{-1}$, 99% of the initial capacity with approximately 100% coulombic efficiency is remained after 2600 cycles in Vo–PNVO (Fig. 4g). The corresponding GCD curves of the 10th, 100th, 1000th, 2000th and 2600th cycles at 4 A g$^{-1}$ also overlap well (Fig. 4i). Moreover, the cycling performance of Vo–PNVO is superior to other electrode materials, such as Na$_{0.33}$V$_{2}$O$_{5}$ (93% retention after 1000 cycles) [8], NH$_{4}$V$_{6}$O$_{10}$ (70.3% retention after 2000 cycles) [47], K$_{2}$V$_{8}$O$_{21}$ (83% retention after 300 cycles) [18], V$_{2}$O$_{5}$@PEDOT nanosheet arrays (89% retention after 1000 cycles) [25], polyaniline intercalated V$_{2}$O$_{3}$ (87.5% retention after 600 cycles) [26] and polyaniline intercalated V$_{2}$O$_{3}$-nH$_{2}$O (80% retention after 800 cycles) [34]. Four possible factors are ascribed to the excellent cycling stability of Vo–PNVO: (1) Positively charged oxygen vacancies reduce the electrostatic interactions between Zn$^{2+}$ and host to inhibit the structural collapse. Oxygen vacancies also provide void sites for fast electron reception/donation, and offer larger lattice distance for Zn$^{2+}$ diffusion, significantly enhancing the kinetics and electrochemical stability of Vo–PNVO. (2) Conductive PEDOT coating may also help to improve overall conductivity as well as preserve the morphology integrity by preventing the dissolution of inner NVO from directly contacting the electrolyte [25,36,48–50]. (3) The structure stability is also related to the chemical bonds between PEDOT and NVO in Vo–PNVO. The robust connection renders PEDOT coating firmly attached in Vo–PNVO after 500 cycles (Fig. S3d), while the pure PEDOT shows serious capacity fade [38]. (4) The electrolyte selection of 3 M Zn(CF$_{3}$SO$_{3}$)$_{2}$ matters in electrochemical stability. Vo–PNVO shows better cycling performance in 3 M Zn(CF$_{3}$SO$_{3}$)$_{2}$ than in 3 M ZnSO$_{4}$ aqueous solution (76% retention after 2000 cycles, Fig. S3e).

3.3. Zn-storage mechanism exploration of Vo–PNVO

Various sweeping rates of CV curves were collected to investigate the reaction kinetics behind the enhanced rate performance of Vo–PNVO (Fig. 5a). The power-law relationship between peak currents (i) and
sweep rates ($\nu$) is described as following:

$$i = a \nu^b$$

(1)

where $a$ and $b$ are variables. A $b$ value of 1 suggests a capacitive process and 0.5 is indicative of diffusion-controlled charge storage. With scan rates increased from 0.2 to 1 mV s$^{-1}$, the $b$ values of peaks 1 and 2 in Vo$^{-}$PNVO are 0.78 and 0.89, whereas these in NVO are 0.57 and 0.62, respectively (Fig. 5b). This indicates the electrochemical reactions in Vo$^{-}$PNVO and NVO are controlled by both surface-controlled capacitance and diffusion-limited redox reactions, and more the capacitive behaviors in Vo$^{-}$PNVO than in NVO at peak potentials. The capacitive and diffusion contribution at various scan rates can be obtained via the equation of

$$i(\nu) = k_1 \nu + k_2 \nu^{1/2}$$

(2)

