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Morphology-driven ionic pathway engineering in CuCo₂O₄/carbon nanotubes for high diffusion hybrid supercapacitors across diverse electrolyte conditions

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In tandem with conductive carbon nanomaterials, redox-active spinel oxides offer a promising strategy to improve the efficacy of electrochemical energy storage devices. Among them, CuCo₂O₄ (CCO) has attracted considerable attention; however, systematic evaluations of its controlled morphology and diffusion dynamics in varied electrolytes remain scarce. In this study, we engineered CCO nanorods, spherical particles, and their nanocomposites with carbon nanotubes (5, 10, and 15 wt%), named CCO-I, CCO-II, and CCO-III, to investigate diffusion behaviour using the galvanostatic intermittent titration technique across different electrolytic conditions, along with key performance parameters. Electron microscopy verified the successful formation of the desired morphologies, where nanorods provided large surface-active sites and spherical particles offered high volumetric energy density. Electrochemical measurements in 1 M KOH, coupled with theoretical investigation using Dunn's model and determination coefficients (R^2) , revealed a mixed capacitive-faradaic charge storage nature of the samples. Among all variants, CCO-II delivered the best performance, with a specific capacity of 1702.01 C $\rm g^{-1}$ along with an energy density of 113.46 Wh kg^{-1} . It also retained 99.94% capacity after 4500 cycles at 0.4 A g^{-1} , while galvanostatic intermittent titration technique showed balanced diffusion coefficients of $3.9 \times 10^{-11} \text{ cm}^2$ $\rm s^{-1}$ in 1 M KOH and $\rm 4.1 \times 10^{-11}~cm^2~s^{-1}$ in 3 M NaOH. Further, the optimized sample exhibited low internal resistance and high ionic conductivity. Overall, these results highlight the potential of the CCO-II as a promising candidate for high-performance energy storage electrodes.

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1 Introduction

The rapid development of energy storage devices (ESDs) has become a key focus of scientific research, driven by an urgent need to meet global energy demands while addressing environmental concerns. ¹⁻⁴ Given the scarcity of fossil fuel reserves and their detrimental effects on the environment, the search for sustainable and alternative energy storage solutions has become essential. ⁵ Various ESDs, including batteries and fuel cells, have been extensively studied in this context. ⁶ The seventh Sustainable Development Goal set by the United Nations aims to provide clean, reliable, and affordable energy by 2030, further fuelling research into effective solutions. ^{7,8} Among the wide range of ESDs, supercapacitors (SCs), also known as electrochemical (EC) capacitors, have emerged as revolutionary

devices that combine features of batteries and traditional capacitors. $^{9-11}$ These devices are vital for numerous applications such as portable devices, electric vehicles, and renewable energy systems because of their long cycle life, superior power density $(P_{\rm d})$, and fast charge/discharge (CD) capabilities. $^{12-15}$ However, their relatively low energy density $(E_{\rm d})$ compared to batteries limits widespread use, prompting the development of new strategies to enhance their performance. $^{13-17}$

The energy storage mechanism of SCs primarily involves two processes: faradaic and non-faradaic. The faradaic process involves quick redox reactions at the surface of the electrode, while the non-faradaic process results from the electrostatic accumulation of ions at the electrode–electrolyte interface. Early on, SCs mainly utilized carbonaceous materials because they offer a large surface area (SA) and high conductivity, which is suitable for electric double-layer capacitors (EDLCs). Nonetheless, the purely electrostatic nature of ion adsorption inherently limits their $E_{\rm d}$. This led to the development of pseudocapacitors (PCs), which employ transition metal oxides (TMOs) like RuO₂ and MnO₂ and conductive polymers to boost energy storage νia faradaic reactions. The technological

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advancement was further accelerated by hybrid capacitors (HCs), which combine PCs and EDLC materials to enhance both $E_{\rm d}$ and $P_{\rm d}$. 20,21

Binary metal oxides, such as CCO, have garnered attention as pseudocapacitive materials because of their high theoretical capacity, excellent rate capability, eco-friendliness, and costeffectiveness.22-24 The mixed oxidation states of cobalt and copper in CCO improve stability and redox activity through their coordinated actions.25 Despite these advantages, CCO faces challenges like low conductivity, limited specific capacitance $(C_{\rm sp})$, and structural degradation during cycling. To overcome these issues, researchers have explored hybridizing CCO with carbon-based materials to create high-performance composites.22,24 Incorporating carbon nanotubes (CNTs) into electrode materials is a promising approach to address these limitations. CNTs are renowned for their exceptional mechanical strength, electrical conductivity, and high aspect ratio, making them ideal for enhancing structural integrity and charge transfer in composites.26-28 Owing to their unique properties, HCs often include CNTs alongside pseudocapacitive materials to improve overall performance.29

Extensive studies have been undertaken to explore the EC characteristics of CCO-based materials for SC applications. For example, BoopathiRaja *et al.* synthesized CCO nanoparticles *via* hydrothermal methods and achieved a maximum specific capacity ($Q_{\rm s}$) of 764.9 mAh g⁻¹ with a capacitance retention ($C_{\rm R}$) of 94.5%. Bhagwan *et al.* enhanced the $C_{\rm sp}$ of CuCo₂O₄ nanoplates by adding multi-walled CNTs using co-precipitation, reaching a $C_{\rm sp}$ of 1052.7 F g⁻¹. Arulkumar *et al.* fabricated CCO/CuO composites on nickel foam (NF), reporting a $C_{\rm sp}$ of

892.7 F g⁻¹ and nearly 100% cyclic stability after 10 000 cycles.³² Liu *et al.* synthesized CCO/CuO nanoflowers combined with CNTs, resulting in a $C_{\rm sp}$ of 1083.7 F g⁻¹ along with a $C_{\rm R}$ of 98% over 9000 cycles.³³ Semerci *et al.* developed a CCO/rGO composite using a stepwise synthesis method, achieving a $C_{\rm R}$ of 99% after 10 000 cycles along with a $C_{\rm sp}$ of 761.9 F g⁻¹.²² These studies highlight how tailoring structural architecture and hybridization with conductive materials can significantly improve the EC characteristics of CCO for high-efficiency SCs.

