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REVIEW



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1. Introduction

The international community has increasingly prioritized addressing global climate change, establishing carbon neutrality as a shared objective.^{1,2} Energy storage systems are essential for advancing energy transformation and supporting low-carbon development, particularly in response to the ongoing growth in global energy demand.³⁻⁶ High-performance energy storage systems play a crucial role in reducing reliance on fossil fuels, optimizing and upgrading the energy infrastructure, and promoting sustainable energy utilization.



Botao Wan,†^a Yajiang Wang,†^a Xiudong Chen,*^a Changchao Zhan, ^b^a Huixiong Jiang,^a Jin-Hang Liu,^a Yun Gao,*^{bc} Xiaoduo Jiang,^a Xiaohua Cao,^a Hang Zhang,*^{bcd} Shi-Xue Dou^b^e and Yao Xiao^b*^{bcd}

Aqueous zinc-ion batteries (AZIBs) are gaining significant attention due to their excellent safety, costeffectiveness, and environmental friendliness, making them highly competitive energy storage solutions. Despite these advantages, the commercial application of AZIBs faces substantial challenges, particularly those related to performance limitations of cathode materials. Among potential candidates, vanadium dioxide (VO₂) stands out due to its exceptional electrochemical properties and unique crystal structure, rendering it a promising cathode material for AZIB applications. The review summarizes the recent research progress on VO₂ in AZIBs, analyzes its crystal structures (tetragonal VO₂(A), monoclinic VO₂(B, D, M), and rutile VO₂(R)), morphology and energy storage mechanisms (Zn²⁺ insertion/extraction, H⁺/ Zn²⁺ co-insertion/extraction, and chemical reaction mechanism), and discusses the relationship between the structure and performance. The review also addresses key challenges associated with VO₂ as a cathode material, including dissolution, by-product formation, and limited ion diffusion kinetics. To overcome these issues, various optimization strategies are systematically discussed, such as ion/ molecule pre-intercalation, composite material fabrication, defect engineering, and elemental doping. Finally, potential research directions and strategies to further enhance the performance and commercial viability of VO₂-based cathodes are proposed.

> Additionally, they facilitate efficient energy allocation, providing robust technical support to ensure the achievement of carbon neutrality.7-9 Lithium-ion batteries (LIBs) are widely recognized as one of the most advanced energy storage technologies available today, valued for their unique advantages.10-13 LIBs offer a long cycle life with minimal capacity decay, ensuring durability over extended periods of working. Additionally, their low self-discharge rate minimizes energy loss during periods of inactivity. These characteristics have led to their widespread adoption in electric vehicles, portable electronic devices, and large-scale energy storage systems.14-22 However, LIBs are prone to overheating during charging and discharging, with a potential risk of thermal runaway and organic electrolyte combustion. Furthermore, the large-scale application of LIBs is heavily influenced by the limited availability of lithium metal, which is affected by geopolitical factors, resource distribution, and other constraints.23 Addressing these issues requires the development of energy storage devices that combine high capacity, environmental sustainability, and superior safety.24-26

> Aqueous rechargeable metal-ion batteries, such as those based on potassium,²⁷ sodium,²⁸ magnesium,²⁹ aluminum,³⁰ and zinc, have garnered significant attention due to their high capacity, safety, and environmental sustainability. Among

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^aSchool of Chemistry and Chemical Engineering, Jiangxi Province Engineering Research Center of Ecological Chemical Industry, Jiujiang University, Jiujiang 332005, China. E-mail: chenxd@jju.edu.cn

^bCollege of Chemistry and Materials Engineering, Wenzhou University, Wenzhou, Zhejiang 325035, China. E-mail: yungao1995@shu.edu.cn; hang@wzu.edu.cn; xiaoyao@wzu.edu.cn

^cWenzhou Key Laboratory of Sodium-Ion Batteries, Wenzhou University Technology Innovation Institute for Carbon Neutralization, Wenzhou, Zhejiang 325035, China

^dKey Laboratory of Advanced Energy Materials Chemistry (Ministry of Education), College of Chemistry, Nankai University, Tianjin 300071, China

^cInstitute of Energy Materials Science (IEMS), University of Shanghai for Science and Technology, Shanghai 200093, China

[†] Botao Wan and Yajiang Wang contributed equally to this work.

Chemical Science

these, aqueous zinc-ion batteries (AZIBs) stand out as a particularly promising contender.31,32 AZIBs offer several advantages, including a high theoretical specific capacity of 800 mA h g^{-1} , a low redox potential of -0.76 V, and abundant natural zinc reserves.³³⁻³⁶ To fully exploit these benefits, the development of cathode materials plays a pivotal role in determining the performance of rechargeable batteries. For the commercialization of AZIBs, structurally stable and high-capacity cathode materials are essential.^{32,37,38} The cathode materials for AZIBs can be broadly categorized into manganese (Mn)-based materials,^{39,40} vanadium-based materials,⁴¹ Prussian blue analogs (PBAs),⁴² organic compounds,³⁷ and other types.^{43,44} Manganesebased compounds offer notable advantages, such as resource abundance and low cost, making them attractive options for AZIB cathodes. However, their practical development is hindered by challenges such as poor electrical conductivity and high solubility, which significantly impact their performance and stability.35 PBAs are distinguished by their open threedimensional framework and high working voltage. Nevertheless, their application is limited by structural instability, which can lead to structural collapse, capacity fading, and performance degradation during cycling. Similarly, the development of organic compounds as cathode materials faces challenges due to their complex synthesis processes and susceptibility to side reactions, which further hinder their practical implementation.45 Vanadium-based compounds are considered promising candidates for next-generation AZIB cathodes due to their open framework structures, low cost, and high specific capacity.46 Among these, vanadium dioxide (VO2) stands out due to its unique crystal structure and high reversible specific capacity. Additionally, the stable structure and excellent ion transport properties of VO2 provide superior performance and enhanced safety compared to other vanadium oxides, making it an especially attractive option for AZIB applications.47

VO₂, with its unique physicochemical properties, has been extensively studied as a cathode material for AZIBs, demonstrating significant potential for practical applications. However, comprehensive and systematic analyses of its progress, challenges, and commercial prospects remain limited. This review focuses on the structure-activity relationship between the VO₂ crystal structure and zinc storage performance, delves into its energy storage mechanisms, and underscores its critical role in energy conversion and storage (Fig. 1). Despite its strengths, VO₂ faces challenges akin to those of other metal oxide cathode materials, including poor structural stability, reduced cycling performance, and the loss of active sites, which collectively constrain its wider application in AZIBs. To overcome these obstacles, researchers have proposed several innovative approaches to stabilize the VO₂ crystal structure and enhance its electrochemical performance. Ion intercalation, for instance, achieves simultaneous improvements in structural stability and ion conductivity through precise control of the VO₂ lattice structure. Molecular intercalation exploits interactions between organic or other molecules and VO₂ layers to effectively enhance the material's electrochemical properties. The composite material strategy integrates VO₂ with other functional materials, leveraging synergistic effects to substantially



Fig. 1 Schematic illustration showing the recent advances in VO₂based cathode materials for AZIBs.⁶¹ Copyright 2021, Wiley-VCH.⁶⁹ Copyright 2019, American Chemical Society.⁷⁴ Copyright 2021, Elsevier.⁷⁶ Copyright 2023, Elsevier.¹⁰⁴ Copyright 2020, American Chemical Society.¹⁰⁸ Copyright 2019, American Chemical Society.

boost overall battery performance. Defect engineering refines the electronic structure and reaction activity by meticulously controlling defect types and concentrations, thereby improving capacity and cycling stability. While these strategies have yielded impressive advancements in VO₂ performance, several technical hurdles persist. Developing composite material systems and exploring how defect engineering affects the electrochemical properties of VO₂ remain crucial. Addressing these challenges is pivotal to advancing the practical application of VO₂-based cathodes in AZIBs.

2. Crystal structure and morphology of VO₂

Vanadium-based cathode materials exhibit rich valence states and diverse crystal structures. These features enable multielectron transfer, maintain local electrical neutrality, and reduce polarization caused by zinc ion embedding. As a result, they deliver excellent zinc storage performance. Vanadium oxide compounds such as V2O3, VO2, V2O5, V6O13, V3O7, and V₅O₁₂ have shown significant potential for zinc storage.⁴⁸ Among them, VO₂ stands out due to its unique and stable crystal structure. This structure allows efficient insertion and extraction of Zn²⁺ ions, resulting in a high theoretical specific capacity, good cycle stability, and high energy and power density. In addition, VO₂ has tunnel and layer structures in which vanadium oxygen layers are stacked on top of each other by weak van der Waals forces. This structure provides a natural pathway for the embedding and deembedding of Zn²⁺ ions, allowing ions to move freely between layers. The crystalline

structure of VO₂ is capable of maintaining a high degree of stability in the processes of Zn^{2+} insertion and extraction. This is due to the good flexibility and plasticity of the VO₂ structure, which can adapt to the lattice changes caused by ion embedding and deembedding, thus effectively reducing the destruction of the material structure and capacity attenuation. These properties make VO₂ an ideal cathode material for AZIBs.⁴⁹

2.1 Crystal structure of VO₂

The diversity of crystal structures of VO₂ has made it a focus of research among the vanadium oxide system. Numerous crystal structures have been studied for VO2, including tetragonal VO₂(A), monoclinic VO₂(B, D, M), and rutile VO₂(R) (Fig. 2).⁵⁰ In spite of the fact that the crystal structures of VO₂ differ greatly according to the coordination of V and O, the open framework and the coordination of O are remarkably similar. In addition to enriching the physical and chemical properties of VO₂, the rich and diverse crystal structures of VO2 also provide broad avenues for its innovative applications in multiple fields like energy storage.⁵¹ The crystal structure of tetragonal VO₂(A) is relatively rare when compared to other crystal structures of VO2. As a result of its lattice arrangement, VO₂(A) can only be synthesized under certain specific synthesis conditions, as a result of its specific symmetry.⁵² Due to the fact that relatively few studies have been conducted on $VO_2(A)$, its physical and chemical

properties are not yet fully understood. While research continues to deepen, people gradually begin to realize that $VO_2(A)$ may be able to play a role in certain specific fields where the material has potential applications.

