Cerium Hexacyanocobaltate: A Lanthanide-Compliant Prussian Blue Analogue for Li-Ion Storage

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1. INTRODUCTION

Rechargeable lithium-ion batteries (LIBs) have dominated the energy storage and conversion sector for decades. Other ion batteries such as Na, K, and Zn are being considered as alternatives to LIBs, primarily because of the high cost of lithium resources. However, we have a long way to go before we can commercialize the other alternatives. In addition, the small size and lower charge density of Li+ ions go before we can commercialize the other alternatives. In high cost of lithium resources. However, we have a long way to consider the other alternatives.

Prussian blue analogues were synthesized as metal–organic frameworks wherein the transition metal ions were alternately bridged to the ends of cyanide linkers. They have served as functional materials in various fields such as gas absorption, drug delivery, ion conductors, and catalysts. The numerous advantages such as an intrinsic robust crystal structure, facile preparation process, reasonable capacity, and low cost make the Prussian blue analogues suitable for use as electrode materials in LIBs.

The general formula of Prussian blue analogues is $A_xP[R(CN)_6]_{1-x}·nH_2O$ where $A$ is an alkali metal mobile cation, $P$ is the nitrogen-coordinated transition metal ion, $R$ is the carbon-coordinated transition-metal ion, and $x$ is the residual $K^+$ ion content in iron(II) hexacyanocobaltate (FeHCCo), as opposed to that in CeHCCo, results in more residual K+ ions, thereby leading to the occupancy of cavities. As a result, the K+ ion-rich FeHCCo exhibits lower capacities of 55 and 15 ± 3 mAh g$^{-1}$ at 0.1 and 1 A g$^{-1}$, respectively, compared with the K+ ion-deficient CeHCCo that exhibits capacities of 242 ± 3 and 111 ± 3 mAh g$^{-1}$ at the same current densities. This work provides a novel contribution for the exploration of new Prussian blue analogues and bestows a newer concept for electrode material design.
(FeHCCo) was greater than that in CeHCCo, resulting in a significant decay in the capacity. In addition, these two electrode materials exhibit a surface-controlled Li-ion storage nature. This work is expected to provide more insights for exploring not only a new type of Prussian blue analogues but also the design of electrode materials for advanced LIBs.

2. RESULTS AND DISCUSSION

The synthesis process and theoretical structures of FeHCCo and CeHCCo are shown in Figure 1, which illustrates that there are eight subunits where Li+ ions can anchor. Fe2+/Ce3+ ions are coordinated to nitrogen atoms of the cyanide linkers for FeHCCo/CeHCCo. In the products, the charge neutrality requirement normally results in some inherent vacancies of \([\text{Co(CN)}_6]^{3-}\) clusters.32 This structural difference is directly manifested in the optical color (Figure 1, glass vials, insets) due to the different optical properties of the metal–cyanide–metal coordination.

The morphology is studied by scanning electron microscopy (SEM) and transmission electron microscopy (TEM). A distinctive difference in the morphologies of FeHCCo and CeHCCo (Figure 2a,d) can be observed. FeHCCo exhibits spherical NPs with a size of ∼20 nm, while polyhedron-shaped CeHCCo NPs are formed with multiple sizes; the smaller FeHCCo NPs have a smooth surface, which is further illustrated in the TEM images (Figure 2b). Correspondingly, the CeHCCo NPs have a size larger than that of FeHCCo NPs, also displayed in a TEM image (Figure 2e). The crystalline particles and diffraction patterns (Figure 2c,f)

Figure 1. Synthesis process and general structural units of FeHCCo and CeHCCo. Eight subunits are present in each unit for Li+ intercalation. Insets show the optical images of the synthesized FeHCCo and CeHCCo nanoparticles (NPs) in aqueous solutions.

Figure 2. (a) SEM, (b) TEM, and (c) HRTEM images of the synthesized FeHCCo NPs and corresponding selected area electron diffraction patterns. (d) SEM, (e) TEM, and (f) HRTEM images of the synthesized CeHCCo NPs and corresponding selected area electron diffraction patterns.

demonstrate the polycrystalline feature. Furthermore, the atomic arrangement can be observed in the high-resolution TEM (HRTEM) images with an interplanar spacing of approximately 0.44 nm for FeHCCo and 0.41 nm for CeHCCo (Figure S1).

Both FeHCCo and CeHCCo NPs exhibit crystallized features, suggesting that manifestation is independent of the initial reactants. However, the large variation in the particle size and morphology implies that the crystal growth is greatly dependent on the initial reactants, which is consistent with other reports.33 The consistent elements are uniformly distributed throughout the NPs (Figure S2), which is in accordance with the qualitatively verified results by the energy-dispersive X-ray (EDX) mapping (Figures S2b,d and S3).