Most existing research on CCO has focused on its synthesis methods and the fabrication of composites with other carbonbased materials and metal oxides, primarily evaluating metrics like Q_s , E_d , and P_d . However, no studies have reported on the ionic conductivity (σ), relaxation time (τ), and diffusion coefficient (D_0) of the CCO/CNTs nanocomposite using the galvanostatic intermittent titration technique (GITT). This study aims to synthesize and optimize a CCO/CNTs composite for advanced SCs. Four samples, such as CCO, CCO-I, CCO-II, and CCO-III, were synthesized with CNT ratios of 0, 5, 10, and 15%, respectively. An optimized hydrothermal process, known for producing uniform morphologies, was used to synthesize pristine CCO. Subsequently, a single-step solvothermal process was employed to produce composites with precise control over particle size and CNTs inclusion (CCO-I, CCO-II, CCO-III). This systematic approach enables a detailed investigation into how different CNT concentrations influence the morphology, structure, and EC properties of the composites. Among all samples, CCO-II delivered the best performance, demonstrating excellent cyclic stability, $Q_{\rm s}$, $E_{\rm d}$, and $P_{\rm d}$. Key EC parameters, including σ , D_0 , and τ , were thoroughly examined to understand their

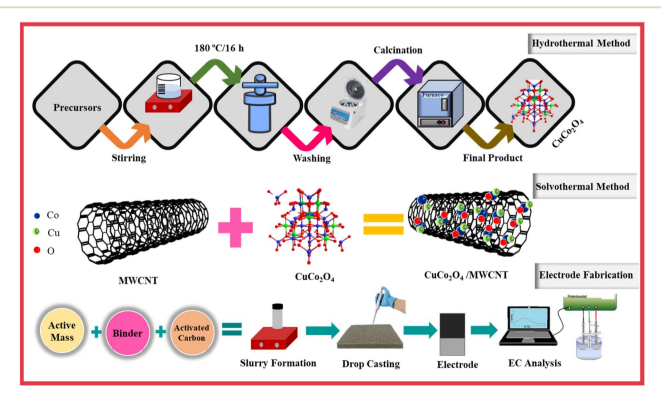


Fig. 1 Schematic representation of the synthesis process of CCO via the hydrothermal method, CCO-I, CCO-II, and CCO-III by the solvothermal method, followed by the electrode fabrication.

impact. These findings underscore the potential of this material for effective and scalable applications in advanced SC systems.

2 Experimental section

2.1 Synthesis of CCO nanoparticles

An optimized hydrothermal approach was used to synthesize the CCO sample, as shown in Fig. 1. The sources and exact quantities of all chemical precursors are provided as SI, and were incorporated as procured, without further treatment, because of their high purity. To form an aqueous solution, stoichiometric quantities of copper(II) nitrate trihydrate, together with cobalt(II) nitrate hexahydrate, were first dissolved in deionized (DI) water. After adding urea (reducing agent) and ammonium fluoride (AF) (fluoride source) to the solution in a 1:1 molar ratio in relation to the metal nitrates, a few drops of ethylene glycol were added. AF was essential in guiding the particular morphology of the final product, while ethylene glycol (solvent) was added to regulate the particle size.16 To facilitate full dissolution and even dispersion of the reactants, the resultant mixture was put on a magnetic stirrer and rapidly agitated for 1 h at room temperature (RT). After that, the uniform mixture was put into an autoclave (180 °C) for 16 h. The hydrothermal process made it easier for the precursors to dissolve and recrystallize under autogenous pressure, resulting in the creation of CCO with distinct morphological and structural properties. After the reaction was complete, the autoclave was cooled to RT. Impurities were eliminated by centrifuging the product and washing it many times with ethanol (dispersing agent) and DI water. The final form of CCO was obtained by drying the material overnight on a hot plate at 60 °C, followed by calcination in a muffle furnace at 550 °C for 3 h to improve crystallinity, eliminate any remaining organic components, and stabilize the material's structure.

2.2 Synthesis of CCO/CNT nanocomposites

Using a facile solvothermal technique, the CCO/CNT nanocomposites were synthesized by integrating different weight percentages of CNTs (5, 10, and 15%) into the CCO framework, as shown in Fig. 1. To ensure equal dispersion, precisely measured amounts of CCO and CNTs were dissolved in 30 mL of ethanol and then ultrasonically treated for 1 h. To create a homogeneous blend, the fluid was then constantly stirred for 2 h at 250 rpm. This thoroughly blended mixture was carefully poured into an autoclave chamber lined with Teflon and sealed tightly. To speed up the solvothermal reaction, the autoclave was heated to 180 °C for 3 h in a muffle furnace. The final product was gathered and repeatedly centrifuged to get rid of any remaining contaminants once it had cooled to RT. After being dried, the refined composite powder was used as the active ingredient in the electrode preparation process.

2.3 Fabrication of the working electrodes

The working electrode (WE) was made by first treating NF (1 $\rm cm^2)$ with DI water and hydrochloric acid solution, and then ultrasonically heating it for 1 h. After sonication, it was washed with DI water and subsequently dried at 60 °C for 1 h to improve

adhesion. To make a clear solution for the electrode slurry, 300 mg of polyvinylidene fluoride (PVDF) was dissolved in 15 mL of dimethylformamide while being continuously stirred. The active mass, PVDF binder solution, and activated carbon (AC) were then combined in a weight ratio of 85:10:5 to produce a homogeneous slurry. To get a uniform consistency, this mixture was agitated for 8 h at 250 rpm. Through the drop-casting method, the slurry was evenly placed on the NF substrate that had been pre-heated, ensuring that the entire surface was covered, as shown in Fig. 1. The coated NF was dried in an oven (80 °C) for 30 min to harden the electrode and evaporate the solvent. The final electrode was made for EC characterization with a 2.5 mg active material loading.