Monoclinic $VO_2(B, D, M)$ is the most common crystal structure type in VO₂. As a result of its good ion conductivity and stability, VO₂(B) has attracted a great deal of attention. The crystal structure of VO₂(B) provides excellent ion conduction properties, making it a potential energy storage material. In the structure of $VO_2(B)$, the V^{4+} ions represent the body center position with each V atom surrounding an oxygen octahedron (VO_6) , which shares edges with the V⁴⁺ ions. VO₂(B) is a space group $P2_1/c$, lattice parameters are a = 0.575 nm, b = 0.452 nm, c = 0.538 nm, $\beta = 122.6^{\circ}$ allowing for layered structures suitable for ion storage.⁵² Using a hydrothermal method and a rapid multi-wave reaction process, Pinnock et al. synthesized tunnelshaped VO₂(B).⁵³ VO₂(B) obtained by the multi-wave reaction process (Fig. 3a) can be decomposed into nanorods and clusters, and the hydrothermal synthesis rule produces nanorods with uniform morphology. In Fig. 3b, Zn²⁺ intercalated VO₂(B) nanorods are embedded and released along b and c. The X-ray diffraction (XRD) spectra of the two synthesized materials are similar, but the multiwave reaction is highly crystallized. According to the rate performance diagrams (Fig. 3c), VO₂(B) exhibits excellent energy storage properties due to its high



Fig. 2 The lattice structures of VO₂: (a) VO₂(A), (b) VO₂(B), (c) VO₂(M), and (d) VO₂(R), each viewed from different facets.



Fig. 3 (a) SEM images of multiwave digested VO₂(B), (b) schematic of Zn^{2+} (de)intercalation in VO₂(B) nanorods projected along the *b*- and *c*directions, and (c) rate capacity tests for AZIBs, with cathodes synthesised *via* hydrothermal and multiwave synthesis approaches.⁵³ Copyright 2025, the Royal Society of Chemistry. (d) XRD pattern of VO₂(M), (e) H⁺ diffusivity *vs.* specific capacity, (f) rate capability at varying current densities and corresponding coulombic efficiencies, and (g) extended cycling performance and the corresponding coulombic efficiency at 20 A g^{-1.56} Copyright 2022, the Royal Society of Chemistry. (h) XRD pattern of VO₂(D), (i) discharge–charge profiles of the VO₂(D) electrode obtained at a current density of 100 mA g⁻¹ and (j) cycling performance at a current density of 0.5 A g^{-1.57} Copyright 2018, the Royal Society of Chemistry. (k) Schematic of the VO₂(M) hollow sphere fabrication processes. (l) XRD pattern of VO₂(D) and (m) the cycling performance of the VO₂(M) and VO₂(R) cathodes at 1.0 A g^{-1.59} Copyright 2024, Elsevier.

specific capacity and good rate performance. This enhancement is attributed to the morphology formed in the hydrothermal samples, increased oxygen vacancies and higher V^{4+} concentrations, which may contribute to more efficient Zn^{2+} ion intercalation compared to multiwave reactor samples.

 $VO_2(M)$ is another monoclinic variation that has received much attention in the recent past. There is a doubling of the unit cell in the M phase of VO₂ as vanadium atoms are paired along the chain, resulting in a doubling of the vanadium atoms. Interestingly, the vanadium atoms have a dislocated lateral displacement, while the oxygen octahedron is twisted and rotated as well.⁵⁴ The crystal structure of VO₂(M) undergoes significant transformation with temperature variation, leading to abrupt changes in its physical properties, such as electrical resistance. With its properties, $VO_2(M)$ is an excellent choice for the preparation of temperature-controlled materials, such as smart windows.55 Jiang's group oxidized V2O5 with H2O2 and compounded it with carbon nanotubes to obtain VO₂(M)/CNT films.⁵⁶ XRD peaks of the obtained product match those of standard card VO₂(M) (Fig. 3d), proving that VO₂(M) was successfully synthesized. Through the carefully designed threedimensional porous structure, VO2 particles can be dispersed well, promoting electrolyte transmission, relieving strain during ion insertion, and reducing by-product formation. In AZIBs, it has a fast ion transmission rate of 10^{-7} (Fig. 3e), exhibits ultrahigh rate performance (Fig. 3f), and maintains a capacity of 232.6 mA h g⁻¹ at a high current density of 20 A g⁻¹. AZIB cathode materials based on VO₂(M) exhibit excellent electrochemical performance, providing a potential strategy for designing high-performance cathodes.

In addition to $VO_2(B)$ and $VO_2(M)$, $VO_2(D)$ is also a variant of the monoclinic system, which is also referred to as $VO_2(D)$. Despite the fact that $VO_2(D)$ has been less studied, it has been demonstrated that under certain conditions it may have unique physical and chemical properties. Consequently, in the future, with the increased depth of scientific research, it is expected that people will gradually become aware of the potential application value of VO₂(D). Wei et al. hydrothermally reacted ammonium vanadate with formic acid to obtain VO₂(D).⁵⁷ The XRD spectrum of the measured sample matches that of the VO₂(D) standard card (Fig. 3h), demonstrating successful synthesis of VO₂(D). It is used as a cathode material to measure electrochemical properties. At a current density of 0.1 Ag^{-1} , the specific capacity can reach 265 mA h g^{-1} and still deliver 249 mA h g⁻¹ after 100 cycles (Fig. 3i). $VO_2(D)$ can also maintain 94.3% after 350 cycles at 0.5 A g^{-1} (Fig. 3j), demonstrating its high zinc storage capacity and long cycle stability. At the time of charging to 1.4 V, the (001) crystal surface of XRD expanded to

12.6 Å, which was consistent with the reported layered vanadium oxide $V_2O_5 \cdot nH_2O_5$, and no reversible disappearance or appearance occurred in the subsequent charging and discharging processes, so it is believed that this was a phase transition reaction. After a series of characterization studies such as in situ XRD, it is proved that the material transforms from $VO_2(D)$ to $V_2O_5 \cdot H_2O$ in the first cycle, thus realizing stable zinc storage. During the phase transition from the ordered structure of VO₂ to the disordered structure of V₂O₅, a unique bilayer lamellar structure is formed, which is composed of double chains stacked face-to-face. It is particularly noteworthy that this newly formed phase plays a crucial role in subsequent electrochemical cycling and exhibits excellent flexibility, enabling it to adaptively adjust the ion diffusion paths to effectively accommodate volume changes, thereby demonstrating remarkable zinc storage stability.

The rutile structure of $VO_2(R)$ is the stable structure of VO_2 when heated above a certain temperature. A phase transition will occur between monoclinic variants such as VO₂(M) and rutile $VO_2(R)$, accompanied by significant changes in the crystal structure and a sudden change in physical properties. Because the vertices and edges of the VO_6 octahedron are shared, the VO₂(R) octahedron forms a more complex network structure. The lattice constants of VO₂(R) are a = b = 0.451 nm and c =0.304 nm, and it belongs to the $P4_2/mnm$ space group.⁵⁸ It is important to note that $VO_2(R)$ is a substance that has excellent chemical and thermal stability, making it suitable for use in harsh environments such as high temperatures and high pressures. Therefore, rutile VO₂(R) can be used for a wide range of applications, including ceramics, coatings, catalysts, and many others. Using oxalic acid and vanadium oxalate, Aydogdu et al.59 produced vanadium oxalate. In an argon atmosphere, they calcined the filtered and dried product and successfully converted VO₂ from the M phase to the R phase, which is suitable for use in AZIBs. VO₂(R) exhibits higher specific capacity/power density and lower charge transfer resistance due to its metalinsulator transition behavior, which result in improved electrochemical performance.

2.2 Morphology of VO₂

The development of VO₂ materials is frequently guided by morphology engineering design. VO2 of the same crystal form can have different morphologies when synthesized using different synthesis strategies. Cathode materials with different morphologies, however, have different cycling stabilities.⁶⁰⁻⁶² On the one hand, it is possible to shorten the diffusion path of Zn²⁺ using some morphological structures, which facilitates the rapid insertion/extraction of Zn²⁺. On the other hand, some materials with a dense stacking or irregular morphology may suffer from performance degradation during the cycle as a result of structural damage, leading to degradation of performance.63 A new type of VO₂ nanosheet has been successfully developed by Cao et al.64 with high uniformity and a circular structure. With a diameter of about 500 nm, a thickness of about 50 nm, and a hole in the middle, the nanosheet resembles an ancient coin (Fig. 4a-c). Circular nanosheets can

be divided into middle and outer parts based on their curvature. The microcrystal's middle region can be considered a smaller circular nanosheet with a higher curvature than its outer region, so it has a higher surface energy than its outer region. Once equilibrium is reached, the structure remains highly stable. For AZIBs with VO₂ as the cathode, a reversible capacity of 394 mA h g⁻¹ is achieved at 0.1 A g⁻¹. According to Bai *et al.*,⁶⁵ curled VO₂(B) nanosheets were produced by adding hydrogen peroxide (H₂O₂) to the commercial V₂O₅ precursor (Fig. 4d–f). The insertion and extraction of Zn²⁺ in/from the precursor do not compromise its structural integrity. The application of VO₂(B) nanosheets in AZIBs showed good electrochemical performance (400.2 mA h g⁻¹ at 0.5 A g⁻¹).

Hierarchical nanoflowers with a larger specific surface area can provide more active sites for zinc storage. A VO₂(B) nanosheet with a petal-like folded top was reported by Bai et al. (Fig. 4g-i).⁶⁶ Electrolytes enter through the gaps between the nanosheets, bringing more ions and facilitating faster storage. The higher pseudocapacitive contribution and partial diffusion in VO₂(B) enhance its electrochemical activity, accelerate electron transfer kinetics, and improve electronic conductivity. $VO_2(B)$ shows a significant capacity of 301.7 mA h g⁻¹ at 1 A g⁻¹. In an in situ self-conversion method, Deng et al.67 synthesized VO₂ with a hierarchical flower-like structure by electrochemically improving the activity of argon-treated VO₂ (AVO) microspheres (Fig. 4j-l). The AVO cathode material with a hierarchical flower-like structure and high crystallinity enables fast ion transfer, thus exhibiting fast reaction kinetics. This AZIB achieves a capacity of 446/361 mA h g⁻¹ at 0.1/5 A g⁻¹. He *et al.*⁶⁸ developed high-performance AZIBs by exposing different crystal faces of $VO_2(B)$ to an aqueous solution. By directionally aligning $VO_2(B)$ channels along the *c*-axis, they achieved rapid ion transport and enhanced performance.

3. Energy storage mechanism of VO₂

Research on various energy storage systems focuses on understanding their reaction mechanisms, making the development of AZIBs one of the most significant challenges. VO2-based AZIBs, with their complex electrolyte system and various forms of VO2, involve more intricate reaction processes. During discharge, zinc from the zinc sheet loses electrons, forming Zn^{2+} ions that enter the electrolyte. Subsequently, Zn^{2+} ions migrate to the cathode, inducing the reduction of V^{4+} to V^{3+} upon insertion. In the charging phase, Zn²⁺ ions leave the cathode and return to the electrolyte, while V³⁺ is oxidized back to V⁴⁺. After migration, V⁴⁺ precipitates onto the zinc electrode. There are several types of VO₂-based cathode materials, each with distinct characteristics for zinc storage in AZIBs (Fig. 5). These materials can be categorized into three primary energy storage mechanisms: (1) Zn^{2+} insertion/extraction; (2) H^+/Zn^{2+} co-insertion/extraction; and (3) chemical reaction mechanisms.