where $k_1$ and $k_2$ are proportional to capacitance and diffusion contributions, respectively. A visual capacitive contribution of 78% in Vo$^{-}$PNVO at 0.8 mV s$^{-1}$ was shown in Fig. 5c. The capacitance contribution increases with sweep rate increased, and it is higher in Vo$^{-}$PNVO than NVO (Fig. 5d). As the high rate performance and energy efficiency of Vo$^{-}$PNVO are attributed to Zn$^{2+}$ diffusion and capacitive behavior, the reasons can be summarized as following: (1) Oxygen vacancies can promote the capacitance-type charge storage with fast charge storage kinetics. Oxygen vacancies enable fast electron transfer with void sites to fast accept/donate electrons, and promote Zn$^{2+}$ diffusion with expanded interplanar distance. (2) Conductive PEDOT enhances charge storage dynamics by enhancing electron transfer and shortening the charge transport distance. [36,48–50] (3) The electrolyte, namely, 3 M Zn(CF$_3$SO$_3$)$_2$, also relates to the Zn-storage capability of Vo$^{-}$PNVO. When tested with 3 M ZnSO$_4$ aqueous solution as electrolyte, Vo$^{-}$PNVO based ZIBs exhibit less satisfactory rate performance and stability than 3 M Zn(CF$_3$SO$_3$)$_2$ aqueous solution, although it is better than in NVO (Fig. 5a–e). Zn(CF$_3$SO$_3$)$_2$ electrolyte favors superior electrochemical properties due to its higher ionic conductivity and facilitated charge transfer, which probably because of the smaller charge density (compared with double-charge SO$_4^{2-}$) and weaker solvation of bulky CF$_3$SO$_3^-$ anions [4,51,52].

Galvanostatic intermittent titration technique (GITT) was conducted to analyze the interfacial charge-transfer kinetics of Vo$^{-}$PNVO. As shown in Fig. 5a–c, the internal reaction resistance (RR) of Vo$^{-}$PNVO (0−0.3 Ω g$^{-1}$) is much smaller than NVO (0.2−1.2 Ω g$^{-1}$). Moreover, the corresponding Zn$^{2+}$ diffusion coefficient ($D_{Zn^{2+}}$) of Vo$^{-}$PNVO is calculated between 10$^{-14}$ and 10$^{-13}$ cm$^2$ s$^{-1}$, higher than that of NVO from 10$^{-15}$ to 10$^{-14}$ cm$^2$ s$^{-1}$. The results suggest that Vo$^{-}$PNVO has fast kinetics for better rate capability than NVO. Electrochemical impedance spectroscopy (EIS) was also used in Fig. 5e. The Nyquist plot consist of a semi-circle at the high-frequency region and a linear tail at low-frequency region, which are related to charge transfer resistance ($R_c$) and ion diffusion. In Table 2, Vo$^{-}$PNVO show a lower $R_c$ value (5.7 Ω) than NVO (25.6 Ω), suggesting faster charge transfer in Vo$^{-}$PNVO. The $D_{Zn^{2+}}$ values of Vo$^{-}$PNVO and NVO are calculated through the relationship between low frequencies and the real part of impedance (Fig. 5f) [13]. Vo$^{-}$PNVO present a larger $D_{Zn^{2+}}$ of 3.5 × 10$^{-14}$ cm$^2$ s$^{-1}$ compared to that of 8.9 × 10$^{-15}$ cm$^2$ s$^{-1}$ in NVO. Consistent with GITT.
result, the lower \( R_s \) value and higher \( D^{2+} \) in Vo−PNVO confirm the contributions of oxygen vacancies and PEDOT in electron transfer and ion diffusion (experimentally and theoretically discussed above). Thus, the resulted fast redox reactions and stable crystal structure endow the above high rate performance, energy efficiency and cycling stability of Vo−PNVO.

