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Copper and conjugated carbonyls of metal–organic polymers as dual redox centers for Na storage†

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Metal–organic polymers (MOPs) are fascinating electrode materials for high-performance sodium-ion batteries due to their multiple redox centers and low cost. Herein, a flower-like π – d conjugated MOP (Cu–TABQ) was synthesized using tetramino-benzoquinone (TABQ) as an organic ligand and Cu^{2+} as a transition metal node under the slow release of Cu^{2+} from $[\text{Cu}(\text{NH}_3)_4]^{2+}$ and subsequent dehydrogenation. It possesses dual redox centers of $\text{Cu}^{2+}/\text{Cu}^+$ and $\text{C}=\text{O}/\text{C}-\text{O}$ to render a three-electron transfer reaction for each coordination unit with a high reversible capacity of $322.9 \text{ mA h g}^{-1}$ at 50 mA g^{-1} in the voltage range of 1.0 to 3.0 V. The flower-like structure enhances fast Na^+ diffusion and highly reversible organic/inorganic redox centers. This results in excellent cycling performance with almost no degradation within 700 cycles and great rate performance with $198.8 \text{ mA h g}^{-1}$ at 4000 mA g^{-1} . The investigation of the Na-storage mechanism and attractive performance will shed light on the insightful design of MOP cathode materials for further batteries.

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Introduction

Sodium-ion batteries (SIBs) have attracted great attention for large-scale energy storage due to the high abundance and uniform distribution of Na resources.^{1,2} A wide range of inorganic materials, such as $\text{Na}_3\text{V}_2(\text{PO}_4)_3$, $\text{Na}_{0.67}\text{MnO}_2$, and $\text{Na}_2\text{-FeFe}(\text{CN})_6$, have been attempted as cathodes for SIBs, to deliver capacities below 200 mA h g^{-1} due to the limited redox of $\text{V}^{4+/3+}$, $\text{Mn}^{4+/3+}$, and $\text{Fe}^{3+/2+}$.^{3–6} Polymer-based organic electrode materials have been actively studied as host materials for Na^+ storage owing to their environmental friendliness and good flexible stability with devisable organic monomers, which could introduce multiple organic/inorganic redox centers to increase theoretical capacity.^{7–10} However, the electrochemical performances of these polymers are governed by their electrically insulating nature (except for conductive polymers) and disordered molecular chain entanglement, leading to the limited

utilization of redox-active sites with low capacity and inferior rate performance.^{11–13} It is rational to design organic cathode materials with more redox sites and great electrical conductivity for Na^+ storage and robust structures for Na^+ (de)intercalation for high energy density and reversibility.

Redox-active π – d coordination metal–organic polymers (MOPs) exist as the framework structure or linear structure by adjusting the type of inorganic metal nodes and organic ligands, which is favorable for eliminating the dissolution of the hybrid materials in organic electrolytes and avoiding the organic chain agglomeration.^{14–17} Moreover, the abundantly delocalized electrons and strong metal–ligand interaction bonds in MOPs guarantee high electrical conductivity and good structural stability, promoting the electron transfer to activate redox-active sites with higher capacity.^{18,19} It is noteworthy that the selection of organic ligands and transition metal centers for MOPs is important to affect the discharge capacity, working potential, and cycling performance. Several typical organic ligands, such as hexaaminobenzene (HAB), 1,2,4,5-tetraaminobenzene (BTA), and 2,3,6,7,10,11-hexaaminotriphenylene (HTTP), have been widely used as the anode, while their electrochemical performance is still limited by the single active site and low discharge capacity.^{20–23} Recently, many studies on conjugated carbonyl-derived linkers have indicated to construct a high-capacity MOP-type cathode material because of its low molecular weight, high potential redox sites (carbonyl group, $\text{C}=\text{O}$), and *ortho*-diamine groups ($-\text{NH}_2$, coordination form of robust metal–nitrogen bonds), like tetramino-benzoquinone (TABQ).^{16,24–32} This is favorable for increasing the discharge

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voltage and enhancing the structural stability upon Na^+ (de) intercalation. For the transition metal centers, the copper cation (Cu^{2+}) shows stronger competitiveness in terms of cost, toxicity, chemical stability, high redox voltages, and extra capacity compared with other metal ions (e.g., Ni^{2+} , Co^{2+} , Sn^{2+} , etc.).^{21,33–37} Therefore, it is desirable to combine TABQ and Cu^{2+} into a MOP skeleton to achieve high energy-density organic cathodes for SIBs and explore the Na^+/e^- transfer mechanism and structural evolution during the charge/discharge process.