3.1 Zn²⁺ insertion/extraction mechanism

Among the new technologies for storing energy, AZIBs have a number of energy storage mechanisms that are similar to



Fig. 4 (a–c) VO_2 nanosheets with an ancient coin structure.⁶⁴ Copyright 2020, Wiley-VCH GmbH. (d–f) Curled VO_2 nanosheets.⁶⁵ Copyright 2023, Elsevier. (g–i) Petal-shaped folded VO_2 nanosheets.⁶⁶ Copyright 2022, Elsevier. (j–l) Hierarchical flower-shaped VO_2 nanosheets.⁶⁷ Copyright 2023, Wiley-VCH.

those of traditional lithium-ion batteries, but also have some differences as well. In AZIBs, VO₂ is considered to be one of the most important types of cathode materials since it exhibits unique energy storage properties. Zinc metal at the negative electrode undergoes an oxidation reaction during the discharge process, resulting in the release of Zn^{2+} accompanied by the release of electrons as a result of the oxidation reaction.⁶⁹ Through the electrolyte, these Zn^{2+} ions migrate to the surface of the VO₂ positive electrode and become embedded in its lattice structure, thus becoming embedded in the lattice structure of VO₂. Vanadium ions (V) in VO₂ are usually partially reduced from a higher valence state (such as V⁴⁺) to a lower valence state (such as V³⁺) during this process, as a result of which a Zn_xVO_2 form is formed, and thus charge storage is

achieved. By passing electrons through an external circuit, electrons flow from the anode to the cathode, converting chemical energy into electrical energy. In contrast, electrons flow through an external circuit, connected to an external power source, from the cathode to the anode during the charging process. The embedded Zn^{2+} ions in the VO₂ cathode are oxidized and re-released, returning to the Zn^{2+} state. After migrating through the electrolyte to the negative electrode surface, these zinc ions are reduced to metallic zinc, completing the charge release process. This process will gradually return the lattice structure of VO₂ to its original state, preparing it for the next discharge.

To investigate the storage behavior of Zn^{2+} in VO₂(B), Ding *et al.*⁷⁰ prepared VO₂(B) nanofibers by a hydrothermal method



Fig. 5 Schematic diagram of three energy storage mechanisms of a VO₂-based cathode for AZIBs.⁶⁹ Copyright 2019, American Chemical Society.⁷² Copyright 2021, Wiley-VCH.⁷⁴ Copyright 2021, Elsevier.



Fig. 6 (a) Crystal structure of VO₂ and battery schematic. (b) Rate performance of VO₂. (c) Charge–discharge curve of VO₂ at 0.5C. (d) *In situ* XRD pattern of VO₂. (e) Schematic diagram of Zn^{2+} embedded in different crystal planes of VO₂.⁷⁰ Copyright 2018, Wiley-VCH.

(Fig. 6a), which showed excellent rate performance when embedded with Zn^{2+} . After cycling the current at a high level for a considerable period of time, it was possible to restore the original capacity (Fig. 6b) when the current was reduced to a small level. Fig. 6c shows a typical charge–discharge curve at 0.5C. *In situ* XRD measurements were performed (Fig. 6d), and

three strong peaks were observed at 45.1°, 49.2° and 59.1°, which correspond to the (-601), (020) and (-404) crystal planes of VO₂(B), respectively. The 2D profiles of the in situ XRD patterns during the charge and discharge process are completely symmetrical, indicating that Zn2+ can be easily embedded and extracted from VO₂(B) nanofibers. Zn²⁺ embedding into VO₂(B) from different crystal planes is represented by the changes in crystal planes (Fig. 6e). There is a slight increase in the lattice spacing of $VO_2(B)$ during complete Zn^{2+} embedding, confirming that the structure of $VO_2(B)$ changes little during the process of Zn²⁺ embedding. During the process of charging from 0.3 V to 1.5 V, these peaks reversibly move back to the initial positions, further proving that the $VO_2(B)$ tunnel has good stability for zinc storage. In the discharge process from about 0.75 V to 0.3 V, a new peak appeared at 58.8°, which is also reversible, coinciding with the appearance of a second redox peak at about 0.65 V in the CV curve. As a result, the electrochemical reaction can be summarized as follows:

Cathode:
$$xZn^{2+} + VO_2 + 2xe^- \leftrightarrow Zn_xVO_2$$
 (1)

Anode:
$$Zn \leftrightarrow Zn^{2+} + 2e^{-}$$
 (2)

Overall:
$$VO_2 + xZn \leftrightarrow Zn_xVO_2$$
 (3)

3.2 H⁺/Zn²⁺ co-insertion/extraction mechanism

In cathode materials, H^+ and Zn^{2+} insert at different sites, resulting in more efficient energy storage and release. The study of the scientific principles and application prospects of the coinsertion/extraction mechanism of H⁺/Zn²⁺ can open up new avenues for the development of battery technology. Nevertheless, further research and exploration are needed to realize this mechanism's practical application.⁷¹ H⁺/Zn²⁺ co-insertion/ extraction is related to different vanadium oxides, and the composition of the electrolyte will also affect chemical reactions. A hydrated vanadium dioxide (Od-HVO@PPy) coating with polypyrrole (PPy) was synthesized by Zhang et al. (Fig. 7a).72 Using a High Resolution Transmission Electron Microscope (HRTEM), Od-HVO@PPy has a lattice spacing of 0.343 nm, which comes from the (110) plane of $VO_2 \cdot xH_2O$ nanocrystals. Vanadium oxide does not dissolve when wrapped in PPy, as shown in Fig. 7b. XPS and EPR measurements show that the Od-HVO@PPy material has an oxygen defect structure (Fig. 7c). The initial specific capacity of the Od-HVO@PPy electrode is as high as 334 mA h g^{-1} at 2 A g^{-1} , which is mainly attributed to the presence of a polypyrrole protective layer to prevent vanadium oxide from dissolving (Fig. 7d). For further investigation of the electrochemical mechanism of Od-HVO@PPy electrodes during cycling, ex situ XRD and XPS characterization was used (Fig. 7e and f). Ex situ XPS characterization reveals symmetry during the charge and discharge processes, indicating that Od-HVO@PPy is reversible during these processes. During the cycle, another significant peak was observed at 12.4°, which was identified as $Zn_4(OH)_6SO_4 \cdot xH_2O$. During the deep discharge process, intensity increased until it reached a maximum value. During the charging process, it decreased until it disappeared in the fully



Fig. 7 (a) Schematic diagram of the synthesis of O_d -HVO@PPy, (b) HRTEM of O_d -HVO@PPy, (c) electron paramagnetic resonance of O_d -HVO@PPy, (d) long-cycle curves of O_d -HVO@PPy and O_d -HVO electrodes at 2 A g⁻¹, (e) *ex situ* XRD of the O_d -HVO@PPy electrode in selected states of the 2nd cycle, and (f) XPS spectra of Zn 2p in the 2nd cycle.⁷² Copyright 2021, Wiley-VCH.

charged state with a high degree of cyclic reversibility. As the OH^- is extracted from water, H^+ participates in embedding, and OH^- forms $Zn_4(OH)_6SO_4 \cdot xH_2O$ with Zn^{2+} .

3.3 Chemical reaction mechanism

During the discharge process, H⁺/Zn²⁺ co-insertion into the VO₂ lattice triggers a change in its chemical structure, thereby generating a new compound. A new material with unique energy storage properties is formed by reshaping the original VO₂ crystal morphology. This new compound, formed by the embedding process, effectively participates in the reversible storage and release cycle of energy during the subsequent charging phase.73 The reversible formation of new products is an important symbol to identify the mechanism of chemical transformation. If the new product can be reversibly formed and disappear during charge and discharge, this is usually due to the H⁺/Zn²⁺ or Zn²⁺ insertion/extraction mechanism. However, if the new product, once formed, directly participates in the subsequent charge and discharge process in this form and does not disappear reversibly, this situation should be classified as a chemical transformation mechanism. The reaction path of VO₂ is a dynamic equilibrium result of the interaction of experimental conditions (ionic environment and preparation process), material properties (crystal phase and surface state) and kinetics (rate and diffusion barrier) and needs to be further verified by specific experiments (such as in situ characterization and crystal relative ratio). Using a sol-gel thermal method, Ding et al.⁷⁴ have prepared VO₂(B) nanorods by hydrothermally treating V2O5 and ethylene glycol for

hydrothermal treatment. XRD measurements under various zinc ion storage states showed that when the zinc ion storage state was charged to 1.4 V, the intensity of the 12.2° characteristic peak began to increase significantly. Upon reaching the maximum value of 1.8 V, the characteristic peak density of $VO_2(B)$ reached its maximum value, which indicates that $VO_2(B)$ was completely converted (Fig. 8a). From the HRTEM images of the original $VO_2(B)$, it can be observed that the lattice spacings of its (110) and (003) planes are 0.352 nm and 0.202 nm, respectively. After being fully discharged to 0.3 V, the lattice spacing of the (003) plane increases to 0.212 nm due to the insertion of some zinc ions. Moreover, the most direct evidence indicating the successful transformation is the emergence of a new lattice spacing of 0.419 nm on the (011) plane of Zn₃(-OH)₂V₂O₇·2H₂O. This is consistent with XRD, again demonstrating the formation of a new product of transformation. According to the charge-discharge curves of the initial two cycles of $VO_2(B)$ nanorods at 0.1 A g⁻¹ in a voltage range of 0.3-1.8 V, a long charging platform is observed when the voltage reaches about 1.5 V for the first time, but disappears with subsequent charging cycles, suggesting that an irreversible conversion reaction has occurred during the first charging process. After the first charge reached 1.8 V, the discharge specific capacity also increased, showing that the active material obtained after the conversion is able to store more zinc ions (Fig. 8b). The reversible capacities of $VO_2(B)$ nanorods are 327, 319, 287, 210 and 130 mA h g^{-1} at 0.5, 1.0, 2.0, 5.0 and 10.0 A g^{-1} , respectively (Fig. 8c). According to the XPS graph of Zn (Fig. 8d), when the electrode is charged to 1.8 V, the signal



Fig. 8 (a) XRD pattern of VO₂(B) during charge and discharge, (b) charge–discharge curves of VO₂(B) at 0.1 A g^{-1} for the 1st and 2nd cycles, (c) rate performance diagram of VO₂(B), (d) XPS of Zn 2p and V 2p under different states of VO₂(B), (e) schematic diagram of the energy storage mechanism of Zn/VO₂(B) during the 1st and 2nd cycles.⁷⁴ Copyright 2021, Elsevier.

peak of Zn 2p is still present. This indicates that zinc-containing compounds were subjected to *in situ* conversion during the initial charging process, and the measured V 2p also corresponds to the embedding of Zn^{2+} in the electrode. As a result of various characterization analyses and electrochemical tests, it was confirmed that the *in situ* conversion reaction product is flaky $Zn_3(OH)_2V_2O_7 \cdot 2H_2O$ (Fig. 8e).