2.4 Characterization techniques

X-ray diffraction (XRD, Shimadzu XRD-6100) was used for crystallographic evaluation of the synthesized materials using a Cu-Kα source with a wavelength of 1.54 Å. A scanning electron microscope (SEM, Nova Nano SEM-450) was utilized for microstructural investigation. A thin layer of gold was applied to the samples to improve conductivity. Nitrogen adsorptiondesorption analysis was performed on a TriStar-II 3020 surface area analyser to evaluate the textural properties, including specific surface area and pore volume, using the Brunauer-Emmett-Teller (BET) method. The EC analysis was carried out using a Corrtest system (CS350M). The complete details of the EC setup are provided in the SI. For EC testing, several techniques were employed: Electrochemical impedance spectroscopy (EIS) to find σ and understand the charge transport, galvanostatic CD (GCD) to evaluate metrics like E_d , P_d , and Q_s , cyclic voltammetry (CV) to investigate charge storage behaviour, and GITT was used to assess the D_0 .

3 Results and discussion

3.1 Structural analysis

XRD was employed to elucidate the crystalline structure, crystallite size, lattice parameters, and degree of crystallinity of the CCO, CCO-I, CCO-II, and CCO-III samples, and the resulting diffraction patterns are given in Fig. 2(a). The peak indexing method outlined by B.D. Cullity was used to interpret these diffraction patterns.³⁴ At the $2\theta = 18.9^{\circ}$, 31.2° , 36.8° , 38.4° , 44.5°, 55.5°, 59.4°, and 65.2°, the diffraction pattern of CCO displayed clear Bragg peaks, which are indexed to the (111), (220), (311), (222), (400), (422), (511), and (440) planes, respectively. The observed diffraction angles and their corresponding peaks exhibited a strong match with the reference pattern (00-001-1155A), thereby confirming the formation of the cubic spinel crystal structure of CCO. The diffraction patterns of the nanocomposites (CCO-I, CCO-II, and CCO-III) showed similar peaks at corresponding angles, with no additional peaks caused by the amorphous nature of the CNTs, indicating consistent and high phase purity. The diffraction pattern of all the assynthesized samples featured sharp and narrow peaks, indicating a high degree of crystallinity. Furthermore, the experimental and reference XRD patterns matched well, confirming

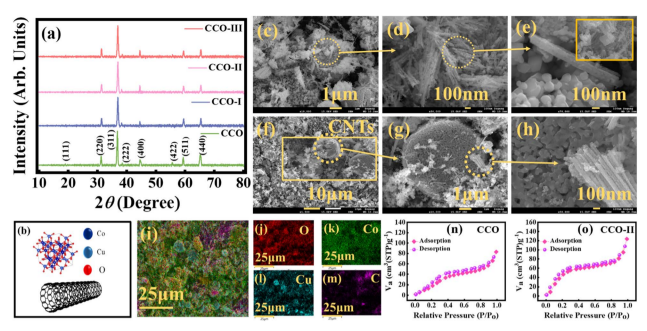


Fig. 2 (a) XRD patterns of CCO with 0, 5, 10, and 15% CNTs, (b) crystal structure of CCO and structure of CNTs, (c–e) SEM images of pristine CCO, (f–h) SEM images of CCO-III, (i) elemental mapping of CCO-III, (j–m) presence of oxygen, cobalt, copper, and carbon detected in the multi-coloured image, and (n and o) BET analysis of CCO and CCO-II.

the material's structural integrity. Lastly, the average crystallite sizes and lattice constants of all the samples were determined employing Scherrer's relation ($D=(\lambda\times 0.9)/(\cos\theta\times\beta)$), and $a=d^2(h^2+k^2+l^2)^{0.5}$, respectively, with the resulting values listed in Table 1.¹⁷ Fig. 2(b) displays the crystal structure of pure CCO and CNTs.

3.2 Morphological analysis

The surface morphology of the as-synthesized samples was examined at an accelerating voltage of 15 kV using SEM at both low and high magnifications, with a working scale bar of 100 nm, and the panoramic view is illustrated in Fig. 2(c–h). The low magnification (100 00×) image of CCO (Fig. 2(c)) shows a heterogeneous morphology, indicating the coexistence of structures with different geometries. High-magnification (700 00×) imaging was employed to better resolve these features, demonstrating the presence of nanorods and nanoparticles in semi-spherical shape as shown in Fig. 2(d and e). The spherical shape of nanoparticles provides high volumetric $E_{\rm d}$, which is beneficial for making lightweight SCs. ¹⁶ In contrast, the nanorods provide several supplementary advantages. Due to their

 $\begin{tabular}{ll} Table 1 & Lattice constants and crystallite size of CCO, CCO-I, CCO-II, and CCO-III composites \end{tabular}$

| | Lattice constants (Å) | | | | |
|---------|-----------------------|-----|-----|-----------------------|--|
| Sample | а | b | с | Crystallite size (nm) | |
| CCO | 8.0 | 8.0 | 8.0 | 15.491 | |
| CCO-I | 8.0 | 8.0 | 8.0 | 15.509 | |
| CCO-II | 8.0 | 8.0 | 8.0 | 15.790 | |
| CCO-III | 8.0 | 8.0 | 8.0 | 15.529 | |

anisotropic geometry, ions and electrons can move more efficiently throughout the electrode surface thanks to shorter diffusion paths. By exposing more active sites, this arrangement enhances ion transport dynamics and stimulates redox reactions. Furthermore, the rod-like shape promotes structural cohesiveness, essential for maintaining mechanical stability over time amid repeated CD cycles. 35,36 Fig. 2(f-h) presents the SEM images of CCO-III. At low magnification (150×), the surface appears more porous than CCO due to CNTs. This increase in porosity enhances ion accessibility and rate performance by facilitating electrolyte penetration. Additionally, a high-magnification (500 00×) image reveals that the CNTs are successfully woven throughout the nanorods and nanoparticles. By providing continuous conductive pathways, this network increases electrical conductivity and reinforces the structure. Overall, the hierarchical morphology positions the composite as the best candidate for advanced SC systems. The spatial distribution of the individual elements forming the nanocomposite material was investigated using elemental mapping. A multicolor composite map, as shown in Fig. 2(i), shows a consistent distribution of elements across the structural matrix, indicating a wellintegrated composition. The individual elemental maps in Fig. 2(j-m), where each element is highlighted with a unique color code, offer even more clarity. These images demonstrate that the elements such as Cu, Co, O, and C are stoichiometrically present in the sample. In mapping images, areas with high concentrations of elements appear brighter, while the darker zones show the absence of a particular element.

