Chemical Science

EDGE ARTICLE



Cite this: Chem. Sci., 2022, 13, 11883

All publication charges for this article have been paid for by the Royal Society of Chemistry

Received 17th July 2022 Accepted 23rd September 2022 DOI: 10.1039/d2sc03980j

rsc.li/chemical-science

Introduction

Lithium ion batteries (LIBs) have been widely used in portable electronics and electric vehicles,^{1,2} owing to their high energy density, low self-discharge, and long-term stability.³⁻⁵ However, explosive growth in the demand for LIBs has led to raw materials becoming more and more scarce and expensive. Especially, conventional electrode materials heavily rely on inorganic materials prepared from raw resources.^{6,7} Additionally, these conventional inorganic electrode materials usually present complex preparation processes and high energy consumption.⁸⁻¹⁰ Meanwhile, serious structural collapse and pulverization for these intercalations or alloying type inorganic materials always lead to fast capacity decay during charge/discharge processes.¹¹⁻¹³ Therefore, it is critical to develop low-cost,

Enhanced cycling stability and rate capability of a graphene-supported commercialized Vat Blue 4 anode for advanced Li-ion batteries†

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Commercialized Vat Blue 4 (VB4) has attracted more attention as a promising anode for large-scale applications in Li-ion batteries (LIBs) due to its high electrochemical activity, low price, and large-scale production. However, its moderate solubility results in severe capacity decay and low utilization of active components. Herein, we present a graphene-supported VB4 composite (VB4/rGO) prepared by a facile sonication and hydrothermal process for long cycling stability and high-rate capability. This design can significantly enhance the Li-storage properties, including high capacity (1045 mA h g⁻¹ at 0.1 A g⁻¹), long cycling stability (537 mA h g⁻¹ even over 1000 cycles at 1 A g⁻¹), and rate capability (315 mA h g⁻¹ at 5 A g⁻¹). Strong π - π interaction derived from the aromatic rings within the π -conjugated system (graphene and VB4) and spatial confinement in-between graphene sheets both can suppress the high solubility of VB4 for superior capacity retention. Moreover, conductive graphene and channels in-between nanosheets can simultaneously facilitate the electron and Li⁺ transfer. This work demonstrates a simple and effective method to improve the electrochemical performance of commercialized Vat dyes and provides a low-cost and large-scale strategy to develop their practical application in the energy storage field.

sustainable, and high-performance electrode materials instead of conventional inorganic compounds for future LIBs.¹⁴⁻¹⁶

Compared to conventional inorganic electrode materials, organic compounds not only present high capacity, abundant and eco-friendly resources, and facile synthesis, but also can easily control the electrochemical properties by molecular structure design.9,17-19 Moreover, based on the rearrangement of chemical bonds during the energy storage process, organic electrode materials can avoid severe structural change, which often exists in inorganic electrode materials due to the insertion/extraction of ions.^{20,21} Benefiting from these advantages, numerous organic electrode materials have been applied including conductive polymers,²² in LIBs, carbonyl compounds,^{19,23,24} imine compounds,²⁵ oxynitride compounds, organic sulfur-containing compounds, and radical compounds.26 Among them, Zhang et al.27 reported conjugated carboxylate (Li₄C₈H₂O₆) and non-conjugated carboxylate (Li₄C₄H₂O₆), which shows high discharge capacities of 251.7 and 251.6 mA h g^{-1} at 0.5 A g^{-1} , respectively. Xia *et al.*²⁸ reported a heteroaromatic hexacarboxylate compound (Li₆-HAT) with a high capacity of 1126.3 mA h g^{-1} , as well as superior cycling stability. Additionally, numerous covalent organic frameworks have also been reported as electrode materials for advanced LIBs.^{29,30} However, it is usually complicated and time-



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[†] Electronic supplementary information (ESI) available. See https://doi.org/10.1039/d2sc03980j

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consuming to prepare these products with high and uniform quality on a large scale for practical applications.³¹