4. Critical issues of VO₂-based cathodes

Due to its excellent electrochemical properties, VO_2 has become a promising AZIB cathode material. Despite the strong potential of VO_2 -based cathode materials, there are still problems such as dissolution, by-product production and ionic diffusion kinetics limitations, which lead to differences between the experimental and theoretical capacity of VO_2 , preventing their widespread commercial application (Fig. 9). These issues primarily arise from the inherent complexities and challenges associated with vanadium oxide itself. To enable the broad use of VO_2 -based cathode materials in AZIBs, extensive research is required to address and overcome these obstacles.

4.1 Dissolution

The crystal structure of VO₂ undergoes significant alterations during the dynamic process of insertion/extraction of Zn²⁺. As Zn^{2+} is inserted into the lattice, the VO₂ structure gradually adapts to accommodate these ions. However, once the amount of embedded Zn²⁺ reaches a certain threshold, the crystal structure can no longer maintain its integrity and stability, leading to collapse and eventual dissolution into the electrolyte. This process is further complicated by the simultaneous embedding of water molecules, which interact with oxygen atoms in the V-O bonds. These interactions create new bonding patterns that weaken the original V-O bonds, destabilizing the crystal structure of VO₂. As the V–O bonds continue to weaken, VO₂ reaches a critical point where it dissolves into the electrolyte due to the continuous embedding of water molecules. Additionally, the charging and discharging cycles can induce phase transitions in VO₂. These transitions significantly alter



g. 9 The main problems and strategies of VO_2 -based cathode materials for AZIBs.

both its crystal structure and chemical properties, further compromising its stability. After phase changes, some VO₂ becomes more soluble in the electrolyte, exacerbating the dissolution issue. Consequently, the insertion and extraction of Zn^{2+} , the incorporation of water molecules, and the phase transition phenomena contribute to the loss of the crystal structure of VO₂, leading to its eventual dissolution.⁴⁶ To enhance the stability and anti-dissolution properties of VO₂, further in-depth research into its crystal structure and chemical properties is essential.

4.2 Generation of by-products

When an aqueous electrolyte comes into contact with VO_2 , a reactive process occurs, leading to the formation of various byproducts depending on the type of electrolyte used. These byproducts, often in the form of hydroxides, salts, and other compounds, can significantly affect the electrochemical performance of VO₂. Initially, the by-products may occupy the active sites intended for Zn²⁺ storage, which reduces the battery's capacity. The deposition of by-products on the electrode surface can also hinder the diffusion of Zn²⁺, impacting the battery's charge and discharge efficiency as well as its cycle stability. As the deposition of by-products continues, they can form a thick passivation layer, which further reduces the electrochemical performance by obstructing successful interaction between Zn²⁺ and VO₂. Current research on AZIBs must focus on addressing these adverse reactions between VO₂ and aqueous electrolytes, with the goal of reducing by-product formation.75 To enhance the overall performance of AZIBs, optimizing the electrolyte composition and electrode structure and exploring new electrode materials are essential areas for future development.

4.3 Limited ion diffusion kinetics

As for the mechanism of reaction in AZIBs, it is certain that whatever mechanism is used, the reaction always involves the

conversion of zinc in the zinc negative electrode to Zn^{2+} , which is then released into the electrolyte, migrates to the positive electrode, and interacts with the cathode. Due to the fact that Zn^{2+} is a divalent cation, it carries two units of positive charge at once. There is a strong electrostatic effect that Zn^{2+} has in an electric field because of this high charge state. Due to this electrostatic effect, it is easy for Zn2+ to interact strongly with negatively charged anions in the electrolyte or electron clouds in the cathode material during the migration process. Zn²⁺ cannot freely diffuse due to this interaction, which acts like an invisible barrier. More complicated, however, is that the lattice structures of different positive electrode materials are unique, and each of these lattice structures corresponds to a significant difference in the diffusion rate of Zn²⁺ within the positive electrode materials. Cathode materials are capable of undergoing phase transitions, lattice strains, or dissolution during a cycle of operation.52,75 Thus, the structure of the cathode material could become unstable, further reducing the diffusion rate of Zn²⁺.

4.4 Others

The inadequate electrochemical performance of cathode materials is also one of the key factors that are hindering the development of AZIBs due to the low electrochemical performance. Due to their poor conductivity and low electrochemical activity, some materials are prone to slow electrode reaction rates and reduced battery charge and discharge efficiency as a result. During the charge and discharge process, it is possible that the cathode material will undergo phase changes, lattice strains, and a variety of other changes that will challenge its structural stability and will further affect the diffusion of Zn²⁺ as well as the ability of the battery to charge and discharge. For the purpose of overcoming these challenges, the structure, composition, and preparation process of the material are optimized to increase the conductivity, electrochemical activity, and structural stability of the material, while reducing dissolution and by-product generation and improving the diffusion rate of

 Zn^{2+} and electrochemical performance of the material. In addition, due to different electrochemical windows, different electrolytes, temperature and other influencing factors, the reported VO₂ cathode capacity varies greatly. Therefore, it is necessary to standardize the test conditions, optimize the electrolyte and electrode design, consider the temperature effect, and conduct a comprehensive comparison and analysis. This will help promote the application and development of VO₂ cathodes in areas such as zinc-ion batteries and realize commercialization at an early date.

5. Optimization strategies of VO₂-based cathodes

The poor electrochemical performance of cathode materials remains a significant challenge for the development of AZIBs. While VO_2 is often considered an ideal cathode material for AZIBs, it still faces several issues, including slow reaction kinetics, low electronic conductivity, structural instability, and vanadium dissolution. To address these challenges, various design strategies have been proposed to enhance the electrochemical performance of VO_2 . These strategies include ion/ molecule pre-intercalation, the use of composite materials, defect engineering, and elemental doping (Fig. 10). By adopting these optimization strategies, the electrochemical performance of VO_2 can be improved. However, it is crucial to select an appropriate strategy based on the specific structure and needs of the system to achieve optimal results.

5.1 Ion pre-intercalation

Electrostatic interactions between Zn^{2+} and cathode materials prevent further insertion and extraction of Zn^{2+} when Zn^{2+} ions



Fig. 10 Schematic diagram of the optimization strategy for VO₂based cathode materials.⁶⁷ Copyright 2023, Wiley-VCH.⁸¹ Copyright 2021, Elsevier.⁹⁸ Copyright 2024, American Chemical Society.¹⁰⁶ Copyright 2023, Elsevier.

are embedded in cathode materials. Additionally, it will cause the material to collapse and dissolve in the electrolyte, resulting in decreased electrochemical performance. Research has shown that the embedding of foreign ions in VO₂ materials not only stabilizes the crystal structure by acting as a "pillar", but also weakens electrostatic interactions during the Zn²⁺ embedding process, preventing the structure from collapsing after Zn²⁺ embedding and extraction, thus improving electrode cycle stability.76,77 The microwave-assisted hydrothermal method was used by Li et al.78 to prepare a nanosphere-cathode containing potassium ions (K^{+}) embedded in VO₂(B). As a pillar, K⁺ has a large ionic radius of 0.138 nm, which greatly improves the diffusion rate of Zn²⁺. The HRTEM image shows a lattice with a spacing of 0.62 nm, which corresponds well to the crystal plane (001) of monoclinic VO₂(B). The nanobelts also showed a lattice fringe of 0.58 nm, which corresponded to the (200) plane of the undoped VO₂(B). Due to the tunnel-like structure of VO₂(B), K^{+} can be accommodated with minimal distortion in the lattice. The battery held a charge capacity of 373 mA h g^{-1} at 0.5 A g^{-1} .

The highly active transition metal Mn, which has a large ionic radius and high activity, is a suitable guest insertion ion.79,80 After expanding the interlayer spacing as tunnel "pillars", the preinserted Mn²⁺ in the VO₂ lattice can store more Zn²⁺ and also provide faster channels and paths. Additionally, metal ions will form oxygen vacancies inside the crystal, which can enhance conductivity and improve redox activity. An oxalic acid solution was added as a reducing agent after dispersing Mn²⁺ in a vanadium solution. MnVO nanobelts with Mn²⁺ pillars were successfully synthesized (Fig. 11a).⁷⁹ As can be seen from XRD, the phases of VO₂ and MnVO are completely consistent with the standard card, indicating that Mn²⁺ ions are doped into the VO₂ bulk phase without creating any new phases. In VO2, Mn2+ ions are embedded in a tunnel structure composed of VO₆ octahedrons with shared edges and oxygen vertices. Zn²⁺ is stored abundantly in an extended tunnel structure that provides a fast transmission channel. Further, Mn²⁺ can improve the conductivity of the VO₂ main phase by optimizing its crystal structure and electron distribution. In comparison to VO₂ as the cathode material for AZIBs, the charge and discharge platform of MnVO is consistent with that of VO₂ (Fig. 11b), with a clear redox platform corresponding to the embedding of Zn²⁺. The MnVO electrode exhibits excellent rate performance (Fig. 11c). Reversible capacities of 432, 418, 390, 364, 338, 281, and 187 mA h g^{-1} can be achieved at 0.1, 0.2, 0.5, 0.8, 1, 2, and 5 A g^{-1} , respectively. Furthermore, they explained the mechanism of insertion and extraction of Zn²⁺ (Fig. 11d). For insertion and extraction, Zn^{2+} ions combine with water to form hydrated zinc ions $(Zn(H_2O)_6^{2+})$. In this combination, the slow kinetics caused by the desolvation process can be suppressed. By optimizing the electronic structure, Mn²⁺ doping also improves the conductivity of the MnVO electrode. In addition, the increased interlayer spacing of MnVO reduces electrostatic interactions, facilitates the rapid transmission of Zn²⁺ within the tunnel, and minimizes irreversible side reactions, resulting in excellent rate and cycle performance. Metal ions can be embedded to maintain charge balance, prevent structural collapse, and improve conductivity. Liu et al.81 introduced Na⁺ and H^+ into VO₂ together. Na⁺ can expand the tunnel, stabilize the



Fig. 11 (a) Schematic diagram of the synthesis of MnVO. (b) Charge–discharge curves and (c) rate performance curves of MnVO and VO₂. (d) Schematic diagram of Zn^{2+} insertion/extraction of MnVO.⁷⁹ Copyright 2023, American Chemical Society. (e) Schematic diagram of the synthesis of Ni_{0.006}Ca_{0.0045}VO₂@C. Structures of (f) Ni_{0.006}Ca_{0.0045}VO₂@C, (g) δ -(Ni,Ca)V₂O₅@C, and (h) Zn₃(OH)₂V₂O₇·3H₂O. (i) CV curve of Ni_{0.006}Ca_{0.0045}VO₂@C, (j) cycle diagram of Ni_{0.006}Ca_{0.0045}VO₂@C, (k) schematic diagram of the structure of a flexible quasi-solid-state battery, and (l) cycle performance diagram of the solid-state battery.⁸² Copyright 2022, Elsevier.

crystal form, and prevent structural collapse when pre-embedded. The structural water surrounding Zn^{2+} also plays an important role, as it can act as a buffer to reduce electrostatic interactions between the ions. With its more stable crystal, it is capable of having an initial capacity of 156 mA h g⁻¹ at 12 A g⁻¹ and maintaining a capacity of 118 mA h g⁻¹ after 3000 cycles.