The crystal structure of FeHCCo determined by X-ray diffraction (XRD, Figure 3a) is well indexed into a cubic structure \((a = b = c = 5.2 \text{ Å})\). However, CeHCCo shows an orthorhombic crystal structure with enhanced lattice constants \((a = 7.4, b = 12.9, \text{ and } c = 13.7 \text{ Å})\) by referring to ErFe(CN)_6 (PDF card: #79-2085, Cmcm (63)), which is expected to facilitate the Li+ ions’ transfer inside the material. The characteristic peaks of both products are well indexed in Figure 3a. A similar occurrence is where crystallization, regardless of the initial reactants, is confirmed by the representative characteristic diffraction peaks in the XRD spectra of both FeHCCo and CeHCCo. The well-constructed frameworks are critical for the Li+ ion intercalation/deintercalation during the charge and discharge processes.

The thermal stability of the synthesized FeHCCo and CeHCCo is examined using thermal gravimetric analysis (TGA, Figure 3b) by varying the temperature from room temperature to 600 °C. The initial weight loss corresponds to the release of zeolitic and coordinated water species.34 A sharp decrease in the weights after 450 °C indicates poor thermal stability for FeHCCo compared with CeHCCo with a platform until 600 °C, demonstrating more bond energy for CeHCCo than that for FeHCCo. The loss of water (∼20% by weight) for FeHCCo is two times more than that of CeHCCo, which is more likely due to the fully bonded \([\text{Co(CN)}_6]^{3-}\) by Ce3+ resulting in less coordinated water.35–37

The molecular formulas based on Fe/Ce are deduced (based on X-ray fluorescence (XRF) results in Table S1) to be \(K_{0.6}\text{Fe}[\text{Co(CN)}_6]_{0.72}\square_{0.28}\) and \(K_{0.6}\text{Ce}[\text{Co(CN)}_6]_{0.99}\square_{0.01}\).
The related qualitative analysis of the constituent elements displayed in Figure S4 agrees well with the EDX results (Figures S2 and S3). A larger amount of the K⁺ impurity, trapped at the subunits to balance the valence state, suggested more cyanide vacancies in the FeHCCo NPs (0.28) than those (0.01) of CeHCCo. The elemental amounts are further conﬁrmed by inductively coupled plasma (ICP, Table S2). As a sequence, the composition of K in the FeHCCo is much higher than that in CeHCCo, which agrees with the XRF results.

To confirm the cyanide-constructed crystal structure, the representative cyanide stretching bands in the Fourier transform infrared spectroscopy (FT-IR) spectra are detected at around 2200 and 500 cm⁻¹ (Figure 3c). The H−O−H stretching band at around 1600 cm⁻¹ and O−H stretching band at around 3400 cm⁻¹ further verify the presence of the residual water molecules and their coordination. The C−N band at around 1210 cm⁻¹ in FeHCCo may be due to the coordination from atmospheric nitrogen.

The Raman spectra (Figure 3d) of the products are obtained from 2000 to 2300 cm⁻¹ to confirm the valence state of cyanide-bonded metal ions as they are sensitive to the oxidation states of the metal ions.39 The obvious bands at 2171 and 2192 cm⁻¹ for FeHCCo and 2163 and 2182 cm⁻¹ for CeHCCo are assigned to the stretching mode of the cyanide bridges. A small Raman shift in the stretching mode of the cyanide bridge is exhibited when the oxidation state of Fe²⁺ changes to Ce³⁺, which results in a higher energy of the stretching band.41

The surface chemical compositions and coordination of the prepared FeHCCo and CeHCCo are measured by X-ray photoelectron spectroscopy (XPS, Figure 4 and Figure S5). In the wide survey spectrum (Figure S5), the O 1s peak in FeHCCo exhibits a higher intensity than that in CeHCCo directly demonstrating more coordinated water molecules due to the [Co(CN)₆]³⁻ vacancies as per the molecular formulas we deduced. There is no obvious difference in the deconvoluted spectra of C 1s and N 1s (Figure S5) where the cyanide functional group within the frameworks is distinctly detected. Two pairs of doublet peaks are exhibited in deconvoluted Ce 3d at binding energies of approximately 885 and 905 eV. The binding energy at 883 eV confirms the existing Ce³⁺ ions inside the frameworks (Figure 4d). The Fe³⁺ in FeHCCo is more likely due to the electron transfer from Fe²⁺ to Co³⁺ through the cyanide bridge.12−46 This electron transfer behavior can be further conﬁrmed by the difference in binding energy of deconvoluted N 1s (Figure S5d,f), together with the formed Co²⁺. Similarly, the Ce⁴⁺ ions in the deconvoluted Ce 3d also suggest the underlying charge transfer.