By contrast, a commercial carbonyl product, Vat Blue 4 (VB4), can be easily obtained from plants or synthesized artificially. More importantly, rich carbonyl and conjugated structures inside simultaneously ensured high theoretical capacity and high electronic conductivity. These advantages demonstrate that VB4 is a desirable electrode and capable of mass production. Unfortunately, VB4 exhibits high solubility in organic electrolytes (carbonate and ether-based), which easily results in performance degradation.32,33 To address these problems, polymerization, salification, and hybridization with insoluble materials have been adopted and proved effective strategies.^{34,35} Among them, constructing a carbon-supported structure is an effective approach to suppress its severe solubility through the strong π - π interaction derived from the highly π -conjugated system and spatial confinement inbetween the graphene sheets. Meanwhile, high conductivity graphene and channels between the nanosheets can further ensure fast electron/ion transport throughout the hybrid,^{36,37} the stability and kinetics of LIBs, thus promoting simultaneously.

Herein, we developed a graphene-supported VB4 hybrid (VB4/rGO) *via* a facile combination of sonication and hydrothermal processes. According to the strong π - π interaction between the aromatic rings within VB4 and graphene, VB4 is uniformly confined in-between the graphene sheets. Benefiting from the synergistic effects of strong π - π interaction and spatial confinement, the high solubility of VB4 can be effectively suppressed, thus enhancing the stability and utilization of active components. Additionally, conductive graphene and numerous channels in a hybrid can further ensure fast electron/ ion transport. Accordingly, when VB4/rGO was evaluated as an anode for LIBs, the VB4/rGO hybrid exhibited enhanced capacity, long-term cycling stability, and high-rate capability.

Results and discussion

Characterization

In this paper, VB4/rGO was obtained by a facile combination of sonication and hydrothermal processes. VB4 is uniformly distributed between the graphene sheets during the procedure according to the strong π - π interaction derived from their aromatic and conjugated systems (Fig. 1a). The molecule structure of VB4 inside the hybrid is not changed after hydrothermal processes, just as shown in the Fourier transform infrared spectra (FT-IR, Fig. 1b). VB4/rGO and VB4 display the same characteristic peaks for C-N, C=O, and C=C bond stretching in Fig. 1b, revealing the successful compounding between VB4 and rGO. Subsequently, scanning electron microscopy (SEM) and transmission electron microscopy (TEM) are performed to observe the morphology of VB4 with and without graphene. Before the hydration, the pristine VB4 monomer presents a densely stacked block consisting of numerous nanorods (Fig. 1c). After sonication and hydrothermal processes, the uniformly dispersed VB4 nanorods are



Fig. 1 (a) Schematic illustration of the VB4/rGO composite preparation. (b) FT-IR spectra of the VB4 and VB4/rGO composites. (c) SEM images of VB4. (d) SEM, (e) TEM, and (f) elemental mapping images of the VB4/rGO composite.



High-resolution XPS of (a) C 1s, (b) N 1s, and (c) O 1s spectra for VB4/rGO, VB4, and GO.

tightly attached to the graphene nanosheets, just as shown in Fig. 1d and e. Importantly, no evident bulk VB4 is viewed, explaining that VB4 has been fully dispersed and absorbed on the surface of graphene sheets. Further energy dispersive spectroscopy (EDS) mapping of VB4/rGO also confirms that C (green), O (red), and N (blue) elements are evenly distributed on the graphene sheets, just as shown in Fig. 1f. This graphenesupported structure can ensure high utilization of active sites inside the VB4 skeleton and enhanced electron/ion transfer kinetics.38