Herein, we synthesized linear Cu-based MOPs using copper (Cu, II) ions as transition metal centers and tetramino-benzoquinone molecules (TABQ, $\text{C}_6\text{H}_8\text{O}_2\text{N}_4$) as organic ligands in a mild process. It is formed with a coplanar π -d delocalization skeleton in each unit of Cu-TABQ to enhance Na^+ diffusion. Active metal ions (Cu^{2+}) and organic ligands (benzoquinone framework) act as the dual redox centers for Na storage, leading to three Na^+ transfers per coordination unit, as revealed in *in situ* FTIR and *ex situ* XPS. This provides a high reversible capacity of $322.9 \text{ mA h g}^{-1}$ at 50 mA g^{-1} within voltages of 1.0–3.0 V for Cu-TABQ. The results show that the Cu-TABQ is an ideal candidate for a high-performance cathode to construct Na-organic batteries and also provides a new direction for the future development of metal-ion batteries.

Results and discussion

The Cu-TABQ MOPs were synthesized *via* coordination reactions between Cu^{2+} and TABQ in dimethylsulfoxide (DMSO) with ammonia under ambient air, as shown in Fig. 1a. The coordination chains are gradually developed and eventually form the coplanar structure and polymeric chain by four N

atoms and one Cu^{2+} coordination, which is accompanied by the oxidative dehydrogenation of TABQ ligands by O_2 and the release of Cu^{2+} from $[\text{Cu}(\text{NH}_3)_4]^{2+}$ in Fig. S1a and b.† Cu-TABQ exhibits π -d conjugated coordination due to the hybridization between the π orbitals of TABQ and the d orbitals of Cu^{2+} , which will effectively enhance the structural stability and electronic conductivity of the coordination compound.³⁸ Compared with TABQ, Cu-TABQ presents a lower LUMO energy level and narrow HOMO–LUMO energy gap (ΔE_g) in Fig. S1c,† indicating enhanced electronic delocalization and increased electron accepting ability. Besides, the Cu-TABQ powder shows an obvious metallic sheen in the optical photo (Fig. S2†). The molecular structure of the designed Cu-TABQ was investigated by various characterization methods including Fourier transform infrared (FTIR) spectroscopy, Raman spectroscopy, and solid-state ^1H nuclear magnetic resonance (^1H NMR) spectroscopy. As shown in Fig. 1b, the characteristic triple vibration signal of $-\text{NH}_2$ in the region of $3000\text{--}3500 \text{ cm}^{-1}$ in the FTIR spectrum of TABQ disappears after coordination, while a new peak ($-\text{NH}-$) appears at around 3276 cm^{-1} for Cu-TABQ due to the reaction of oxidative dehydrogenation. The coexistence of benzoid amine ($\text{C}-\text{N}$) and quinoid imine ($\text{C}=\text{N}$) peaks at 1264 and 1450 cm^{-1} in the FTIR spectra of Cu-TABQ indicates the deprotonation of TABQ to form a new coordination bond with Cu^{2+} and the possession of unpaired electrons around N atoms. This can be confirmed by the red-shifted peak at 1664 cm^{-1} for the carbonyl bond ($\text{C}=\text{O}$) in the organic ligand to 1579 cm^{-1} for TABQ with coordination and attraction between chains.

The chemical state of N and O in the Cu-TABQ structure is analyzed by X-ray photoelectron spectroscopy (XPS) in Fig. S3 and S4.† The high-resolution N1s and O1s spectra confirmed