The Zn-ion storage mechanism of Vo−PNVO was studied by means of ex-situ XRD. The XRD patterns of full charge and discharge states in the initial three cycles are displayed in Fig. 6a. In the first full discharge state, the main (200) peak shifts slightly to a high angle from the pristine state, and the (001), (401), (−202) and (−204) peaks move to the low angle regions. The peak shifts suggest the (001) facet is enlarged from 9.72 Å to 9.92 Å, while the (200) and (−202) plane is contracted slightly, which is because Zn\(^{2+}\) intercalated into the structure interior and have electrostatic interaction with the host Vo−PNVO [47,53]. At the same time, new peaks (marked with club) located at 6.4°, 12.9°, 19.4°, 26.2°, and 32.8° emerge, matching well with the patterns of Zn\(_2\)(CF\(_3\)SO\(_2\))\(_3\)OH\(_{2x-y}\)\(\cdot\)nH\(_2\)O [51,54]. It is reported as the co-insertion mechanism of a proton and Zn\(^{2+}\), where OH− reacts with Zn(CF\(_3\)SO\(_2\))\(_2\) and Zn\(_2\)(CF\(_3\)SO\(_2\))\(_3\)OH\(_{2x-y}\)\(\cdot\)nH\(_2\)O forms on the surface of the electrode. [54,55] The proton insertion corresponds to the capacitive contributions from Eq. (2). Note that the insulating Zn\(_2\)(CF\(_3\)SO\(_2\))\(_3\)OH\(_{2x-y}\)\(\cdot\)nH\(_2\)O is undesirable as the battery internal resistance will be increased and the electrochemical reactions would be impeded with decreased active area. [54] Thus, the formation of Zn\(_2\)(CF\(_3\)SO\(_2\))\(_3\)OH\(_{2x-y}\)\(\cdot\)nH\(_2\)O should be mainly responsible for the capacity decay from 355 to 295 mA h g\(^{-1}\) in the first discharge process. In the following charge state, the Zn\(_2\)(CF\(_3\)SO\(_2\))\(_3\)OH\(_{2x-y}\)\(\cdot\)nH\(_2\)O phase disappears and Vo−PNVO peaks recover to original states, accompanied with another series of peaks belonging to Zn\(_2\)V\(_2\)O\(_7\)(OH\(_2\))\(\cdot\)2H\(_2\)O (JCPDS No.87-0417, marked with diamond) [26]. Then, the above two byproducts reversibly and alternately formed in the following charge and discharge states. The subsequently formed Zn\(_2\)V\(_2\)O\(_7\)(OH\(_2\))\(\cdot\)2H\(_2\)O probably shows certain electrochemical activity and contribute partially to the capacity by involving in the Zn\(^{2+}\) storage, which has been reported and will not be further investigated [26,47]. In brief, the byproducts of Zn\(_2\)(CF\(_3\)SO\(_2\))\(_3\)OH\(_{2x-y}\)\(\cdot\)nH\(_2\)O and Zn\(_2\)V\(_2\)O\(_7\)(OH\(_2\))\(\cdot\)2H\(_2\)O can explain the decay in the initial capacities (355, 295, 290 and 285 mA h g\(^{-1}\)), as well as the difference in the CV curves of Vo−PNVO. Note that, although the byproducts form, the main crystalline structure is still Vo−PNVO which deliver high rate performance and energy efficiency due to oxygen vacancies and PEDOT (discussed above).

The reaction mechanism of Vo−PNVO was further examined by ex-situ XPS measurements, as shown in Fig. 6b–d. Na 1 s is observed in both charge and discharge processes of Vo−PNVO (Fig. 6b). Zn 2p peaks observed in full charge states imply incomplete extraction of Zn\(^{2+}\) or new phase form (Fig. 6c). The peak gap between O 1 s and V 2p\(_{3/2}\) gets

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**Table 2**

<table>
<thead>
<tr>
<th>Sample</th>
<th>( R_s ) (Ω)</th>
<th>( R_{ct} ) (Ω)</th>
<th>( D^{2+} ) (cm(^{2}) s(^{-1}))</th>
</tr>
</thead>
<tbody>
<tr>
<td>Vo−PNVO</td>
<td>1.4</td>
<td>5.7</td>
<td>( 3.5 \times 10^{-14} )</td>
</tr>
<tr>
<td>NVO</td>
<td>2.6</td>
<td>25.6</td>
<td>( 8.9 \times 10^{-15} )</td>
</tr>
</tbody>
</table>