Afterward, the detailed composition and groups of these products are tested by X-ray photoelectron spectroscopy (XPS). Significantly, the existence of the N 1s signal at 400 eV in the XPS survey mainly contributes to the VB4 molecules, indicating the introduction of VB4 in the composite (Fig. S1[†]). Subsequently, the high-resolution C 1s XPS spectra analyze the evolution of the C-N group and C-O group originating from VB4 and rGO, respectively, as shown in Fig. 2a. The co-existence of C-N and C-O groups in the hybrid undoubtedly confirms the combination of VB4 and graphene. More importantly, a π - π * satellite peak at about 291 eV can be observed in the hybrid, which implies conjugated systems and π - π stacking,^{38,39} ensuring that VB4 strongly interacts with graphene. Moreover, the highresolution N 1s and O 1s XPS spectra also prove the evolution of the C-N and C-O groups among them (Fig. 2b and c), which are well consistent with the C 1s XPS spectra. Additionally, the highresolution C 1s and O 1s XPS spectra both show a decreased C-O group content in the composite after the hydrazine hydrate reduction process (Fig. 2a and c), indicating the reduction of GO.

Subsequently, the composition and interaction between VB4 and rGO were further explored. Fig. 3a exhibits the X-ray diffraction (XRD) patterns of VB4, rGO, and the VB4/rGO hybrid. For the VB4/rGO hybrid, the diffraction peaks located at 6.4° , 11.3° , 13.1° , 22.6° , and 27.2° can be attributed to the characteristic peaks of VB4 and rGO, further revealing the successful compounding between VB4 and rGO without any other peaks for impurities. Note that the above diffraction peaks of the hybrid significantly shift towards a low degree compared to those of VB4, demonstrating that rGO interacted with the VB4 molecule. Meanwhile, the Raman spectra of VB4, rGO, and the VB4/rGO hybrid present broad D and G bands (Fig. 3b). Compared with VB4 and rGO, the hybrid also presents an obvious shift of the D and G bands towards lower wavenumbers, which highly supports the charge transfer from VB4 to graphene, thus revealing the π - π interaction between VB4 and graphene.²⁸ Owing to the strong π - π interaction and spatial confinement in-between graphene, this graphene-supported structure can effectively suppress the solubility of VB4 in the electrolyte, just as shown in Fig. 3c. Significantly, VB4 is dissolved in the electrolyte while VB4/rGO is insoluble even over four months, confirming the stability of VB4/rGO in the electrolyte. Subsequently, to analyze the structure-property relationships of VB4/rGO in-depth, density functional theory (DFT) simulation is carried out (Fig. S2⁺). Fig. 3d presents a high VB4 adsorption energy of -15.34 eV for the graphene-supported structure, proving the strong interaction against the high solubility of VB4 as well. Moreover, the charge density difference is introduced to explore the charge distribution of VB4 adsorption models. The charge accumulation region around the interfaces is more pronounced (Fig. 3d and e), indicating strong VB4 binding with graphene. Meanwhile, Fig. 3f represents different models' density of states (DOS). The DOS of VB4/rGO around the Fermi level is greatly enhanced compared to that of VB4, revealing a high electronic conductivity.29 Thus, benefiting from the high VB4 adsorption ability and electronic conductivity, the graphene-supported VB4 hybrid could achieve a higher stability and faster reaction kinetics compared to pristine VB4.

Fig. 3g shows the nitrogen adsorption-desorption isotherms of VB4/rGO, which reveal a Brunauer-Emmett-Teller (BET)



Fig. 3 (a) XRD patterns and (b) Raman spectra of rGO, VB4, and VB4/rGO. (c) Photograph of VB4 and VB4/rGO soaked in the electrolyte. (d and e) Adsorption energy and charge density differences of VB4 molecules in-between graphene sheets. (f) Density of states of VB4 and VB4/rGO models. (g) N₂ sorption isotherms and pore size distribution curve of VB4/rGO.

surface area of 184.85 m² g⁻¹. Moreover, the features of type IV and H3 hysteresis loops illustrate the existence of mesoporous structures. Afterward, an average pore size of 3-5 nm is confirmed according to the pore-size distribution of VB4/rGO (the inset of Fig. 3g). By contrast, owing to the stacked block morphology (Fig. 1c), the surface area of VB4 is calculated to be

only 20.05 $m^2 g^{-1}$ (Fig. S3[†]). Therefore, the construction of a graphene-supported structure can form fast ion-diffusion channels, and provides large contact areas between the electrolyte and active components, finally enabling higher reaction kinetics and more active sites compared to pristine VB4.