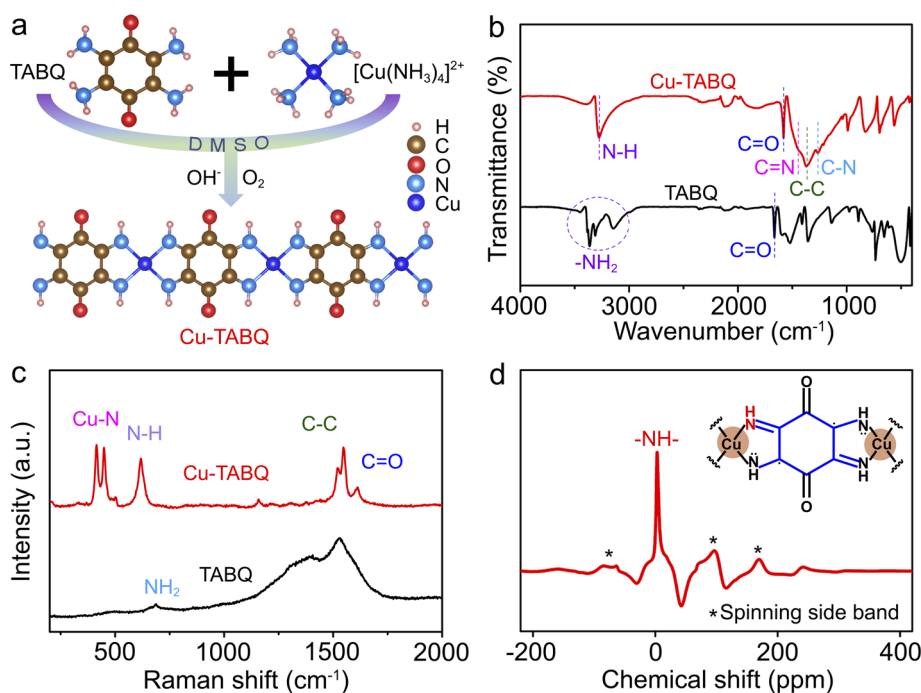


Fig. 1 (a) Synthetic scheme of Cu-TABQ; (b) FTIR and (c) Raman spectra of TABQ and Cu-TABQ; (d) solid-state ^1H NMR spectrum of Cu-TABQ.

the coexistence of C–N and C=N bonds and the presence of C=O in Cu-TABQ.^{30,39} The chemical moieties of coordination between Cu²⁺ and the TABQ ligand can be probed by the emergence of stretching vibrations of Cu–N (414 and 449 cm^{−1}) in the Raman spectrum (Fig. 1c).^{30,40} The structure information of Cu-TABQ was further verified using the solid-state ¹H NMR spectrum. In Fig. 1d, an obvious ¹H signal at 3.1 ppm is assigned to hydrogen in the –NH– bond, further confirming the successful synthesis of MOPs with TABQ and Cu²⁺. The satellite peaks in the Cu 2p spectrum in Fig. S5† reveal the divalent state of Cu. The addition peak at slightly lower binding energy in Cu-TABQ compared to the main Cu 2p_{3/2} peaks indicates the presence of Cu(I).²⁰ Furthermore, a strong EPR signal with a g-factor of 2.0002 was observed in the sample (Fig. S6†), indicative of the presence of ligand radicals which is consistent with the previously reported MOPs.^{17,20,32} These suggest that Cu²⁺ is bonded to TABQ *via* the coordination bond of Cu–N for Cu-TABQ.

The crystal structure of Cu-TABQ was studied by powder X-ray diffraction (PXRD) and simulated by Rietveld refinement in Fig. 2a. The diffraction peaks located at 22.96°, 28.12°, 35.74°, and 40.20° are assigned to the (210), (111), (401) and (332) facets, respectively. The refined unit cell parameters for Cu-TABQ are shown in Table S1,† specifically, *a* = 6.98253 Å, *b* = 6.98253 Å, *c* = 3.22703 Å, $\alpha = \gamma = 92.2166^\circ$, $\beta = 67.5636^\circ$, and *V* = 145.2689 Å³. Fig. 2b shows the planar and layered packing motif of Cu-TABQ, in which there exists a large planar structure through Cu–N square-planar coordination and hydrogen bonds

between chains along the *ab* plane and the stack cambiums into layered structures along the *c*-axis. Density functional theory (DFT) calculation was used to investigate the charge configuration and coordination bonds between Cu²⁺ and TABQ in Cu-TABQ. There is significant overlapping between the d orbitals of Cu atoms and the p orbitals of N atoms in the projected density of states (PDOS) of Cu-TABQ (Fig. 2c), suggesting their strong orbital hybridization. The π –d conjugation of Cu²⁺ and N atoms with square planar coordination manner guarantees the rigid and coplanar nature of the Cu-TABQ chains, which favors the electronic conductivity of Cu-TABQ and hence enhances the redox activity.