The role of a single metal cation is often insufficient to maintain the stability of the material, and thus double metal cations are embedded into VO₂ through the pre-intercalation method in order to improve its electrochemical performance. The calcium ions have a larger ionic radius, which facilitates the expansion of interlayer spacing and the stabilization of crystal structures. Nickel ions can not only support a material but also enhance its conductivity. Zhao et al. synthesized double cation pre-intercalated vanadium dioxide (Ni_{0.006}Ca_{0.0045}VO₂@C) (Fig. 11e).⁸² XRD analysis of the prepared material indicated that VO2 was monoclinic in nature. The XRD spectrum revealed no peaks attributed to carbon, indicating that VO₂ and amorphous carbon were successfully prepared. Ca²⁺ and Ni²⁺ were successfully inserted, causing a slight positive shift in 2θ , indicating an enlarged interlayer. The first cycle CV curve of Ni_{0.006}Ca_{0.0045}VO₂@C shows an irreversible oxidation peak at 1.60 V, indicating that the vanadium(IV)-based oxide is in situ converted to a V₂O₅-based double-layer structure (Fig. 11f-h). The δ -V₂O₅-based double-layer structure gives the

Ni_{0.006}Ca_{0.0045}VO₂@C electrode a higher specific capacity (433.8 mA h g^{-1} at 0.1 A g^{-1}) (Fig. 11i). Ni_{0.006}Ca_{0.0045}VO₂@C electrodes have a reversible capacity of 115 mA h g^{-1} at 5 A g^{-1} and can maintain 74 mA h g^{-1} after 4000 cycles (Fig. 11j). Under various charge and discharge conditions, the zinc storage mechanism of the Ni0.006Ca0.0045VO2@C electrode was investigated by ex situ XRD. After the first charge to 1.4 V, vanadium(IV)-based oxide remains the main vanadium species. Upon initial charging to 1.6 V, the characteristic peaks of the VO₂phase disappear, while those of the V₂O₅ phase emerge, coinciding with the charge plateau near 1.55 V. This observation confirms the effective phase transition achieved through electrochemical activation. In subsequent cycles, the zinc ion storage mechanism primarily involves reversible Zn²⁺/H⁺ coinsertion/extraction in the host lattice of double-layer structured δ-V₂O₅. During the charge-discharge process, a new phase of $Zn_xV_2O_5 \cdot nH_2O$ was also detected due to the conversion reaction and H₂O co-insertion behavior. Fig. 11k and l show its application to a flexible quasi-solid-state battery. The first discharge specific capacity was 158 mA h g^{-1} , and it remained at 73.2% after 100 cycles, indicating long-term cycle stability.

5.2 Molecule pre-intercalation

By introducing molecules into the VO_2 material matrix in advance, as opposed to directly embedding cations, the intrinsic

crystal structure of VO₂ and the Zn²⁺ transport kinetics can be significantly regulated by forming chemical bonds or coordination with the main structural framework. Molecule embedding plays a crucial role as a "lubricant" in this process. Besides expanding the interlayer distance and creating favorable conditions for Zn²⁺ migration, it can also reduce the electrostatic interaction strength between the embedded Zn²⁺ and the active material by acting as a charge shielding layer. A mixture of NH₄VO₃ and polyacrylamide is combined with hydrochloric acid and hydrogen peroxide. After hydrothermal treatment, $VO_2 \cdot xH_2O$ and (NH₄)₂V₁₀O₂₅·8H₂O can be obtained (Fig. 12a).⁸³ Both VO₂- $\cdot xH_2O$ and $(NH_4)_2V_{10}O_{25}\cdot 8H_2O$ have tunnel structures (Fig. 12b), while the (001) plane of $(NH_4)_2V_{10}O_{25} \cdot 8H_2O$ has a larger lattice spacing at a smaller angle, which may lead to its structural instability. In SEM, (NH₄)₂V₁₀O₂₅·8H₂O exhibits a cross-linked and unclear nanobelt structure, while VO2 · xH2O exhibits a 10-30 nm nanobelt structure. $VO_2 \cdot xH_2O$ nanobelts possess excellent order and can be interwoven to form a stable nanonetwork structure, which is beneficial for Zn^{2+} transport and storage. According to TG, each VO₂ contains 0.35 grams of crystalline water. These water molecules act as a lubricant, weakening the electrostatic interaction between Zn2+ and the active material and enhancing transport. The $VO_2 \cdot xH_2O$ electrode can still store 376 mA h g⁻¹ after 200 cycles at 1 A g⁻¹. The VO₂ \cdot *x*H₂O electrode retained specific capacity after 2000 cycles at 5 A g^{-1} was 289 mA h g^{-1} (Fig. 12c). Fig. 12d depicts full and enlarged colormap 3D waterfall plots with the projection of *in situ* XRD of the $VO_2 \cdot xH_2O$ electrode during the first three cycles. The diffraction peak at $2\theta = 14.4^{\circ}$ is significantly shifted during discharge due to the insertion of Zn²⁺ and the decrease of electrostatic bond strength, which increases the lattice spacing. During the subsequent charging process, the diffraction peak of $VO_2 \cdot xH_2O$ undergoes the opposite process. The in situ XRD test results of the $VO_2 \cdot xH_2O$ electrode show that its zinc storage process is based on



Fig. 12 (a) Schematic diagram of the synthesis of the $VO_2 \cdot xH_2O$ material, (b) XRD pattern of $VO_2 \cdot xH_2O$, (c) long-cycle curve of the $VO_2 \cdot xH_2O$ electrode at a current density of 5 A g⁻¹, (d) the full and enlarged corresponding 3D colormap surface with the *in situ* XRD pattern of the $VO_2 \cdot xH_2O$ electrode, (e) line spectrum from the result of *in situ* XRD of the $VO_2 \cdot xH_2O$ electrode and corresponding discharge/charge curve.⁸³ Copyright 2022, the Royal Society of Chemistry.

the H^+/Zn^{2+} co-insertion/extraction (Fig. 12e). During the discharge process, H⁺ and Zn²⁺ will synergistically embed into the main structure of $VO_2 \cdot xH_2O$, and the increase in OH^- concentration in the electrolyte will lead to the formation of $Zn_x(CF_3SO_3)_y(OH)_{2x-y} \cdot nH_2O$. There are two phases in the charging process that are responsible for the extraction of H⁺ and Zn^{2+} from the main structure of $VO_2 \cdot xH_2O$ into the electrolyte, resulting in the decomposition of by-products. The energy storage mechanism revealed that some by-products $Zn_x(CF_3SO_3)_v(OH)_{2x-v} \cdot nH_2O$ were produced during discharge, but the by-products underwent reversible reactions during the cvcle.

A vanadium-based oxide containing organic molecules exhibits excellent properties in terms of energy density, structural stability, and conductivity. Ye's group successfully prepared monoclinic VO2 intercalated with polyaniline (PANI) using a simple hydrothermal method.⁸⁴ L-PANI-VO₂ and PANI-VO₂ are the compounds prepared by adding 0.16 and 0.32 mL of hydrochloric acid, respectively. The VO₂ crystal structure can be optimized, with the layered L-PANI-VO₂ exhibiting an interlayer spacing of 10.2 Å. A larger interlayer spacing allows for the embedding of more Zn2+ ions. The specific capacity of the L-PANI-VO₂ electrode can reach 415 mA h g^{-1} at 0.1 A g^{-1} . Moreover, the intercalation of the conductive polymer enhances the electronic cloud structure of the material, which improves the diffusion rate of Zn²⁺ ions. This, in turn, boosts the charge/ discharge rate performance of the battery. The specific capacity of L-PANI-VO2 is 415, 380, 365, 340, 320, 290, 270 and 175 mA h g⁻¹ at 0.1, 0.2, 0.3, 0.5, 1.0, 2.0, 3.0 and 5.0 A g⁻¹, respectively. The SEM image reveals that L-PANI-VO₂ possesses a layered structure with well-defined edges and a smooth surface.

The L-PANI–VO₂ sheets typically measure over ten microns in length and range from 100 to 200 nm in thickness. The CV curve of the L-PANI–VO₂ composite material displays four pairs of redox peaks at 0.35/0.48, 0.55/0.70, 0.92/1.04, and 1.32/1.43 V, which correspond to the insertion and extraction of Zn^{2+} ions. This indicates a multi-step charge storage process at a low scan rate. In contrast, the CV curve of the PANI–VO₂ composite material shows only two pairs of redox peaks. Electrochemical testing further confirms that the L-PANI–VO₂ composite material exhibits a significantly higher specific capacity compared to the PANI–VO₂ composite material, as indicated by the larger peak areas in the cyclic voltammetry curves.