Fig. 4 (a) The CV curves of VB4/rGO anodes at a scan rate of 0.1 mV s^{-1} . (b) The cycling performance of VB4 and VB4/rGO anodes at 0.1 A g^{-1} . (c) Charge-discharge voltage profiles of VB4/rGO at various current densities. (d) Rate performances of VB4 and VB4/rGO anodes. (e) Cycling performance of VB4 and VB4/rGO anodes at 1 A g^{-1} . (f) Ex situ FT-IR spectra of the VB4/rGO anode in different cycles.

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Electrochemical performance

Benefiting from these advantages, the resultant hybrid could possess improved electrochemical performances. Subsequently, the CV curves of the VB4/rGO anode for LIBs are first collected. As shown in Fig. 4a, two redox peaks are observed in the first cycle. Among them, an irreversible peak appears at 1.4–1.6 V due to solid electrolyte interface (SEI) formation and disappears in the subsequent cycles.^{30,40,41} Note that the CV curves during the cycles all present obvious couples of redox peaks at 0.75/ 1.2 V, indicating good reversibility and great stability during the lithiation/delithiation processes.

Fig. 4b shows the cycling performance of the VB4 and VB4/ rGO anodes at 0.1 A g⁻¹. The VB4/rGO anode presents a high reversible discharge capacity of 1045 mA h g⁻¹ even after 100 cycles, which is far superior to that of VB4 (626 mA h g^{-1}) and other reported electrode materials.^{28,42-44} Subsequently, the rate performances of the VB4 and VB4/rGO anodes at different current densities ranging from 0.1 A g^{-1} to 5 A g^{-1} are explored (Fig. 4c and d). With the increase of current densities from 0.1 A g^{-1} to 5.0 A g^{-1} , VB4/rGO exhibits a high capacity of 1097, 906, 714, 565, 444, and 315 mA h g^{-1} , respectively. Particularly, the capacity of VB4/rGO can still recover to 1000 mA h g^{-1} , when the current density is back to 0.1 Ag^{-1} . By contrast, the capacity of pristine VB4 drops quickly from 617 to 172 mA h g⁻¹ owing to its high solubility. These enhanced properties of VB4/rGO can be attributed to the introduction of a graphene-supported structure and multi-electroactive centers inside the VB4 skeleton.

Moreover, the long-term cycling stability of these products at a high current density of 1.0 A g^{-1} is displayed in Fig. 4e. VB4/rGO exhibits an excellent reversible capacity of 537 mA h g⁻¹ even over 1000 cycles. Meanwhile, an average coulombic efficiency of 100%

can be obtained throughout the cycles. In contrast, the discharge capacity of VB4 is only 265 mA h g^{-1} after 400 cycles, much lower than that of VB4/rGO, indicating that the introduction of graphene is beneficial for Li-storage properties. To explore the stability of VB4/rGO, electrochemical impedance spectroscopy (EIS) plots upon cycling are collected, as shown in Fig. S4.[†] With increasing cycle numbers, the R_{ct} corresponding to charge transfer impedance significantly decreases, indicating a fast charge transfer process throughout the cycling. Additionally, the slope of the EIS plots within the low frequency region is always related to the Li ion diffusion. In the initial cycles, the decreased slope usually corresponds to sluggish ion diffusion, which is related to the initial activation process. Afterward, the slopes remain basically unchanged, indicating stable Li ion diffusion, which is well consistent with the evolution of cycling performance. Impressively, compared to the reported electrodes (Table S1[†]), VB4/rGO can still exhibit competitive cycling stability and rate performance. Afterward, to further investigate this cycling stability, the ex situ FT-IR spectra of VB4/rGO in different cycles are collected in Fig. 4f. Throughout the long-term cycling, these characteristic peaks attributed to VB4 remain consistent, ensuring excellent reversibility and stability. These remarkably enhanced performances of VB4/rGO are mainly attributed to its unique graphene-supported structure, which can significantly suppress the high solubility of VB4 according to the strong interaction and enhance its electronic conductivity. Additionally, the large specific surface area and the porous structure can increase the contact area with the electrolyte and expose numerous electroactive sites of the VB4 molecule.