The morphology of Cu-TABQ can be controlled by different reaction temperatures. Previous reports have demonstrated that oxygen could increase the possibility of coordination between Cu²⁺ and TABQ molecules, and finally promote the formation of the cross-linked coordination polymer.²¹ As shown in Fig. S7–S9,† the coordination temperatures facily affect the crystallinity and morphology of Cu-TABQ. The integrated SEM and TEM images of Cu-TABQ present 0-dimensional nanoparticles and one-dimensional short rod structures at 30 °C and 60 °C respectively, with a low reaction rate. Whereas the flower-like structure of two-dimensional nanosheets formed at 90 °C for Cu-TABQ because of the fast reaction rate and self-assembly characteristics (Fig. 2d, e, and S10†).⁴¹ The TEM image in Fig. 2f shows that Cu-TABQ has higher crystallization at 90 °C with a flower-like shape, which is consistent with XRD results in Fig. S7.† Moreover, the high-resolution TEM (HRTEM) image of

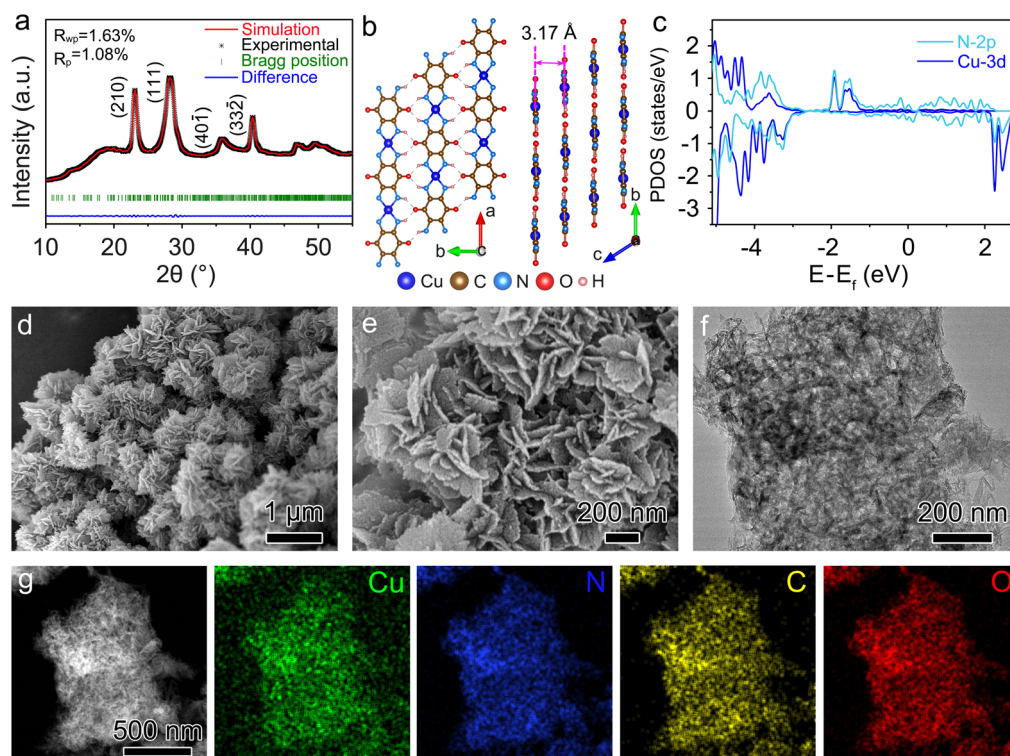


Fig. 2 (a) PXRD patterns and Rietveld refinement; (b) diagram of the planar and layered packing motif; (c) PDOS; (d) and (e) SEM images; (f) TEM image; (g) EDS mappings of Cu-TABQ.



Cu-TABQ reveals periodic lattice fringes with an interplanar spacing of 0.32 nm (Fig. S11[†]), which is assigned to the (001) facet and consistent with the XRD pattern in Fig. 2a. The TEM-EDS-mapping images of Cu-TABQ show the uniform distribution of Cu, C, N, and O elements in Fig. 2g, which is consistent with the elemental analysis results (Table S2[†]). The Brunauer–Emmett–Teller (BET) curve of Cu-TABQ is obtained from the N₂ adsorption isotherm and shown in Fig. S12[†], which displays a type II isothermal curve with a large specific surface area of 118.49 m² g⁻¹. This promises more Na⁺ storage sites and favors the penetration of electrolytes into cathode materials. The thermal stability of Cu-TABQ was investigated using a thermogravimetric analyzer (TGA) in Fig. S13[†]. There is ~3% weight loss at about 180 °C due to the residual DMSO solvent evaporation. Cu-TABQ shows better thermal stability than that of TABQ, which is attributed to the π -d conjugation of Cu²⁺ and TABQ after coordination. Diffuse reflectance ultraviolet-visible (UV-vis) spectroscopy was used to further explore the electron delocalization and metal-ligand conjugation of Cu-TABQ. As shown in Fig. S14[†], Cu-TABQ shows a strong absorption around 439 nm, which could be attributed to the ligand-centered (LC) π - π^* transition.⁴² Its maximum absorption is red-shifted with extended absorption compared with the pristine TABQ revealing electron delocalization between Cu²⁺ and TABQ and the metal-ligand d- π conjugation.^{32,42,43} Further, the electronic conductivity of Cu-TABQ was obtained based on *I*-*V* curves in Fig. S15[†] to show a high value of 1.02×10^{-3} S m⁻¹ ensuring good rate capability in batteries.