3.3 Brunauer-Emmett-Teller investigation

The EC behavior of the synthesized materials is significantly influenced by their specific surface area. The BET analysis,

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which involves the physical adsorption of the N_2 gas, was employed to evaluate the textural properties of pristine CCO and CCO-II, including surface area, pore volume, and pore size distribution. These parameters were obtained from the analysis of the N_2 adsorption–desorption isotherms, as illustrated in Fig. 2(n and o). The pristine and optimized samples exhibit typical N_2 adsorption/desorption profiles along with their corresponding pore size distribution. Based on the isotherm analysis, the specific surface areas of CCO and CCO-II were determined to be 56.92 and 75.01 m^2 g^{-1} , respectively. The comparatively high surface area of CCO-II indicates its enhanced EC performance capability.

At lower relative pressure $(P/P_0 = 0.0-0.1)$, a linear increase in N₂ adsorption reflects the formation of the monolayer along the internal pore surfaces. The subsequent plateau region, followed by a linear portion, signifies the coexistence of the mesoporous and nonporous structures.20 The steep uptake in adsorption across the wider pressure range $(p/p_0 = 0.0-1.0)$ confirms interlayer gas adsorption within the porous framework. The pore size distribution was further evaluated using the Barrett-Joyner-Halenda (BJH). The integrated pore volumes (within the mesoporous range) were calculated to be 42 cm³ g⁻¹ for CCO and 0.27 cm³ g⁻¹ for CCO-II, demonstrating their effectiveness in facilitating ion transport and diffusion. The appearance of type IV adsorption isotherms accompanied by well-defined hysteresis loops confirms the mesoporous nature of both samples. Such structural features make these materials highly suitable for electrode applications, enabling rapid faradaic reactions and efficient charge transport processes.

3.4 Electrochemical analysis

3.4.1 Redox reversibility and kinetic study. CV was employed to elucidate redox behaviour and charge storage mechanisms of CCO, CCO-I, CCO-II, and CCO-III. An EC workstation operating in potentiostatic mode was used to

record CV loops under varying scan rates (SR), specifically 2.5, 5, 7.5, 10, 20, 50, and 100 mV s⁻¹, within an optimal potential window ranging from 0 to 0.5 V. The CV profiles for CCO are shown in Fig. 3(a). The voltammograms exhibit conspicuous redox peaks, indicating faradaic redox processes that lead to pseudocapacitive behaviour.³² The reactions occurring during oxidation and reduction are given in eqn (1)–(3).

$$Cu^{2+}Co_2^{3+}O_4 + 2H_2O + e^- \leftrightarrow 2Co^{3+}OOH + Cu^+OH + OH^-$$
 (1)

$$Co^{3+}OOH + OH^{-} \leftrightarrow Co^{4+}O_2 + H_2O + e^{-}$$
 (2)

$$Cu^{+}OH + OH^{-} \leftrightarrow Cu^{2+}(OH)_{2} + e^{-}$$
 (3)

These successive reactions show the multistep redox behaviour of both Cu and Co species within the spinel framework and correlate with the anodic peaks, spanning the potential range between 0.36–0.45 V as observed in the CV curves. The reduction peaks, which fell between 0.23–0.18 V in the CV profiles, show these processes and validate the reversibility of the redox transitions involving $\mathrm{Co^{4+}/Co^{3+}}$ and $\mathrm{Cu^{2+}/Cu^{+}.^{23,31,37}}$ The Q_{s} values have been derived from the CV data using eqn (4), with CCO exhibiting a value of 1630.74 C g⁻¹.

$$Q_{\rm s} = A \ (m^{-1} \times \nu) \tag{4}$$

here, m is the active mass, A is the area under the curve, and ν is the SR. As the SR increases, the empirically noted alteration in peak potentials indicates kinetic constraints and polarization effects at higher SRs. ¹⁶ Fig. 3(b and c) displays the voltammograms of CCO-I and CCO-II, respectively. Both samples exhibit redox peaks that are consistent with faradaic behaviour. CCO-II showed higher Q_s (2050.59 C g^{-1}), followed by CCO-I (1990.35 C g^{-1}). This enhancement is attributed to the rising CNT concentration in the nanocomposites. CNTs facilitate

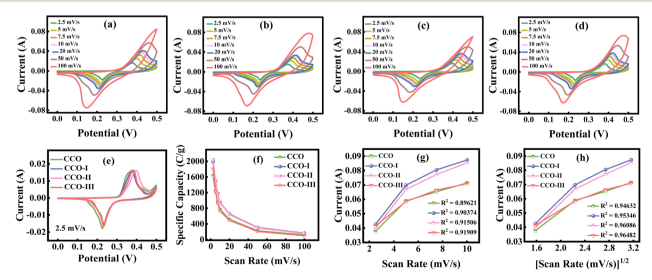


Fig. 3 CV profiles at different scan rates for (a) pristine CCO, (b) CCO-I, (c) CCO-II, and (d) CCO-III, (e) comparison of CV profiles of all samples at 2.5 mV s⁻¹, (f) specific capacity of all samples at different scan rates, (g) current vs. scan rate plot for capacitive contribution, and (h) current vs. (scan rate) $^{1/2}$ plot for diffusive contribution.

conductive pathways, improve electron transport, and reduce total electrode resistance, contributing to greater electrical conductivity. Furthermore, the composite's porous structure, enabled by CNTs, provides effective channels for the transport of electrolyte ions, thereby accelerating redox processes. The increased SA resulting from CNTs also creates more active sites for charge storage.37 The presence of redox peaks, even at high SRs, suggests exceptional rate capability. Additionally, the voltammograms reveal symmetric redox peaks, indicating strong EC reversibility. 16,38 In contrast, CCO-III displays a Q_s of 1815.02 $C g^{-1}$, as evidenced by the CV curves in Fig. 3(d). This decline in Q_s may be due to the uneven distribution of CNTs in the composite material. CNTs may form aggregates due to uneven dispersion, reducing their effective SA and impairing their ability to facilitate efficient charge transfer across the electrode.39 Fig. 3(e) superimposes the CV profiles of all the samples at 2.5 mV s⁻¹ to further clarify the optimum CNTs content. It is clear that the area under the CV curve first rises and reaches a maximum for CCO-II as the CNT concentration increases, before declining for CCO-III. This finding demonstrates that CCO-II has an optimal concentration of CNTs, which improves the EC response. Fig. 3(f) plots the Q_s modulation relative to the SR for all samples. It is evident that as the SR increases, the Q_s noticeably decrease because the reduced diffusion time restricts ion access primarily to the electrode surface. Yet, the electrolyte ions have more time to enter the electrode material's interior pores at lower SRs, the active sites can be fully utilized, leading to larger Q_s values. 16 The Randles-Sevcik relation given in eqn (5) was used to evaluate the dominant charge storage mechanism which explains a diffusion-limited response in which the $I_{\rm p}$ varies according to the $v^{1/2}$.