Afterward, to explore the reaction kinetics of VB4/rGO, EISis first performed to analyze the electronic/ionic conductivity. As



Fig. 5 (a) EIS plots and (b) GITT curves of VB4 and VB4/rGO. (c and d) Li⁺ diffusion coefficients during charge/discharge processes for VB4 and VB4/rGO. (e) CV curves of VB4/rGO at different scan rates. (f) log *i* vs. log v profiles of VB4/rGO.

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shown in Fig. 5a, EIS curves are composed of a semicircle and sloping line. The smaller diameter of the semicircle assigned to VB4/rGO indicates a lower charge transfer resistance ($R_{\rm ct}$) than that of VB4, demonstrating that the introduction of graphene is beneficial for the improved conductivity of VB4. Additionally, the plot of VB4/rGO in the low-frequency region is closer to the real impedance axis, confirming the better Li⁺ diffusion compared to VB4. Afterward, the detailed Li⁺ diffusion coefficients are estimated according to the galvanostatic intermittent titration technique (GITT, Fig. 5b) and the following equation,⁴⁵

$$D_{\mathrm{Li}^+} = \left(\frac{4L^2}{\pi\tau}\right) \times \left(\frac{\Delta E_{\mathrm{s}}}{\Delta E_{\mathrm{\tau}}}\right)^2$$
 (1)

where *L* represents the transfer distance of Li⁺, τ is the relaxation time, and ΔE_s and ΔE_{τ} stand for the difference of voltages after adjacent relaxations and the voltage change during the current pulse, respectively (Fig. S5 and S6†). Fig. 5c and d show the evolutions of D_{Li}^+ for VB4/rGO and VB4 during the charge and discharge processes. They both present the same shift trend of the diffusion coefficient (10⁻¹² to 10⁻⁹ cm² s⁻¹) in the range from 3.0 V to 0.01 V. Noting that, during the charge/

discharge processes, the D_{Li}^+ of VB/rGO is higher than that of VB4. Moreover, the diffusion coefficient of VB4/rGO is within or better than that of other reported organic materials, such as HAB-COF ($10^{-12.5}$ to $10^{-9.5}$ cm² s⁻¹),⁴⁶ rCTF (7.00×10^{-11} cm² s⁻¹),⁴⁷ and E-FCTF (2.43×10^{-11} to 5.09×10^{-10} cm² s⁻¹),⁴⁸ ensuring better ion diffusion kinetics.

Afterward, the electrochemical mechanism of VB4/rGO is tested by coupling the CV curves at different scan rates (Fig. 5e) with the following equations,³⁸

$$i = av^b \tag{2}$$

$$\log i(V) = b \log v + \log a \tag{3}$$

where the current (*i*) complies with the power-law relationship with the scan rate (v), and *a* and *b* are adjustable parameters. *b* represents the domination of a totally diffusion-controlled mechanism (b = 0.5) or capacitive-controlled process (b =1).³⁹ Then, according to the relationship between $\log(i)$ and $\log(v)$ (Fig. 5f), the values of *b* are determined from the slopes of the redox peaks. The calculated 0.9 and 0.8 indicate that the Li-



Fig. 6 (a and b) The *ex situ* FT-IR spectra of the VB4/rGO anode under different charge/discharge states. (c) DFT calculated electronegativity of VB4. (d) Illustration of the structural advantages for the VB4/rGO hybrid.

storage of VB4/RGO is mainly controlled by a capacitive process along with fast reaction kinetics.