The Na-storage performances of Cu-TABQ were tested by assembling CR2032 coin cells using Na metal anodes. Cyclic voltammetry (CV) was performed in the voltage range from 1.0 to 3.0 V at 0.1 mV s⁻¹ in Fig. 3a. The irreversible phenomenon in the

first cycle is mainly caused by the formation of the SEI film.¹⁹ One broad reduction peak located at around 1.55 V in the first cathodic scan originated from the reduction of both Cu²⁺ and the organic ligand. There are two broad oxidation peaks at 1.7 and 2.4 V attributed to oxidation of C=O bonds and Cu⁺ in the initial cycle. In the subsequent cycles, two pairs of overlapped redox peaks located at 1.42/1.68 and 1.73/2.31 V are observed, which can be assigned to the redox reaction of the C=O/C-O bonds and Cu²⁺/Cu⁺, respectively. Fig. 3b shows sloping charge/discharge curves of Cu-TABQ with no clear plateaus at 50 mA g⁻¹ between 1.0 and 3.0 V in selected cycles, which is consistent with the CV curves. It delivers an initial discharge capacity of 426.8 mA h g⁻¹ with a coulombic efficiency of 75.6% in the first cycle. The first 50 cycles show similar overlapping charge/discharge curves, indicating the stable structure of Cu-TABQ during cycling.

The rate performance of the Cu-TABQ electrode is tested at current densities from 50 to 4000 mA g⁻¹ (Fig. 3c and S16[†]). It shows the specific capacity of 350.8 mA h g⁻¹ at 50 mA g⁻¹, revealing a three-electron redox process for every coordination unit. Upon cycling at 4000 mA g⁻¹, Cu-TABQ exhibits a capacity of 198.8 mA h g⁻¹, corresponding to 56.7% of its capacity at 50 mA g⁻¹. This excellent rate performance can be attributed to the unique flower-like structure ensuring fast electrolyte infiltration and the inherent π -d conjugated structure providing high conductivity, and thus fast reaction kinetics of Cu-TABQ. The cycling performance was evaluated at 50 mA g⁻¹ to achieve a high capacity of 318.8 mA h g⁻¹ after 50 cycles with a capacity retention of 98.7% in Fig. 3d. After cycling, there is no obvious color change from the disassembled cells (Fig. S17[†]), which indicates insolubility of the discharged/charged products in the electrolyte. Remarkably, it delivers a capacity of 238.3 mA h g⁻¹ without obvious degradation after 700 cycles when tested at 2000 mA g⁻¹ in Fig. 3e, indicating a high rate stability. The Cu-TABQ electrode shows excellent electrochemical performance in terms of charge/discharge capacity, capacity retention, and rate capability, as compared to the reported metal-organic polymer cathode electrodes (Table S3[†]), suggesting its promising application in SIBs.

To further understand the chemical states and Na⁺ storage mechanism of Cu-TABQ, *in situ* FTIR, *ex situ* XPS, *ex situ* Raman, and computational simulations were carried out. *In situ* FTIR was performed in the voltage range from 1.0 to 3.0 V in the first cycle at 50 mA g⁻¹, as shown in Fig. 4a. During the whole oxidation/reduction process, Cu-TABQ shows a sloping charge/discharge curve, corresponding to a fast reaction kinetics of the Na⁺ (de)intercalation process due to the dual inorganic (Cu²⁺) and organic (C=O) active sites. In the initial discharge process, the signal peak of C=O (1577 cm⁻¹) gradually decreases and eventually disappears, while a new stretching vibration peak is formed around 1199.3 cm⁻¹ that is assigned to C-O bonds. After being fully charged, the intensity of the stretching vibration peak of C=O gradually increases and resumes its original positions without any new phases, suggesting the highly reversible redox reactions of C=O bonds. The peaks of -N-H at 3276 cm⁻¹ and the C-N bond at 1081 cm⁻¹ exhibit similar behaviour to that of C=O bonds, because the insertion of Na⁺ changes the coordination environment of N and Cu atoms and