5.3 Composites with other materials

To prevent VO₂ crystals from changing shape and dissolving, they can be coated with carbon-based materials, metal oxides, conductive polymers, etc. Long-term cycle stability is achieved by stabilizing the crystals and preventing volume changes caused by ion insertion and extraction during charging and discharge. The abundance of free electrons inside the carbon coating can improve conductivity, which facilitates the fast transfer of electrons. Additionally, in some cases, carbon coatings have been shown to reduce the interfacial resistance between the cathode and the electrolyte, thereby increasing the overall conductivity of the system. By calcining organic matter in a specific atmosphere and retaining a carbon layer after the calcination process, the carbon coating can be obtained.85-87 For carbon coating, acetylene (C_2H_2) gas with a high carbon content is a good raw material. VO2 can be calcined in a mixture of C2H2 and argon to produce VO2@C nanobelts with different microstructures (Fig. 13a).⁸⁸ As can be seen from the SEM, there is



Fig. 13 (a) Synthesis route and structural diagram of VO₂@C and (b) VO₂@C cycle performance diagram.⁸⁸ Copyright 2023, the Royal Society of Chemistry. (c) Synthesis route of N–VO₂@NC@CNTF, (d) CV diagrams at different scan rates and (e) rate performance of N–VO₂@NC@CNTF// Zn@CNTF.⁹⁰ Copyright 2023, Wiley-VCH. (f) Synthesis route of H–VO₂@GO, and (g) Schematic diagram of the H–VO₂@GO reaction mechanism⁹² Copyright 2024, Elsevier. (h) Synthesis diagram of VMC and (i) rate performance of the VMC electrode.⁹⁹ Copyright 2024, Elsevier.

a coating layer of about 2 nm on the VO₂ nanobelt. According to further analysis, the coating is a carbon layer, and the crystal structure of VO₂ has not changed significantly. The addition of oxygen vacancies to the AZIBs through carbon coating also increases active sites, contributing to higher electrochemical performance compared to AZIBs with pure oxygen. After low current activation, VO2@C nanobelts can achieve an initial capacity of 438/342 mA h g^{-1} at 1/2 A g^{-1} (Fig. 13b). Moreover, He's group successfully prepared carbon-coated VO₂ (H_{1.99}VO@C) by a hydrothermal reaction of VO₂ with glucose and subsequent annealing.89 Carbonized carbon is provided by glucose to achieve the coating of rod-shaped VO₂ with a carbonized carbon shell. Carbon shells can improve conductivity and mechanical properties. H1.99VO@C electrodes have a specific capacity of 425 mA h g^{-1} at 0.1 A g^{-1} .

Due to their excellent conductivity, carbon nanofibers have been widely used in energy storage devices in recent years. The use of these materials can enhance electron transmission capacity, reduce battery internal resistance, and increase the power density and charging efficiency of batteries. During the charge/discharge process, the volume change of the cathode material can cause stress, which can be relieved with the high strength and toughness of carbon nanofibers, which also prevent the collapse of the cathode material, improving the battery's cycle stability. Carbon nanofibers, in addition to having a high specific surface area, also increase the contact area between the positive electrode material and the electrolyte. Increasing the contact area increases the number of active sites, which increases the capacity and energy density of the battery. N-VO2@NC@CNTF was obtained after calcination by Guo et al.90 through growing nitrogen-doped vanadium dioxide on carbon nanotube fibers (CNTFs) (Fig. 13c). During calcination, the polypyrrole (PPy) deposited on CNTF releases ammonia, which results in the introduction of N. On CNTFs, densely deposited PPy nanowires are observed. PPy nanowires enhance the conductivity of carbonized nanocomposites and provide conductive scaffolds for regulating the loading of the active materials in the outer layer. Even after calcination, scanning electron microscopy images show that N-VO2@NC@CNTF retains its structural integrity and the unique characteristics of high porosity and a large specific surface area. They assembled an entangled quasi-solid FAZIB with the N-VO2@NC@CNTF cathode and Zn@CNTF anode coated with ZnSO4/CMC gel electrolyte. Charge-discharge curves demonstrate the superiority of the NC NW scaffold and N doping, where VO2@-NC@CNTF//Zn@CNTF is significantly more specific than VO2@CNTF//Zn@CNTF, and N-VO2@NC@CNTF//Zn@CNTF is significantly more specific than VO₂. Moreover, the CV curves of the FAZIB at different scan rates clearly show that the curve has two pairs of oxidation and reduction peaks, which are caused by the multi-step insertion/deinsertion mechanism for Zn²⁺ (Fig. 13d). The specific capacity of the FAZIB is 441.383 mA h cm^{-3} at 0.2 A cm^{-3} . This indicates that the FAZIB has excellent rate performance due to its improved conductivity (Fig. 13e). The advantages of carbon nanotubes are also similar. The unique flower-like structure of the $\ensuremath{\text{VO}}_2$ and $\ensuremath{\text{CNT}}$ composites facilitates the contact between the electrode and electrolyte,

thereby reducing the diffusion distance of Zn^{2+} . The VO₂/CNT composites exhibit high reversible capacity (410 mA h g⁻¹ at 0.1 A g⁻¹), excellent cycling stability (reversible capacity of 221 mA h g⁻¹ after 2000 cycles at 5 A g⁻¹), and good rate capability.⁹¹

The graphene oxide (GO) material is a novel type of carbon material that exhibits excellent performance. GO contains many oxygen-containing functional groups, including epoxide, carbonyl, carboxyl, and hydroxyl, which can be reduced to obtain reduced graphene oxide (rGO). Although some oxygencontaining functional groups are reduced in graphene, its partial conjugated structure is restored, making it conductive. As rGO can be well compounded with metal oxides, it is added to VO2 to achieve carbon coating and defect formation. Li et al.92 used a hydrothermal method and high-temperature calcination method to pre-prepare hollow VO2 microspheres and then composited them with GO to form redox graphene-anchored hollow VO₂ microspheres. The hollow morphology of microspheres provides a large specific surface area and a large number of electrochemical reaction sites, enabling them to store energy efficiently and resist volume stress during Zn²⁺ deintercalation, which greatly enhances their microstructural stability. Additionally, GO enhances the material's conductivity and increases its ion diffusion rate. The 400 nm nanospheres (S-VO₂) exhibit obvious lattice fringes. After calcination, a hollow sphere structure (H-VO₂) is formed, which reduces the crystallinity, providing more active sites for ion storage (Fig. 13f), making it a highly efficient electrochemical material. As a result, the specific discharge capacities of the H-VO₂ electrode are 400.1/261.5 mA h g^{-1} at 0.5/10 A g^{-1} , respectively. Ex situ testing was conducted to investigate the energy storage mechanism of the H-VO₂ electrode. Zn₃(OH)₂V₂O₇·2H₂O is formed during the charge and discharge process of the first cycle and participates in subsequent energy storage (Fig. 13g). Directly dispersing VO₂(B) powder on rGO sheets can also achieve carbon coating.93,94 Polystyrene nanospheres and redox graphene nanosheets were also added to the spray solution, and spherical electrode materials were formed by spray pyrolysis. VO2 nanosheets were anchored in the layers of redox graphene microspheres, resulting in a composite of rGO and vanadium oxide (P-VO2@rGO). The P-VO2@rGO electrode can achieve a specific capacity of 342 mA h g^{-1} at 0.1 A g^{-1} .⁹⁵ In addition to redox graphene, expanded graphite with good conductivity and stability can also be composited with VO2.96

As an alternative coating material for VO₂, MXene is a 2D transition metal carbide, nitride, or carbonitride that has excellent electrical conductivity, outstanding mechanical properties, and adjustable surface end groups.⁹⁷ The most troublesome thing about MXene, however, is the fact that it needs to be pretreated before use.⁹⁸ Wang's group successfully used MXene nanosheets as conductive adhesives for anchoring VO₂·*x*H₂O nanorods and single-walled carbon nanotubes (SWCNTs) (Fig. 13h).⁹⁹ Compared to the traditional method of scraping materials on flexible substrates, this technique provides a flexible and stable electrode by preventing the active material from falling off as happens with traditional scraping electrodes when repeatedly deformed under repeated deformation conditions.

Chemical Science

SEM images reveal that VMC consists of interconnected onedimensional (1D) nanorods, 1D nanotubes, and 2D nanosheets. The microstructure can be revealed through SEM images of VMC cross sections. VO2 · xH2O nanorods are preferentially distributed on MXene sheets by van der Waals interactions, forming a hierarchical structure. Additionally, SWCNTs provide mechanical support and flexibility for the cathode, facilitating rapid charge transfer and long-term cycling. In combination with VO₂, this low-dimensional carbon material can improve conductivity and increase active sites, thereby improving electrochemical performance and rate performance (Fig. 13i). The VMC electrode exhibits an ultrafast ion diffusion rate $(10^{-10} \text{ to } 10^{-7} \text{ cm}^2 \text{ s}^{-1})$. This is mainly due to the unique 3D network structure constructed from MXene sheets, $VO_2 \cdot xH_2O$ nanorods and SWCNTs, which provides an efficient electron transmission path and ion diffusion channel.

In various battery systems, atomic layer deposition (ALD) is one of the most effective methods to solve the interface problems of electrodes because it is one of the most precise coating methods.¹⁰⁰ ALD layers cannot only protect the electrode from adverse side reactions at the interface in order to inhibit cathode dissolution, but can also serve as an artificial cathode electrolyte interface to facilitate the transfer of Zn²⁺ between the electrode and electrolyte. Boruah et al.¹⁰¹ have applied a layer of ZnO to a carbon fiber (CF) current collector as a hole and electron transport layer. Growing VO2 directly on the ZnO layer improved charge separation and the interface between the active materials, resulting in an increase in light conversion efficiency. Shuai et al.¹⁰² inhibited vanadium dissolution and side reactions using an ALD zinc oxide layer (ZnO@VOH) on a hydrated vanadium dioxide nanosheet cathode. The XRD analysis showed that the original crystal structure did not change much after deposition on ZnO. In electrochemical tests, the ZnO@VOH electrode retained 74% to 89% of its capacity after 100 cycles at 0.5 A g^{-1} and 29% to 71% after 990 cycles at 5 A g⁻¹. An XRD analysis of the ZnO@VOH electrode at different cycle numbers revealed a very stable crystal structure, indicating that the material is cycle stable. By comparing the morphology of the ZnO@VOH electrode in the original state and after 20 cycles at 0.2 A g^{-1} , it was found that the morphology of the nanosheets was well preserved, further confirming that the ZnO on the surface protected the vanadium oxide. Moreover, Cheng's group has successfully prepared a material containing a heterojunction structure by a microwave-assisted hydrothermal method, which is composed of bismuth vanadate (BiVO₄) and vanadium dioxide (VO₂).¹⁰³ Due to the dual-phase structure, the interlayer spacing is enhanced, more pathways are provided for Zn²⁺ transport, and excellent stability is ensured, resulting electrochemical in outstanding performance.