$$I_{\rm n} = 0.436 \times A \times n \times C^{\rm o} \times F(D_{\rm o} \times v \times F \times n/R \times T)^{0.5}$$
 (5)

here, $I_{\rm p}$ is the peak current (*A*) from the CV, and *A* is the area of the working electrode's surface (cm²). $C^{\rm o}$ is the concentration of the electroactive species in bulk (mol cm⁻¹), and *n* is the number of electrons that are involved in the redox process. $D_{\rm o}$ is the diffusion coefficient of the electroactive species (cm² s⁻¹), and F (96 485 C mol⁻¹) is the Faraday constant. *R* is the universal gas constant (8.314 J mol⁻¹ K⁻¹), ν is the scan rate (mV s⁻¹), and *T* is the temperature in kelvin (K). These traits are very important for figuring out how the redox process works and how the system as a whole behaves in an electrochemical way. The formula above shows how the peak current in a CV curve changes when the scan rate and the diffusion of electroactive species change.

Linear fits were produced for both I_p vs. ν and I_p vs. $\nu^{1/2}$ to identify the dominant charge storage mechanism, and are shown in Fig. 3(g and h). The determination coefficients (R^2) of these plots were then compared. The I_p vs. $\nu^{1/2}$ plot showed a somewhat larger R^2 , indicating that diffusive effects were more prevalent, despite both mechanisms being active. Nonetheless, the high R^2 values in the I_p vs. ν plot also indicate significant capacitive contributions, which show that the material exhibits a hybrid charge storage behaviour. To further validate this, a theoretical model based on the power law given

in eqn (6) was used to verify the hybrid nature of the charge storage mechanism.

$$I_{\rm p} = a v^b \tag{6}$$

In this expression, b is the variable that provides important details about the EC process. If the b-value is around 1.0, it shows capacitive-controlled contribution (like EDL capacitance or surface redox reactions), and if it is near 0.5, it indicates diffusion-limited contribution (like faradaic intercalation reactions). The logarithmic form of the power law, as given in eqn (7), was utilized to compute the b-value. 23,29

$$Log(I_p) = log(a) + b log(v)$$
 (7)

The b-value can be found by linearly fitting $log(I_p)$ against $\log(v)$ and calculating the slope of the line. The slope in this situation represents the predominant charge storage mechanism. A greater capacitive contribution is indicated by a slope that is closer to 1, whereas diffusion-controlled behaviour is more prominent when the slope is closer to 0.5. The calculated b-values for CCO, CCO-I, CCO-II, and CCO-III were about 0.59, 0.65, 0.77, and 0.55, respectively, and their graphs are shown in Fig. 4(a-d). These intermediate values, which fall between 0.5 and 1.0, imply that both diffusion-controlled intercalation and capacitive surface reactions are present in the energy storage process. This result supported the existence of hybrid charge storage behaviour in the system and was in good agreement with the R^2 values. The Dunn's model was used to further quantify the relative contributions of these processes. This approach is particularly useful for differentiating capacitive and diffusive current responses at different SRs. The formula given in eqn (8) is used to deconvolute the total current at a fixed potential:31,41

$$I_{p} = k_{1}v + k_{2}v^{1/2} \tag{8}$$

here, the first term (k_1v) represents the capacitive contribution, and the second term $(k_2v^{1/2})$ denotes the diffusion-controlled contribution. Using this relation, these contributions were quantitatively separated at 2.5 mV s⁻¹ and are graphically shown in Fig. 4(e-h). Furthermore, the contributions were also evaluated at various SRs, as shown in Fig. 4(i-l). The diffusion contribution percentages for CCO, CCO-I, CCO-II, and CCO-III were 84, 90, 92, and 87%, whereas the capacitive contributions were determined to be 16, 10, 8, and 13%, respectively. These outcomes highlight the materials' hybrid charge storage properties and support the previous conclusions drawn from the CV analysis. The diffusive component is due to faradaic redox processes involving intercalation, in which ions permeate into the bulk of the electrode material. However, surface ion adsorption at the interface of the WE and electrolyte is the main source of the capacitive component.28 Furthermore, it was found that the diffusive contribution dominates in all samples at lower SRs. Nonetheless, a discernible increase in the capacitive contribution is observed as the SR increases. A common reason for this phenomenon is the limited time available for ions to diffuse into the electrode bulk at higher SRs.

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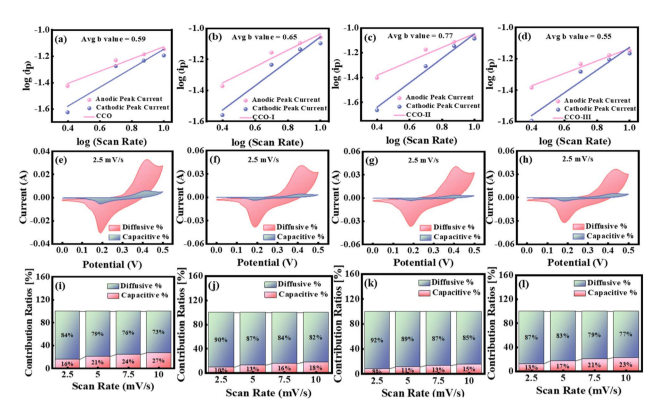


Fig. 4 (a-d) Linear fit curve of log of peak current and log of scan rate for CCO, CCO-I, CCO-II, and CCO-III, respectively, (e-h) percentage contribution of capacitive θ diffusive limited kinetics at 2.5 mV s⁻¹, and (i–l) percentage contribution of capacitive θ diffusive limited kinetics at all

Consequently, charge storage tends to minimize the role of slower intercalation processes by favouring quick surface-level reactions.25,28,31 This change in dominance from diffusioncontrolled to capacitive behaviour as the SR increases demonstrates the electrode materials' exceptional rate flexibility. Taken together, the CV responses, characterized by discrete and well-defined redox peaks, illustrate the hybrid charge storage mechanism and excellent EC reversibility, supporting the electrodes' promise in high-performance energy storage systems.