Based on the above analysis, a feasible growth mechanism for CeHCCo NPs can be obtained. Ce³⁺, [Co(CN)₆]³⁻, and K⁺ are the primary existing species in the initial solution. To lower the total energy raised by the opposite charges, the crystal growth proceeds in a directional manner for CeHCCo. The high-valence Ce−N is expected to be more stable because of its higher crystal field stabilization energy.17,49 With increasing time, CeHCCo with six-fold-coordinated Ce³⁺ possesses a promising and robust structural stability, and its growth proceeds toward the polyhedral crystal morphology. The directional growth can be achieved by taking advantage of the oriented assembly enabled by the high selectivity of facet bonding as pursued by a previous report.49 Due to the ﬁxed vector angles of every two facets within one family such as 90° or 180° for {100} facets, etc., we thus infer that the CeHCCo product directionally grows along the <110> crystallographic orientation with a 60° vector angle. Furthermore, a similar crystal structural morphology was found in a previous report by Lin et al., which was formed via a controllable epitaxial growth by tuning the F⁻ ion additive concentration in the reaction system.50 After a comprehensive analysis, we infer that our CeHCCo product directionally grows along the <110> crystallographic orientation and is enclosed with 12 side facets; thus, the hexagonal structure was ﬁnally formed.

The effect of vacancies in the FeHCCo and CeHCCo nanocrystals on their electrochemical properties is determined by assembling these NPs as the active materials of the cathode in half cells. A redox reaction is exhibited by a pair of peaks at around 3.3 V versus Li⁺/Li (cathodic) and 3.6 V versus Li⁺/Li (anodic) in the cyclic voltammetry (CV) curves (Figure 5a,b) of both FeHCCo and CeHCCo. To ensure that the capacity is mainly contributed by the active materials in the subsequent measurement, a cycling test for the naked graphite current

Figure 4. Deconvoluted XPS spectra of (a) Co 2p and (b) Fe 2p for FeHCCo and (c) Co 2p and (d) Ce 3d for CeHCCo.

Figure 5. CV curves of (a) FeHCCo and (b) CeHCCo in the voltage range of 2.2−4.5 V vs Li⁺/Li with a scan rate of 0.5 mV s⁻¹. (c) Rate performances at current densities of 0.1, 0.2, 0.4, 0.6, 0.8, and 1 A g⁻¹ and (d) cycling measurements (discharge capacity) at a current density of 0.1 A g⁻¹ for FeHCCo and CeHCCo.
collector is performed. The result shows a negligible capacity value (Figure S6c,d). The distinct increases in capacity from 55 ± 3 to 242 ± 3 mAh g⁻¹ at 0.1 A g⁻¹ and 15 ± 3 to 111 ± 3 mAh g⁻¹ at 1 A g⁻¹ are achieved when the initial reactant Fe²⁺ ion is changed into a Ce³⁺ ion (Figure 5c). Furthermore, an enhanced capacity retention for CeHCCo (46%) is achieved compared with that of FeHCCo (27%) with increasing current density from 0.1 to 1 A g⁻¹. The charge/discharge cycling stability displays the improved capacity for CeHCCo compared with that of FeHCCo (Figure 5d). Furthermore, the high performance of CeHCCo can also be demonstrated by comparing with other reported Prussian blue analogues as shown in Table S3 where a comprehensive improvement in both capacity and life span can be attained at a higher current density. Typically, higher capacities are exhibited for CeHCCo than those for both CeHCoFe II with the same cation and Mn₃[Co(CN)₆]₂ with the same anion. This implies that the improved electrochemical capacities are mainly attributed to this combination. In addition, one may ask the case of FeIIIHCCo. For which, FeFeIIHCCo also donates highly improved capacities compared with FeIIIHCCo as shown in Table S3. This agrees that the well-matched valence states are favorable for Li⁺ ion storage. Furthermore, the residual K⁺ impurity and impedance result in the capacity extraction.

To further confirm the final capacity of FeHCCo and CeHCCo, CV curves are scanned at various rates (Figure 6a,b). Well-preserved CV curves are exhibited for both FeHCCo and CeHCCo without obvious distortions. An obvious hysteresis for FeHCCo when scanned at higher rates is clearly shown by the negative shift of cathodic peaks and positive shift of anodic peaks. In other words, the hysteresis in FeHCCo can reduce the capacity and stability of electrochemical reactions. An evaluation of the capacity contribution is analyzed by the relationship between response currents and scan rates at a fixed voltage (3.1 V vs Li⁺/Li) in CV curves (Figure 6c). The quasi-linear relationship for both FeHCCo and CeHCCo demonstrates the surface-controlling Li⁺ ion storage mechanism. Furthermore, the higher response current for CeHCCo demonstrates the lower impedance as measured in the EIS spectra (Figure S8a−c) assisting to increase the capacity.