5.4 Defect engineering

By adjusting the electronic structure, increasing active sites, optimizing ion transport paths, and improving structural stability, it is possible to improve the electrochemical performance of electrode materials. Traditional defect engineering

includes oxygen defects and vacancy defects, with oxygen defect engineering being widely used in vanadium oxides. Introducing oxygen vacancies into vanadium oxides modifies the electronic structure, band structure, charge distribution, and density of states. This alteration enhances conductivity, boosts activity, and improves energy storage capabilities, while also inhibiting undesirable phase transitions. Unfortunately, defects can alter the reaction kinetics of a material by acting as an active site that promotes or hinders oxidation reactions. They also affect the stability of the material, which can lead to faster degradation and reduced electrochemical performance over time. Therefore, it is important to consider appropriate deficiencies and develop strategies to minimize their adverse effects. Cao's group used multiple phase transition methods to successfully prepare VO₂(B) rich in oxygen vacancies.¹⁰⁴ Due to oxygen defects increasing the active site and improving ion diffusion and transport, the oxidation reaction rate of VO₂ during charging may be increased. This means that in the same amount of time, more VO₂ can be oxidized to higher-priced vanadium ions. As a result, it showed a discharge capacity of 325.6 mA h g^{-1} at 0.05 Ag^{-1} , while the capacity retention rate was 86% after 5000 cycles at 3.0 A g⁻¹. As graphene exhibits excellent electrical conductivity, Li et al. created oxygen vacancies in VO2/G nanobelts by attaching vanadium oxide to redox graphene.105 Incorporating graphene can maintain vanadium oxide's morphology and increase the diffusion rate of Zn^{2+} by accelerating charge transfer. Yuan et al. also combined VO2 with GO through a hydrothermal route to construct a vanadium dioxide/GO composite material rich in oxygen vacancies (VO2(B)/GO).106 VO₂(B)/GO showed excellent electrochemical performance (423/ 227 mA h g^{-1} at 0.5/15 A g^{-1}). The one-step hydrothermal method to in situ prepare defective VO2 is economical and effective in comparison with most methods that synthesize VO₂ first and then introduce defects. Ascorbic acid and V2O5 can be hydrothermally treated to produce VO₂ rich in oxygen defects $(O_d$ -VO₂). By adjusting the amount of ascorbic acid, oxygen vacancies can be controlled. Od-VO2 presents a layered stacking structure in SEM. The Od-VO2 electrode exhibits excellent zinc storage properties (92.6% capacity retention after 2000 cycles at 3.0 A g^{-1} .¹⁰⁷

Nanostructures provide high surface areas, short ion diffusion paths, and porous structures that facilitate electrolyte diffusion and maintain volume changes during the insertion and extraction of ions. The use of graphene sheet nanocomposites can enhance cycle stability and rate performance at high currents. By weakening the strong interaction between Zn²⁺ and the vanadium oxide skeleton, a lattice defect can enhance zinc ion transport. As a result of the above three optimization strategies, Huang et al.¹⁰⁸ were able to fabricate 3D sponge-like hydrated vanadium dioxide composites that contain abundant oxygen vacancy defects and graphene that has been modified by one-step hydrothermal treatment. In Fig. 14a, 2D hydrated vanadium dioxide nanorods with oxygen vacancies are intertwined with rGO nanosheets to form a nanowall (Od-HVO/rG). The rGO nanosheets cover a large number of hydrated vanadium dioxide nanorods with oxygen vacancies, preventing them from stacking. Due to their 3D



Fig. 14 (a) Schematic diagram of the synthesis of O_d -HVO/rG materials, (b) XRD patterns of all samples, (c) TGA curve of O_d -HVO/rG, (d) rate performance of HVO, O_d -HVO and O_d -HVO/rG, (e) *ex situ* XRD of O_d -HVO/rG, and (f and g) schematic diagram of the energy storage mechanism of O_d -HVO/rG.¹⁰⁸ Copyright 2021, American Chemical Society.

structure, sponges are less susceptible to structural collapse and damage during electrochemical reactions. The XRD spectra of HVO, Od-HVO, and Od-HVO/rG are consistent with those of VO₂·xH₂O. O_d-HVO and O_d-HVO/rG show weaker intensities than HVO, indicating that oxygen vacancy defects have decreased the crystallinity of Od-HVO and Od-HVO/rG. The thermogravimetric analysis in Fig. 14c indicates that a 2.1% weight loss occurs when adsorbent water evaporates at 100 °C; when the temperature is between 100 °C and 500 °C, it is 2.7% of the crystal structure water loss, and 4.5% of the weight loss, which can be converted to 11.8% for reduced graphene. In addition to increasing capacity, oxygen vacancies in the Od-HVO/rG electrode can reduce electrostatic interactions between Zn²⁺ and the host material. rGO enhances conductivity, suppresses crystal distortion, and achieves high stability in the Od-HVO/rG electrode. Od-HVO/rG electrodes exhibit a high initial charge capacity of 428.6 mA h g^{-1} at 0.1 A g^{-1} (Fig. 14d). The energy density and power density of Od-HVO/rG are compared with those of other reported cathodes in Fig. 14c. Od-HVO/rG exhibits a superior energy density (245.3 W h kg⁻¹) and power density (71 W kg⁻¹). To clarify the reaction mechanism

and structural evolution of the Od-HVO/rG electrode during cycling, an ex situ XRD analysis was performed (Fig. 14e). A number of obvious characteristic peaks were observed during the discharge process of the O_d-HVO/rG electrode at 8.1°, 16.2°, 24.4° and 29.3°, which can be attributed to the characteristic peak of $Zn_4SO_4(OH)_6 \cdot 5H_2O$. After charging at 0.4–1.4 V, Zn_4 - $SO_4(OH)_6 \cdot 5H_2O$ gradually disappeared, indicating that Zn_4 - $SO_4(OH)_6 \cdot 5H_2O$ was reversibly and continuously generated and decomposed. The embedding and spacing reduction of Zn²⁺ during the discharge process resulted in a noticeable diffraction peak shift on the (110) crystal plane of Od-HVO/rG. As a result of charging, the diffraction peak returned to its original position, indicating that Zn²⁺ had been removed. Od-HVO/rG contains continuously precipitated H⁺ and Zn²⁺, as well as OH⁻ that is formed when water decomposes (Fig. 14f and g). A special atmosphere can also be used to produce oxygen vacancies in defect engineering by calcining vanadium oxides with special substances. To create oxygen vacancies, Liu et al.109 used hydrothermal method to grow VO₂ nanorods on polydopamine-coated gourd-shaped carbon and then calcined them with sodium borohydride under an argon atmosphere.

The uniform gourd carbon matrix can enhance electronic conductivity by supplying additional Zn^{2+} storage sites and weakening the electrostatic interaction between Zn^{2+} and VO₂.

5.5 Elemental doping

Different from ion pre-intercalation, elemental doping here mainly refers to the doping of mostly non-metallic elements and a small proportion of metallic elements. These non-metallic elements replace the original position of oxygen, which not only changes the electronic structure and spatial configuration of the crystal, but also accelerates electron/ion dynamics, thereby playing a crucial role in improving the material. Metal doping replaces the original position of V, increases the lattice parameters of the material, enhances its thermal and chemical stability, and thus extends the service life of the battery. In AZIBs, the low conductivity of $VO_2(B)$ and the slow migration rate of Zn²⁺ hinder its application. The crystal structure and electrochemical performance of vanadium oxide can be improved by nitrogen doping. By calcining tunnel-type $VO_2(B)$ in an ammonia atmosphere, Gu et al. prepared nitrogen-doped VO₂ (Fig. 15a).¹¹⁰ Heat treatment at different temperatures in NH₃ preserved the VO₂ nanobelt structure well. As the calcination temperature increased, the edges of the VO₂ nanobelts gradually melted and aggregated. As a result of hightemperature reactions, nanobelt aggregates were also confirmed by TEM. The HRTEM images of the material (VO₂-N400) at 400 °C revealed lattice fringes with a spacing of 0.353 nm, which corresponds to the (110) plane of $VO_2(B)$. The lattice of nitrogen atom incorporation also exhibited a variety of disordered structural regions. VO2-N400 exhibits excellent rate performance and cycle stability due to the introduction of N (Fig. 15b). Structural and electrochemical characterization studies have demonstrated that the introduction of N into $VO_2(B)$ results in lattice expansion, grain size reduction, and partial structural disorder. The electrochemical performance of $VO_2(B)$ in AZIBs is superior to that of this product due to its fast ion transport, additional ion storage sites, and inhibition of structural degradation. The VO₂-N400 electrode has a specific capacity of 373.7 mA h g⁻¹ at 0.1 A g⁻¹ (Fig. 15c). *Ex situ* XRD characterization was conducted to determine how Zn²⁺ ions are stored in VO₂-N400. The ex situ XRD pattern of the VO₂-N400 electrode in the first cycle revealed four monoclinic VO₂(B) peaks at 25.2°, 30.0°, 35.8°, and 44.9°. These peaks suggest that VO2-N400 has good reversibility and structural stability for the storage of Zn²⁺. In addition, a new phase, Zn₂(OH)₂V₂O_{5.72}- $N_{1.28}{\cdot}2H_2O$ (12.4° and 22.7°), was detected, which appeared with the insertion of Zn²⁺ and disappeared with the extraction of Zn²⁺, indicating that the zinc storage process is based on the H⁺/Zn²⁺ co-insertion/extraction.

As opposed to nitrogen doping, metal atoms (such as tungsten, molybdenum, and niobium) can be introduced into the center of the lattice polyhedron to replace vanadium atoms, thus improving the crystal structure and performance.¹¹¹ A hydrothermal method was used by Li *et al.*¹¹² to dope tungsten (W) into VO₂ crystals. Using tungstic acid (H₂WO₄) and ethylene glycol, ammonium metavanadate (NH₄VO₃) was first



Fig. 15 (a) Schematic diagram of the synthesis of VO₂–N nanobelts, (b) long cycle performance curves of VO₂–N400 at 2/5 A g⁻¹, and (c) cycle performance curves of VO₂–N with different calcination temperatures at 0.1 A g^{-1,110} Copyright 2023, Elsevier. (d) Schematic diagram of the synthesis of W–VO₂, (e) XRD patterns of VO₂ and W– VO₂, and (f) contribution ratio between capacitance and diffusioncontrolled capacitance.¹¹² Copyright 2021, American Chemical Society.

decomposed to form V₂O₅ nanoparticles, which were then selfassembled into a three-dimensional network structure (W- VO_2). W-VO₂ content was controlled by the concentration of tungstic acid in the precursor solution. Ultrathin nanosheets of W-VO₂ were observed. The thickness of W-VO₂ was significantly lower than that of pure VO_2 (nanoparticles). On the ultrathin nanosheets of W-VO₂, mesopores can be seen due to the introduction of tungsten atoms and the generation of a large amount of gases (such as NH_3 and CO_2). The XRD pattern of W-VO₂ shows that W doping does not alter the VO₂ crystal phase diagram (Fig. 15e). Interestingly, the addition of W does not cause a significant shift in the diffraction peak but rather increases its intensity, which is explained by the low W doping content of W-VO₂, which does not cause a noticeable shift. However, introducing W leads to stronger V-O bond stretching vibration, so the intensity increases. When W-VO2 is tested in the voltage range of 0.2-1.4 V, an incomplete CV curve can be seen during the first cycle, which may be attributed to the

relatively low open circuit voltage. The pseudocapacitive effect becomes more obvious as the scan rate increases, indicating that the electrochemical behavior of the W-VO₂ electrode is primarily influenced by the pseudocapacitive effect (Fig. 15f). In W-VO₂, the stronger W-O bond results in a weaker electrostatic interaction between zinc and oxygen, which is why W-VO₂ is the preferred positive electrode for AZIBs because of its superior zinc storage performance. At 0.1 A g^{-1} , W-VO₂ has an initial reversible capacity of 346 mA h g^{-1} ; it retains 281.3 mA h g^{-1} after 100 cycles. To investigate the zinc storage mechanism in W-VO2, ex situ XRD and XPS characterization studies were conducted. In the ex situ XRD pattern, the peak at 25.6° shifted slightly downwards, corresponding to the expansion in the distance between VO2 interlayers during discharge, which could be explained by the embedding of zinc atoms, which resulted in a decrease in the distance between W and VO₂. As shown by the ex situ XPS spectrum of V 2p, the peak changes of V⁴⁺ and V³⁺ are consistent, indicating the stability of the cycle.