3.4.2 Galvanostatic charge-discharge analysis. GCD testing was conducted under controlled conditions using a 1 M KOH aqueous electrolyte to gain insights into Q_s , E_d , and the rate capabilities of the synthesized electrode materials, with the workstation operating in galvanostatic mode. The experimental setup and a visual depiction of the GCD mechanism are shown in Fig. 5 and are explained in the SI. Comprehensive GCD curves of CCO, CCO-I, CCO-II, and CCO-III in Fig. 6(a-d) display characteristic nonlinear plateaus, contrasting with linear capacitive systems, indicating that pseudocapacitive faradaic processes mainly govern charge storage.23 This aligns with the EC properties demonstrated by CV. Various current densities $(I_{\rm m})$ (7.0, 9.4, 11.7, 14.1, and 16.4 A g⁻¹) were applied to thoroughly evaluate ion diffusion kinetics and rate performance. Besides the plateau behaviour, an abrupt voltage drop, known as the IR drop, occurs at the start of the discharge curve. This decline reflects the internal resistance of the electrode and electrolyte system and is calculated by measuring the voltage difference between the end of charging and the beginning of discharging on the GCD curve. 42 The GCD curves clearly show that adding CNTs gradually decreases the IR drop, indicating enhanced conductivity. CCO-II had the lowest IR drop value (0.048 V), which is beneficial because a lower IR drop correlates with improved charge transport and reduced internal resistance. Using the IR drop values for each sample, the internal resistance was also calculated via Ohm's law, given in eqn (9).

$$R = V/I \tag{9}$$

The calculated values of resistance for CCO, CCO-I, CCO-II, and CCO-III are 4.07, 3.53, 3.25, and 3.67 Ω , respectively. As the Im increases, the IR drop also increases due to higher resistive losses. Furthermore, the CD behaviour of the composite electrodes changed notably with the addition of CNTs to the CCO matrix. These variations in duration, CD symmetry, and plateau shapes provide critical evidence of how nanostructural changes influence the materials' energy storage capacity. The values of Q_s , E_d , and P_d were calculated with the help of discharge time (Δt) , potential window (ΔV) , and the applied current using eqn (10)-(12):5,43

$$Q_{\rm s} = (I \times \Delta t)/m \tag{10}$$

$$E_{\rm d} = Q_{\rm s} \times \Delta V / (2 \times 3.6) \tag{11}$$

$$P_{\rm d} = (E_{\rm d} \times 3600)/\Delta t \tag{12}$$

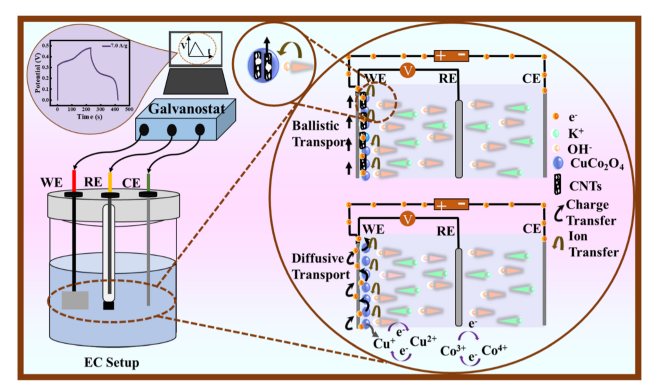


Fig. 5 Schematic illustration of experimental setup and charging-discharging cycles in GCD working mechanism.

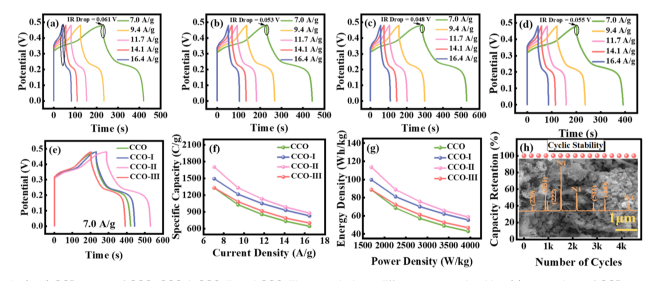


Fig. 6 (a-d) GCD curves of CCO, CCO-I, CCO-II, and CCO-III, respectively, at different current densities, (e) comparison of GCD curves at $7.0 \, \text{A} \, \text{g}^{-1}$, (f) specific capacity of all samples at different current densities, (g) Ragone plot, and (h) cyclic stability, post-cycling XRD, and SEM image of CCO-II electrode.

CCO had the lowest Q_s (1330.40 C g^{-1}) and E_d (88.69 Wh kg⁻¹) values, as well as the quickest discharge time. The comparatively low electrical conductivity and small active SA of CCO may be the causes of its poor performance, which leads to ineffective ion transport during redox cycling. Incorporating CNTs into CCO-I increased its EC response, as evidenced by a longer discharge period and more distinct redox plateaus, resulting in a higher Q_s of 1495.05 C g⁻¹ and E_d of 99.67 Wh kg⁻¹. CCO-II had the best EC performance, with the longest

discharge period and the most pronounced redox characteristics, resulting in an impressive $Q_{\rm s}$ of 1702.01 C g $^{-1}$ and an $E_{\rm d}$ of 113.46 Wh kg $^{-1}$. This improvement can be ascribed to the CNTs' optimal dispersion, which improved ion diffusion and charge transfer throughout the electrode matrix. Nevertheless, performance declined when the CNT contents in CCO-III were increased further; $Q_{\rm s}$ dropped to 1331.77 C g $^{-1}$ and $E_{\rm d}$ to 88.78 Wh kg $^{-1}$, most likely due to CNTs' aggregation that blocked active sites and decreased ion accessibility. In all cases,