3. CONCLUSIONS

In summary, we have synthesized CeHCCo and studied the effect of cation nodes with different valence states on the structural and Li⁺ ion storage electrochemical properties of both FeHCCo and CeHCCo. Both spherical FeHCCo and polyhedral CeHCCo exhibit a polycrystalline nature. The reduced K⁺ impurity and impedance result in the capacity increase from 55 ± 3 to 242 ± 3 mAh g⁻¹ at 0.1 A g⁻¹ and 15 ± 3 to 111 ± 3 mAh g⁻¹ at 1 A g⁻¹. This work presents more detailed insights in terms of exploration of new Prussian blue analogues and reactant selection for electrode material fabrication, which undeniably pave the way for the development of advanced Li⁺ ion storage devices.

4. EXPERIMENTAL SECTION

4.1. Preparation of FeHCCo and CeHCCo. FeHCCo NPs were synthesized by a coprecipitation method by simultaneous dropwise addition of 100 mL of FeCl₃ (0.03 M) (Sigma-Aldrich) and 100 mL of K₂[Co(CN)₆] (0.03 M, Sigma-Aldrich) to 100 mL of H₂O under constant stirring; a milky white precipitate ensued immediately. After sonication for 30 min, the suspension was allowed to sit for 1 h. The precipitate was separated and rinsed several times with a large amount of deionized water and ethanol. It was then dried in an oven at 60 °C for further use. The synthesis of CeHCCo was carried out in a similar manner, except that cerium(III) nitrate hexahydrate (Samchun, 0.03 M) was used as the initial
reactant and the codeposition process was performed at 80 °C. During the synthesis of CeHCCo, a transparent solution, without any deposition, was obtained at room temperature, which suggested the requirement of heating treatment. In the deliberation for the appropriate heating temperature, the maximum value attainable for aqueous solutions is 100 °C. However, a transparent solution obtained at 60 °C indicates that a higher temperature is warranted. We thus synthesized CeHCCo at 80 °C, and a white deposition was obtained.

4.2. Physical Characterization. Structural study was performed using XRD (D8-Advance) equipped with Cu Kα radiation at a fixed incident angle of 2θ, XPS (PHI 5000 VersaProbe) using an Al Kα source (Sigma probe, VG Scientifics), Raman spectroscopy (inVia Raman microscope), and FT-IR (Nicolet iS50) spectroscopy. The morphology and constitution were studied using FE-SEM (SUPRA 55VP), and EDX. The constitutions of the prepared FeHCCo and CeHCCo were studied using XRF (ZSX-PRIMUS) and TGA, which was carried out under a N2 flow from room temperature to 600 °C at a heating rate of 10 °C min⁻¹. ICP spectroscopy was performed for confirming the elemental amounts of the products. Each product (0.1 g) was dissolved into 35 mL of NaOH solution for ICP measurements.

4.3. Electrode Preparation. Briefly, a slurry containing FeHCCo or CeHCCo NPs, carbon black (Super P Li), and poly(vinylidene) difluoride in a mass ratio of 7:2:1 was prepared by manually grinding the materials and dispersing them in N-methyl-2-pyrrolidinone (NMP). Before the injection of NMP, the ground mixture was sufficiently dried overnight in vacuum at 80 °C. The weight of the ground mixture was measured before and after vacuum drying to ensure that water was eliminated as much as possible. The working electrode with a mass loading of ~3 mg cm⁻² was prepared by spreading the slurry on a graphite paper current collector (Alfa Aesar) followed by drying overnight in vacuum at 60 °C.

4.4. Electrochemical Characterization. To perform the electrochemical measurements, we prepared a two-electrode half-cell containing the working electrode and sufficient lithium metal (reference electrode) in an argon-filled glove box. LiPF6 was dissolved into 35 mL of NaOH solution for ICP measurements.

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\text{FeHCCo; CeHCCo; XRF mapping for the as-prepared FeHCCo and CeHCCo; XPS for the as-prepared FeHCCo and CeHCCo; XPS spectra for FeHCCo and CeHCCo; CeHCCo; representative charge/discharge curves for FeHCCo and CeHCCo; charge/discharge cycling measurement and representative charge/discharge curves of graphite paper current collector; schematic illustration of the discharge process of FeHCCo and CeHCCo; EIS spectra of FeHCCo and CeHCCo; the corresponding equivalent circuit; XRD spectra of FeHCCo and CeHCCo before and after charge/discharge cycling measurements; quantitative elemental characterization for the as-prepared FeHCCo and CeHCCo; quantitative elemental characterization for the as-prepared FeHCCo and CeHCCo by ICP; and a comparison in capacity of the CeHCCo product and other reported Prussian blue analogues (PDF)}
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