The material structure can be intrinsically adjusted through doping strategies in order to enhance performance. The dopant atoms are not introduced into the center position of the lattice polyhedron. Incorporating dopant atoms into the gaps of the lattice polyhedron can increase tunnel size and stabilize the lattice at the same time. The element boron (B) usually exists as B^{3+} , which is small and low in atomic weight, so using B element for interstitial doping makes sense.¹¹³ Through the use of H₃BO₄ as a precursor, an interstitial boron-doped tunnel-type $VO_2(B)$ was successfully prepared by a hydrothermal method. The XRD spectrum of boron-doped VO₂(B) shifts to the left because of the interstitial doping of boron. The lattice spacing of the (110) plane of the boron-doped product increases to 0.012 nm as boron occupies the interstitial position and forms boron-oxygen bonds, which cause oxygen atoms to shrink and the lattice spacing to increase. The specific capacity of borondoped VO₂(B) is 281.7 mA h g^{-1} at 0.1 A g^{-1} . The tunnel-type $VO_2(B)$ retains a capacity of 133.3 mA h g⁻¹ after 1000 cycles at 5 A g^{-1} , indicating that interstitial boron doping can improve its Zn²⁺ storage capacity. To verify the universality of boron doping, different concentrations of boron were added to tunneltype MnO₂. It is interesting to note that boron doping is universally used to increase zinc storage capacity in manganesebased products.

6. Conclusion and practical prospects

Recent research on VO₂-based cathode materials for AZIBs has shown remarkable progress. When the zinc anode and electrolyte are optimized to match each other, this battery system could serve as a promising candidate for large-scale applications. However, like all metal oxide cathode materials, VO₂ still faces significant challenges such as dissolution, byproduct formation, and slow diffusion rates, which hinder its widespread use. Various modification strategies have been employed to enhance the electrochemical performance of AZIBs, including ion intercalation, molecular intercalation, composite materials, and defect engineering. Ion intercalation stabilizes the cathode's structure and improves its kinetics, while



Fig. 16 Schematic diagram of the strategy for the application of VO_2 -based cathodes in AZIBs.

molecular intercalation enhances the overall conductivity and further stabilizes the crystal structure. The use of composite materials can help prevent the collapse of the cathode structure, improving both conductivity and electrochemical properties. Moreover, composite materials offer a new avenue for developing more effective cathode materials. Defect engineering plays a critical role in enhancing the structural stability and electrochemical performance of VO₂-based cathodes. By introducing or controlling defects, it is possible to modify the crystal structure of the material to improve its properties. Additionally, elemental doping can optimize electrochemical performance by tuning the energy band structure and enhancing the ion diffusion rate, achieved by incorporating foreign elements into the VO₂ crystal lattice.

 VO_2 , as a cathode material for AZIBs, necessitates comprehensive research and innovation from various perspectives to reach commercialization and further broaden its application potential. This review highlights key areas for future exploration, including the innovation of synthesis methods, enhancement of optimization strategies, refinement of electrolytes, and the development of advanced characterization techniques (Fig. 16).

6.1 In-depth exploration of the VO₂ synthesis strategy

Currently, VO_2 is mainly synthesized through redox reactions starting from V_2O_5 . However, it must be noted that this traditional method may suffer from several disadvantages such as high energy consumption, low yield, or insufficient purity of the final product. As a consequence, to improve the performance of VO_2 and reduce the cost of VO_2 , it is crucial to explore new synthesis routes. Due to their unique structure and properties, metal-organic frameworks (MOFs) provide new ideas for synthesizing VO_2 with their unique structure and properties as a new type of compound. The characteristics of MOFs are that they have a large specific surface area, high porosity, diverse structural tunability, and excellent versatility. The MOF derivatives obtained through calcination can maintain the structural advantages of the original MOFs. In addition to controlling the morphology and size of VO₂ using MOF precursors or templates, it is also possible to introduce new properties or functions into VO₂ using MOFs as precursors or templates. By adjusting the structure and composition of MOFs, it is possible, for example, to dope or composite VO₂ to improve its electrochemical efficiency.

6.2 Multi-strategy collaborative optimization

To improve one aspect of performance, a single modification often comes at the expense of another. It is necessary to integrate multiple optimization measures to achieve a highperformance VO₂-based cathode. By carefully regulating the crystal structure, morphology, and size of VO₂, its ion transmission path and charge storage capacity can be significantly improved. Modification methods such as ion intercalation and carbon coating can also be used to enhance the electrochemical activity, structural stability, and cycle stability of VO2. Ion intercalation can increase the interlayer spacing by introducing specific ions into the crystal structure of VO₂, thereby providing a greater channel for the insertion and extraction of zinc ions. This results in a reduction in the resistance to ion transmission and an increase in the rate of charging and discharging and the energy density of the battery. To improve the overall conductivity of the material, we can use carbon coatings to create a layer of carbon on the surface of VO2 particles. By doing so, we are able to extend the life cycle of the battery by preventing the VO₂ structure from being damaged during charging and discharging processes. It is through this technique that AZIBs can be developed with higher storage capacities and longer cycle lives, both of which can contribute greatly to the rapid advancement of AZIB technology in practical applications.

6.3 Improvement of electrolyte adaptation

In AZIBs, the electrolyte plays a crucial role in the performance and stability of the battery. Aqueous electrolyte is another important factor affecting the reversibility of the zinc anode within a specific electrochemical working window. Additionally, aqueous electrolyte controls the dissolution and transfer kinetics of cathode materials. AZIBs' performance can be improved by optimizing the electrolyte. The most commonly studied zinc salts are zinc sulfate (ZnSO₄), zinc chloride (ZnCl₂), and zinc trifluoromethanesulfonate $(Zn(OTF)_2)$. Generally, research focuses more on the type of salt used, but ignores the effect of the amount. From the perspective of energy density, the less electrolyte used, the higher the energy density. As a result of the continuous electrolyte consumption caused by selfcorrosion inside the battery, it is necessary to maintain an appropriate amount of electrolyte to maintain normal operation of the battery. Electrolytes must precisely match cathode and anode requirements. Zinc ions must be transported quickly and smoothly in the electrolyte by choosing an electrolyte with high ionic conductivity and low viscosity. To prevent decomposition or performance degradation during battery charging and

discharging, the solvent and salt components in the electrolyte need to be chemically and thermally stable. A variety of strategies can be adopted to further optimize the electrolyte's performance. Adding the right amount of additives, such as surfactants or corrosion inhibitors, can improve the stability of the electrolyte and its compatibility with electrode materials. The pH value of the electrolyte is also an important optimization method for improving the electrolyte's overall performance. This new electrolyte system can achieve a balance between a wider working window, better compatibility between positive and negative electrodes, and better safety performance by using dual solvents or even ternary solvents. Choosing this option not only improves the overall performance, but also guarantees a more stable battery system operation.

6.4 Advanced characterization methods and theoretical calculations

In the process of studying VO_2 as a cathode material for AZIBs, advanced characterization methods (in situ Raman, synchrotron radiation, scanning tunneling microscopy, etc.) can be used to gain a deeper understanding of its performance and structure. During electrochemical reactions, in situ Raman can monitor the structural changes in materials in real time, which is crucial for understanding phase change and structural evolution. The in situ Raman method reveals the energy storage mechanism of VO₂ materials during the charge and discharge process. Synchrotron radiation light sources can provide deep insights into the microstructural characteristics and evolution mechanisms of VO₂ positive electrode materials. Synchrotron radiation technology can track the dynamic evolution of the material structure and provide direct evidence for understanding VO2's structural changes during charging and discharge. For understanding the electrochemical properties of VO2, scanning tunneling microscopy (STM) can be used to study its atomic arrangement and electronic structure. By utilizing this information, researchers can develop cathode materials with higher energy density and better cycle stability. Theoretical calculations, as well as simulations and predictions, can provide a deeper understanding of the advantages and disadvantages of VO₂ cathode materials. For example, first-principles calculations can predict the electronic structure and energy band structure of VO₂. Besides providing theoretical support for experiments, DFT can also simulate the structural changes of VO2 during charging and discharging.

The exploration of VO_2 as a cathode material for AZIBs presents both challenges and opportunities. Every stage of the process, from synthesis strategies to the enhancement of modification techniques and the adaptation and optimization of electrolytes, is essential and demands continuous innovation and breakthroughs. With deeper research, more efficient and cost-effective methods for synthesizing VO_2 are likely to emerge. Furthermore, applying various modification strategies can improve its electrochemical properties, leading to higher and more stable performance. Additionally, the electrolyte, being a critical component of the battery system, must also be optimized. Enhancing AZIB technology by developing a more adaptable and efficient electrolyte system will be key. VO_2 -based AZIBs hold great potential in energy storage and are poised to make significant contributions to sustainable development.

Data availability

No primary research results, software or code have been included and no new data were generated or analysed as part of this review.

Author contributions

Botao Wan: data curation, writing – original draft. Yajiang Wang: investigation, writing – original draft. Xiudong Chen: conceptualization, project administration, funding acquisition, writing – review & editing. Changchao Zhan: supervision, formal analysis. Huixiong Jiang: conceptualization, supervision. Jin-Hang Liu: investigation, formal analysis. Yun Gao: formal analysis, project administration. Xiaoduo Jiang: resources, formal analysis. Xiaohua Cao: supervision. Hang Zhang: data curation, formal analysis, project administration. Shi-Xue Dou: conceptualization, supervision. Yao Xiao: supervision, visualization, formal analysis, writing – review & editing.

Conflicts of interest

There are no conflicts to declare.

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