increasing $I_{\rm m}$ resulted in a shorter discharge time because higher rates reduced electrolyte ion penetration into the active material's inner pores, limiting charge storage to the outside surfaces.¹⁷ The patterns shown in the samples highlight how crucial it is to maximize the use of electroactive materials by balancing structural integrity with the conductivity enhancement through CNTs content optimization. In Fig. 6(e), all samples' GCD profiles are plotted at 7.0 A g^{-1} for comparison, and Table 2 lists the calculated values of Q_s , E_d , and P_d at the same $I_{\rm m}$. Next, for all electrodes, the change in $Q_{\rm s}$ with $I_{\rm m}$ was studied, as shown in Fig. 6(f). By increasing the $I_{\rm m}$, there is a progressive decline in Q_s values for all the samples. This behaviour is explained by the shorter ion diffusion time at higher $I_{\rm m}$, which restricts electrolyte ions' full access to the inner active sites and reduces their capacity to store charge overall. On the other hand, larger Q_s values are achieved at lower $I_{\rm m}$ because there is enough time for complete ion intercalation across the electrode matrix.17,44,45 Table 3 provides a thorough list of exact Q_s values for each electrode at different I_m . A Ragone plot was used to systematically assess the E_d and P_d of the assynthesized samples, as shown in Fig. 6(g). There is a clear

trade-off behaviour that shows as P_d rises, E_d gradually

decreases. This phenomenon occurs because the full active material capacity cannot be used due to the restricted time available for ion diffusion and charge storage at higher discharge rates.43 CCO-II exhibited exceptional performance, achieving the highest E_d (113.46 Wh kg⁻¹) at a P_d of 1694.11 W kg⁻¹. CCO-II retained significant energy storage capacity at high $P_{\rm d}$, demonstrating its robust rate capability. Table 3 displays the comprehensive comparative E_d and P_d values for the remaining samples. Continuous CD experiments at $I_{\rm m}$ of 0.4 A ${\rm g}^{-1}$ were used to assess the CCO/CNTs electrodes' long-term cyclic capability. The CCO-II hybrid showed exceptional EC durability, retaining 99.94% of its initial Q_s after 4500 cycles, as shown in Fig. 6(h). In essence, the GCD profiles validate the material's exceptional rate capability and long-lasting charge storage behaviour, enhancing its promise for high-performance SC devices. Table S1 provides a comparative analysis of the EC performance of CCO-based electrodes.

Additionally, the phase purity, structural stability, and morphological integrity of the optimized sample (CCO-II) were evaluated through post-cycling XRD and SEM analysis. The XRD pattern in the revised manuscript, presented in Fig. 6(h), show excellent agreement with the pre-cycling data, with no

Table 2 Specific capacity, energy density, and power density at the current density of 7.0 A g^{-1} for CCO, CCO-I, CCO-II, and CCO-III composites

| Sample | Current density (A g ⁻¹) | Discharge time (s) | Specific capacity (C g ⁻¹) | Energy density (Wh kg ⁻¹) | Power density (W kg ⁻¹) |
|--------|--------------------------------------|-----------------------|--|--|-------------------------------------|
| CCO | 7.0 | 188.47 | 1330.40 | 88.69 | 1694.11 |
| CCO- I | 7.0 | 211.79 | 1495.05 | 99.67 | 1694.11 |
| CCO-II | 7.0 | 241.11 | 1702.01 | 113.46 | 1694.11 |
| CCO- | 7.0 | 188.66 | 1331.77 | 88.78 | 1694.11 |
| III | | | | | |

Table 3 Specific capacity, energy density, and power density of CCO, CCO-I, CCO-II, and CCO-III from GCD analysis

| Sample | Current density (A g^{-1}) | Discharge time (s) | Specific capacity (C g^{-1}) | Energy density (Wh kg ⁻¹) | Power density $(W kg^{-1})$ |
|--------|-------------------------------|-----------------------|---------------------------------|--|-----------------------------|
| CCO | 7.0 | 188.47 | 1330.40 | 88.69 | 1694.11 |
| | 9.4 | 109.34 | 1029.15 | 68.61 | 2258.82 |
| | 11.7 | 72.85 | 857.13 | 57.14 | 2823.52 |
| | 14.1 | 52.04 | 734.69 | 48.97 | 3388.23 |
| | 16.4 | 39.12 | 644.40 | 42.96 | 3952.94 |
| CCO-I | 7.0 | 211.79 | 1495.05 | 99.67 | 1694.11 |
| | 9.4 | 129.28 | 1216.81 | 81.12 | 2258.82 |
| | 11.7 | 89.40 | 1051.87 | 70.12 | 2823.52 |
| | 14.1 | 65.95 | 931.19 | 62.07 | 3388.23 |
| | 16.4 | 50.41 | 830.28 | 55.35 | 3952.94 |
| CCO-II | 7.0 | 241.11 | 1702.01 | 113.46 | 1694.11 |
| | 9.4 | 141.52 | 1332.04 | 88.80 | 2258.82 |
| | 11.7 | 96.47 | 1135.04 | 75.66 | 2823.52 |
| | 14.1 | 69.80 | 985.43 | 65.69 | 3388.23 |
| | 16.4 | 53.35 | 878.73 | 58.58 | 3952.94 |
| CCO- | 7.0 | 188.66 | 1331.77 | 88.78 | 1694.11 |
| III | 9.4 | 115.25 | 1084.73 | 72.31 | 2258.82 |
| | 11.7 | 78.08 | 918.69 | 61.24 | 2823.52 |
| | 14.1 | 55.97 | 790.21 | 52.68 | 3388.23 |
| | 16.4 | 42.63 | 702.23 | 46.81 | 3952.94 |

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additional peaks, peak broadening, or peak shifts, confirming the material's phase purity, with no observable degradation and consistent crystallinity. The characteristic diffraction peaks at $2\theta = 44.58^{\circ}$, 51.91°, and 76.74°, corresponding to the (111), (200), and (220) planes, respectively, match well with JCPDS card no. 00-04-08850, confirming the presence of the Ni foam substrate used during electrode fabrication. The post-cycling SEM analysis of CCO-II, shown in Fig. 6(h), demonstrates the retention of its nanorods and semi-spherical nanoplates-like morphology without noticeable structural collapse or surface degradation, indicating excellent morphological stability after the prolonged cycling. These results collectively confirm that the optimized electrode maintains its structural and morphological integrity, supporting its outstanding EC performances.