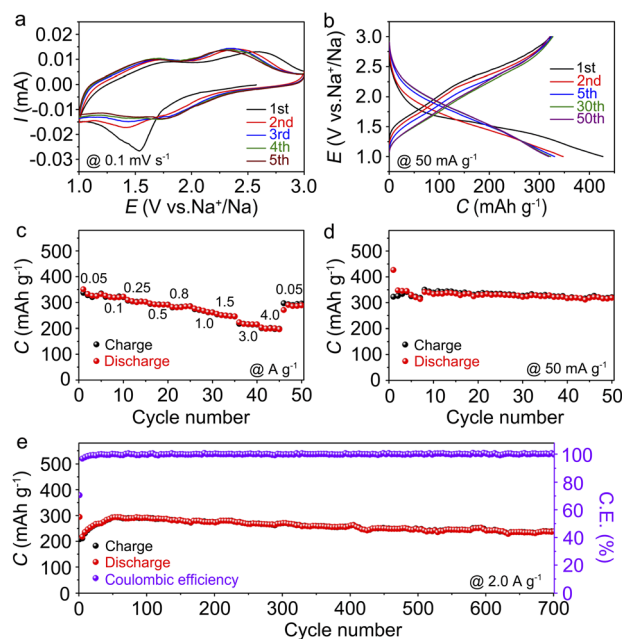


Fig. 3 (a) CV curves at 0.1 mV s⁻¹; (b) discharge/charge curves of initial 50 cycles at 50 mA g⁻¹; (c) rate capability from 0.05 to 4.0 A g⁻¹; (d and e) cycling performance of Cu-TABQ at 50 and 2.0 A g⁻¹.

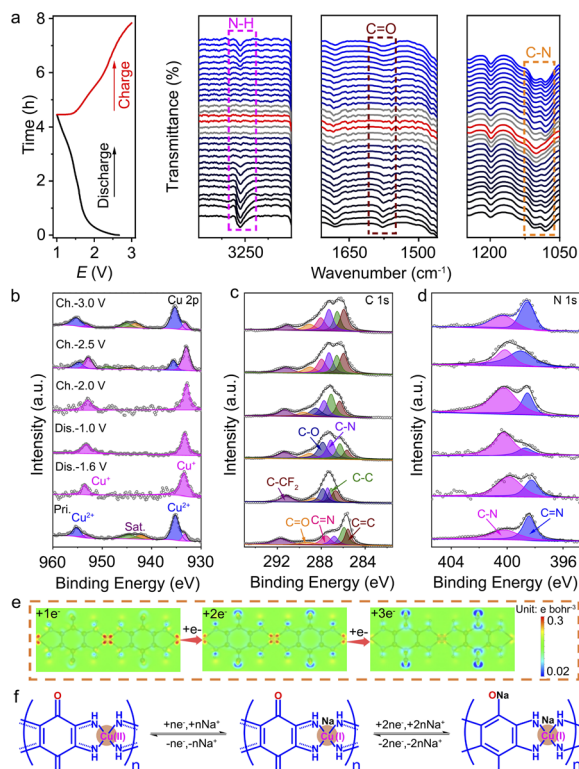


Fig. 4 (a) Discharge/charge profile and *in situ* FTIR spectra of Cu-TABQ; (b–d) *ex situ* Cu 2p, C 1s, and N 1s XPS spectra of Cu-TABQ; (e) extra charge distribution upon reduction of Cu-TABQ after injection of one, two, and three electrons; and (f) schematic diagram of the Na-storage mechanism.

weakens the N–H...Cu(Na⁺)/N–H...N(Na⁺) hydrogen bonding interaction. During the charging process, the vibrational peak of the amine group appeared again with the extraction of Na⁺. At the same time, the disappeared peak at 1081 cm^{−1} and the redshift C–N stretching band after discharging were observed in FTIR spectra, which is related to the π -d conjugation electron rearrangement of the C–N/C=N bond after discharge.