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Fig. 6. Ex-situ of Vo−PNVO at different full discharge and charge states: (a) ex-situ XRD patterns, ex-situ XPS patterns of (b) full survey, (c) Na 1 s, (d) Zn 2p and (e) V 2p spectra. (f) Illustration of the energy storage mechanism of Vo−PNVO.
larger in discharge states than in charge states (Table S1), and the V2p3/2 spectra appear at lower energy regions at full discharge states than full charge states (Fig. S6a). The above result suggests more low vanadium valences in discharge states reduced along with the intercalation of Zn2+. Thus, the V 2p3/2 spectra were deconvoluted to explore the vanadium oxidation states. In the discharge-charge process (Fig. S6b–d), the pristine V5+ and V4+ were reduced into V4+ and V3+ at discharge states and recovered to V5+ and V4+ at charge states. The stable discharge-state D and charge-state E electrodes were further investigated. The SEM images (Fig. S7a and b) show that there are flakes on the surface of discharge-state D electrode. Based on the XRD and the EDS results (Fig. S7c), these flakes should be Zn2(CF3SO3)2(OH)2x·nH2O. Moreover, XPS V 2p3/2 spectra shows that there are V4+ and V5+ at the surface of discharge-state D electrode, while V5+, V4+ and V3+ are collected when the D electrode was etched (Fig. 6d). This confirms that high valence vanadium was reduced with the Zn2+ intercalation. As XPS measurements focus on surface characterizations, the result also indicates the surface-layer location of oxygen vacancies in Vo-PNO. The charge state E electrode present large ZnV2O5(OH)2·2xH2O sheets at the surface (SEM images in Fig. S7d and e), which are confirmed by EDS analysis in Figure S7f. In XPS V 2p3/2 spectra in Fig. 6e, only V5+ is detected (most likely comes from the Zn2V2O5(OH)2·2H2O at the surface), while V5+ along with a small amount of V4+ exist in inner part after surface etching. Based on the ex-situ XRD, XPS and SEM results, the possible redox reactions are illustrated in Fig. 6f: Zn2(CF3SO3)2(OH)2x·nH2O and Zn2V2O5(OH)2·2H2O reversibly and alternately formed in the discharge and charge states after the first discharge.

4. Conclusions
Oxygen-vacancy rich Na0.28V6O15/PEDOT nanocables (Vo-PNCO) were synthesized as cathode materials for aqueous ZIBs. Both experimental results and DFT calculations elucidated that the bridge-oxygen sited oxygen vacancies in Vo-PNCO provide large interplanar distance, remarkably enhancing the electron transfer and Zn2+ diffusion and weaken the electrostatic interactions. With the synergistic effects of the oxygen vacancies and conductive PEDOT coating, Vo-PNCO-based ZIBs deliver excellent performance with high energy conversion efficiency, high energy and high power, and excellent cycling stability (99% capacity retention over 2600 cycles). The results demonstrated that introducing oxygen vacancies is an effective strategy for high-performance ZIB cathodes. In addition, the mechanism study offers detailed information for better understanding of Zn2+ ion storage.

Declaration of Competing Interest
The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper

CRedit authorship contribution statement

Wenchao Bi: Conceptualization, Methodology, Software, Investigation, Writing – original draft, Writing – review & editing, Visualization. Guohua Gao: Software, resources, Supervision, Writing – original draft, Funding acquisition. Guangming Wu: resources, Supervision, Project administration, Funding acquisition. M. Atif: resources. MS AlSalhi: resources. Guozhong Cao: resources, Supervision, Writing – review & editing, Visualization, Funding acquisition.

Data availability
Data will be made available on request.

Acknowledgement
The authors are grateful for the financial support from National Natural Science Foundation of China (grant No. 51872204, No. 52072261, No. 22011540379), the National Key Research and Development Program of China (grant No. 2017YFA0204600), Shanghai Social Development Science and Technology Project (grant No. 20DZ1201800), Shanghai Sailing Program (21YF1430900). This work is also supported by the National Science Foundation (1803256). The authors are grateful to the Deanship of Scientific Research, King Saud University for funding through Vice Deanship of Scientific Research Chairs.

Supplementary materials
Supplemental material associated with this article can be found, in the online version, at doi:10.1016/j.ensm.2021.05.003.

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