Subsequently, ex situ FT-IR corresponding to the different potential states of discharge/charge profiles is studied to explore the Li⁺ storage mechanism of VB4/rGO (Fig. 6a and b). As shown in Fig. 6b, with the battery discharge from 3.0 to 0.01 V, the characteristic peaks of C=C (1580 and 1450 cm⁻¹) and C=O (1640 cm^{-1}) gradually weaken due to the Li⁺ reaction with C=C and C=O to form C-C-Li and C-O-Li, respectively.49 Then, during the charging process, the tendency of enhanced peaks of C=C and C=O can be observed, indicating the removal of Li^+ from C-C-Li and C-O-Li. Such reversible evolutions can ensure the electroactive centers (C=C and C=O) and Li⁺ storage mechanism in the VB4 molecule. Besides, further ex situ XPS was also performed to analyze the Li-storage mechanism of VB4/rGO, just as shown in Fig. S7.† During the discharge process, the peaks of C=O and C=C gradually weaken in the high-resolution C 1s and O 1s spectra. By contrary, the peaks of C-C and C-O gradually increase inside, indicating the reaction of Li⁺ with C=C and C= O. During the subsequent charging process, the peaks of C=C and C=O bonds gradually recover, demonstrating the highly reversible process of Li-storage. Meanwhile, by DFT simulation, the results of electron density distribution around the C=C and C=O groups exhibit obvious negative values (Fig. 6c), further proving the highly reactive centers.⁵⁰ In summary, the electrochemical merits of VB4/rGO are mainly due to its graphenesupported structure and molecular composition, as illustrated in Fig. 6d. (1) Strong π - π interaction within the conjugated systems and spatial confinement in-between graphene sheets both can suppress the high solubility of VB4 for high cycling stability. (2) Highly conductive graphene and the porous structure can facilitate electronic and Li⁺ transfer. (3) Multi-electroactive centers can coordinate Li⁺ for high theoretical capacity.

Conclusions

This paper reports a simple and universal strategy to prepare a graphene-supported VB4 anode for LIBs by assembling VB4 molecules and graphene. For this design, multi-electroactive centers within the VB4 skeleton can result in high Li-ion storage capacity. The introduction of conductive graphene can not only suppress the high solubility of VB4 according to the strong π - π interaction and spatial confinement in-between graphene sheets, but also exposes numerous electroactive sites and provide fast electron/ion transfer highways. Therefore, VB4/rGO presents outstanding electrochemical performance, such as a high reversible capacity of 1045 mA h g^{-1} at 0.1 A g^{-1} , good cycling stability (537 mA h g^{-1} after 1000 cycles at 1 A g^{-1}), and good rate performance (315 mA h g^{-1} even at 5 A g^{-1}). More importantly, this work provides an effective method to improve the Li-ion storage performance of commercialized dyes and a low-cost, large-scale strategy for practical applications.

Data availability

The data that support the findings of this study are openly available on line.

Author contributions

H. W. K. and C. F. Z. designed and guided this experiment. Q. W. M., R.W., X.R.W. and H. W. K. carried out the experiment and materials characterization. S. S. C. and L.H.Z. contributed to the data analysis and co-wrote the paper. H. W. K., S. S. C., and C. F. Z. proposed and supervised the project.

Conflicts of interest

There are no conflicts to declare.

Acknowledgements

We thank the financial support from the National Natural Science Foundation of China (51872071 and 52172173), Anhui Provincial Natural Science Foundation for Distinguished Young Scholar (2108085J25), Anhui Province Key Laboratory of Environment-Friendly Polymer Materials, Open Project of Anhui Provincial Key Laboratory for Degradation and Monitoring of Pollution of the Environment (FSKFKT001D), Natural Science Research Projects of Universities in Anhui Province (KJ2020A0021 and KJ2020A0547), and Talent program of Fuyang Normal University (2020KYQD0015). This work was also supported by the Open Fund of Guangdong Provincial Key Laboratory of Advance Energy Storage Materials (AESM202106) and Science-Technology Development Program of Henan Province (202102210030). The authors also acknowledge the High-performance Computing Platform of Anhui University for providing computing resources.

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