Ex situ XPS of Cu-TABQ was conducted in Fig. 4b–d, S18, and S19[†] to investigate the redox reaction and chemical state of the organic ligand. In its original state, a small amount of Cu⁺ ions at 933.4 eV was detected due to the partially reduced Cu²⁺ in Cu-TABQ during electrode preparation. When the electrode is discharged to 1.6 V, strong satellite peaks of the Cu 2p spectrum in the pristine electrode disappear, and the peak values of Cu 2p_{3/2} and Cu 2p_{1/2} shift to the smaller binding energies of 933.4 and 953.4 eV, indicating the reduction of Cu²⁺ to Cu⁺ in Cu-TABQ during the Na⁺ intercalation process. The Cu 2p spectrum is still dominated by Cu⁺ further discharged to 1.0 V, which is similar to the redox reaction of Cu²⁺/Cu⁺ in other conjugated coordination compounds.^{31,37} After being fully charged to 3.0 V, the Cu-TABQ can be returned to the initial state. Besides, the C 1s spectra are presented in Fig. 4c, and the peak of the C=O bond disappears while the peak of the C–O bond becomes stronger during discharge from 1.6 to 1.0 V, which confirms that C=O groups were involved in the redox process. This suggests that the C=O bond site acts as one redox centre for Na⁺ storage.

In contrast, the C=N bond was strengthened and the C–N bond was weakened after discharge, which is caused by the electron redistribution of C–N/C=N bond upon Na⁺ intercalation.^{14,44} An Auger electron peak of Na at 536.3 eV can be observed in the O 1s spectrum in Fig. S19[†] after discharge, corresponding to the sodiation reaction in Cu-TABQ. The N 1s spectrum of Cu-TABQ is shown in Fig. 4c. At the end of discharge, the peak value of the C–N single bond at 400.2 eV increases, while the peak value of the C=N double bond at 398.4 eV decreases, which is mainly related to the electron changes in the whole molecular structure, caused by Na⁺ intercalation in Cu-TABQ.

Fig. S20[†] shows the Raman spectra of Cu-TABQ in the charging and discharging process. The tensile vibration of the quinone ring remains relatively stable, while the Cu–N bond is weakened and disappears, as in the case of the C=O double bond. This indicates that Cu is involved in the redox reaction as the active metal site, and in agreement with the *ex situ* XPS in Fig. 4b. The *ex situ* XRD patterns of the Cu-TABQ electrode at different charge/discharge states were recorded to study the structural evolution. As shown in Fig. S21,[†] the diffraction peaks located at 28.1° (111) shift slightly leftward to 26.8° when discharged to 1.0 V, indicating the increased interlayer (3.3 Å) caused by the Na⁺ inserted into the space of Cu-TABQ. When it was recharged to 3.0 V, the lattice parameters returned to the initial states, indicating a good reversibility of Cu-TABQ. The molecular electrostatic potential (MESP) of electron-injected Cu-TABQ was further calculated in Fig. 4e to reveal the redox reaction. When one extra electron is injected into the Cu-TABQ unit, the extra electrons tend to locate around N atoms, suggesting the priority of Cu²⁺ reduction. After another two extra electrons are injected into the Cu-TABQ unit, the extra electrons tend to locate around C and O atoms, indicating the subsequent reduction of C=O. Fig. 4f schematically illustrates the Na⁺ storage pathway of Cu-TABQ *via* a three-electron redox pathway that involves Cu²⁺/Cu⁺ and C=O/C–O. Furthermore, Cu-TABQ can undergo a 5 electron transfer reaction and display a high reversible capacity of 776.3 mA h g^{−1}, when the voltage region is extended to 0.1–3.0 V in Fig. S22 and S23,[†] which is similar to Ni-TABQ.¹⁶

The Na⁺ kinetic behaviour of Cu-TABQ is investigated using CV curves at different scanning rates (Fig. 5a). The shape of the CV curve is almost identical as the scan rate increases. However, the cathodic peaks move towards higher values, and the anodic peaks move negatively, which are caused by polarization. The relationship between the peak current (*i*_p) and scan rate (ν) can be described using the following equation: $i_p = a'\nu^b$ ($\lg i_p = \lg a + b \lg \nu$), where *a* and *b* are adjustable parameters. The linear relationship between the peak current and scan rate is depicted in Fig. 5b. The *b* values of O1, O2, and R1 are 0.86, 0.80, and 0.88, respectively, illustrating that the charge storage process is collaboratively controlled by faradaic intercalation and surface pseudocapacitance.⁴⁵ The electrochemical reaction kinetics of Cu-TABQ is studied using the galvanostatic intermittent titration technique (GITT), as shown in Fig. 5c. The calculated Na⁺ diffusion coefficient of the Cu-TABQ cathode is 1.51×10^{-13} to 1.51×10^{-11} cm² s^{−1}, which is consistent with that based on CV curves at different scan rates in Fig. 5a and EIS in Fig. S24.[†] DFT calculations were further applied to analyse the diffusion

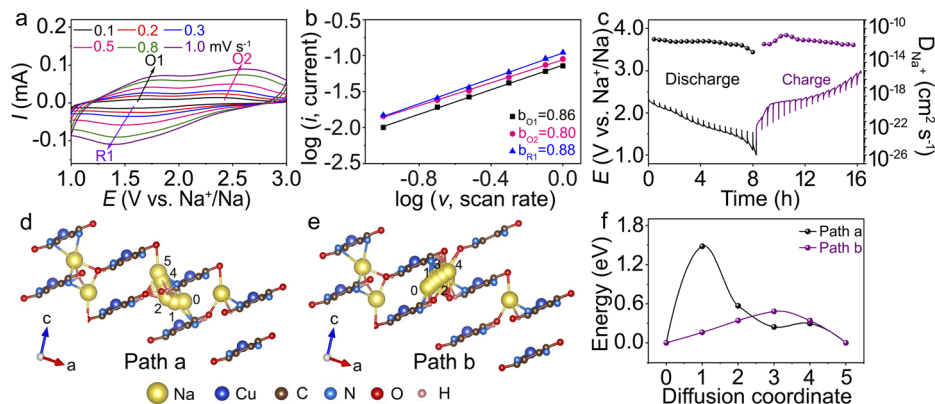


Fig. 5 (a) CV curves of Cu-TABQ at scan rates from 0.1 to 1.0 mV s⁻¹; (b) relationship between different peak currents and scan rates of Cu-TABQ; (c) GITT curves and Na⁺ diffusion coefficient of Cu-TABQ at 50 mA g⁻¹; (d and e) Na⁺ diffusion pathways in Na₃Cu-TABQ (path a across chains, path b along the layers) and (f) diffusion energy barrier.

behavior. The Na⁺ diffusion pathways in crystal structures of Na₃Cu-TABQ are shown in Fig. 5d and e. There are two Na⁺ diffusion pathways in Na₃Cu-TABQ, one is across chains (path a) and the other is along the layers (path b). The calculated migration activation energies of the pathways are plotted in Fig. 5f. The diffusion energy barrier along the layers is much smaller than that across chains, resulting in a faster Na⁺ diffusion and agreeing with the GITT in Fig. 5c. This suggests that the fast diffusion kinetics of Na⁺ in Cu-TABQ is attributed to the distinctive migration pathways and the large interlayer spacing between MOP chains.

Conclusions

In summary, a two-dimensional π -d conjugated coordination polymer (Cu-TABQ) is successfully synthesized by a simple wet chemical method and used as a cathode material for Na⁺ storage. The physicochemical properties, electrochemical properties and sodium storage mechanism of Cu-TABQ are exhaustively studied. It delivers a high reversible capacity of 322.9 mA h g⁻¹ with capacity retention up to 98.7% after 50 cycles. At a high current density of 4000 mA g⁻¹, the capacity can still reach 198.8 mA h g⁻¹. This can be attributed to the stable coordination between Cu²⁺ and the TABQ ligand with π -d conjugation, which greatly improves the electronic conductivity and hence redox activity for Na⁺ storage in Cu-TABQ. By means of FTIR, Raman, XPS, and other characterization methods, the sodium storage mechanism is revealed. It is found that Cu-TABQ has dual redox centers of Cu²⁺/Cu⁺ (one electron) and C=O/C-O (two electrons) for three-electron transfer for each unit and increasing the specific capacity. This shows the great potential of constructing high-performance coordination polymer cathode materials with multiple redox centres and will facilitate the development of advanced cathodes for SIBs.

Data availability

All experimental supporting data and procedures are available in the ESI.†

Author contributions

Liubin Wang: conceptualization, supervision, software, writing – review & editing, project administration. Ningbo Liu: investigation, validation, data curation, formal analysis, writing – original draft. Xiaoying Zhao: software, methodology, formal analysis. Xiaohan Wang: methodology, formal analysis. Tong Zhang: formal analysis. Zhiqiang Luo: supervision, formal analysis, writing – review & editing. Fujun Li: supervision, writing – review & editing.

Conflicts of interest

There are no conflicts to declare.

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