3.4.3 Electrochemical impedance spectroscopy analysis. EIS was performed to examine the interfacial charge transport, ion movement behaviour, and the impedance response of CCO, CCO-I, CCO-II, and CCO-III. Fig. 7 shows the experimental setup and a schematic illustration of the EIS mechanism. It is shown that at first, a tiny sinusoidal AC input signal is supplied between the WE and RE, and the resulting current response is recorded. The impedance is then determined by the ratio of these two values. After this, a graph of the real vs. imaginary parts of this impedance (Nyquist plot) is subsequently created, providing insights into impedance behaviour. The Randles circuit was utilized to simulate the EC process. The measurements spanned a broad range of frequency (10⁻² to 10⁵ Hz) at 10 mV, and BioLogic EC-Lab software (v11.60) was used for data fitting with a Randles circuit model. Table 4 compiles the extracted fitting parameters. The selected equivalent (Randles)

Table 4 Values of elements for the corresponding fitted circuit, $R_1 =$ R_{s} , $R_{4} = R_{ct}$, $Q_{4} = constant$ phase elements, and $M_{a6} = restricted$ diffusion element

| Samples | $R_1\left(\Omega\right)$ | $R_4\left(\Omega\right)$ | Q_4 (F s ^(a-1)) | $M_{\mathrm{a6}}\left(\Omega\right)$ |
|---------|--------------------------|--------------------------|-------------------------------|--------------------------------------|
| CCO | 1.37 | 5.07 | 0.01 | 0.27 |
| CCO-I | 1.04 | 5.59 | 0.04 | 0.22 |
| CCO-II | 1.02 | 4.65 | 0.08 | 0.18 |
| CCO-III | 1.30 | 6.36 | 0.03 | 0.44 |
| | | | | |

circuit included a charge transfer resistance (R_4) , a double-layer capacitor (C_2) , a Warburg diffusion element (M_{a6}) , a solution resistance (R_1) , and a constant phase element (Q_4) to represent non-ideal capacitive behaviour. This circuit's schematic is shown in Fig. 8(a). Each element corresponds to specific EC processes at the electrode-electrolyte interface. Fig. 8(b) shows the combined Nyquist plot of all samples, while Fig. 8(c-f) displays the individual Nyquist plots of each sample, respectively. The charge transfer kinetics are illustrated by the high-tomid frequency semicircle in the Nyquist plots, while the ion diffusion limitations within the porous electrode are represented by the low-frequency linear tail. R_1 corresponds to the intercept with the abscissa, while the semicircle's width indicates R_4 , which can be calculated using eqn (13).

$$R_4 = (R \times T)/(n \times F \times i_0) \tag{13}$$

here, R_4 is the charge-transfer resistance (Ω), which shows how hard it is to transfer electrons at the interface between the electrode and the electrolyte. The exchanging current density (A cm⁻¹), which is shown by the symbols i_0 , represents how fast

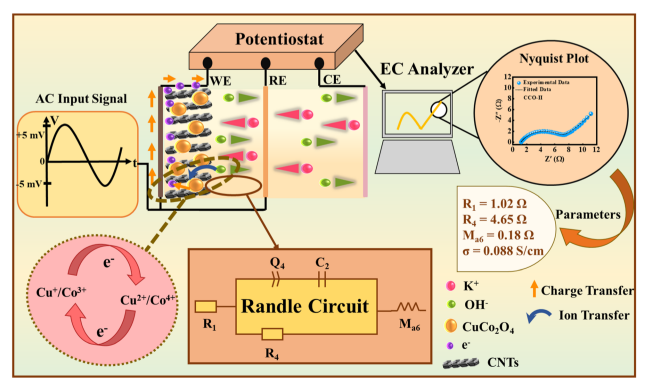


Fig. 7 Schematic illustration of experimental setup, EIS mechanism, resultant Randles circuit, Nyquist plot, and performance parameters.

the redox reactions are happening at equilibrium. 40 We have already established that n stands for the number of electrons, and R stands for the universal gas constant. The slope of the inclined line at lower frequencies, related to the ion transport efficiency via the electrode matrix, can be utilized to estimate the contribution of the Warburg element, which is given in eqn (14).

$$Z_{\rm w} = (A_{\rm w}/\omega^{1/2}) + (A_{\rm w}/i\omega^{1/2}) \tag{14}$$

herein, $Z_{\rm w}$, $A_{\rm w}$, and ω indicate the Warburg impedance, Warburg coefficient, and angular frequency. When ions pass between electrodes and electrolytes, the Warburg impedance (Ω) occurs, and it is represented by $Z_{\rm w}$. The amount of resistance to ion movement through the electrochemical system is indicated by the Warburg coefficient (Ω s^{-1/2}). The complex impedance is measured in imaginary units, and the angular frequency is denoted as ω (rad s⁻¹). These features highlight the diffusion behavior of ions at the electrode-electrolyte interface, as well as impedance diffusion. A frequency-dependent impedance is produced by ions in motion via the electrolyte and electrodes. Because it reveals the rate of connection between surface area, capacitive processes, and slower diffusion-controlled faradaic processes, the Warburg term is significant for hybrid supercapacitors. The angle formed by the horizontal axis and the low-frequency tail is a critical parameter for assessing ion mobility; larger angles predict more effective diffusion pathways. 25,28,46 However, when examining the total impedance response, CCO-II performed the best as it had a favourable M_{a6} (0.18 Ω), a decreased R_1 (1.02 Ω), and a minimized R_4 (4.65 Ω). Furthermore, as seen in Table 4, CCO-II also had the highest Q_4 value (0.08 Fs^(a-1)), representing deviation from ideal capacitive behaviour. The superior EC performance of CCO-II is further supported by a higher Q_4 value, indicating a larger effective SA and an enhanced capacitive response.²⁸ The Bode phase plots (Fig. 8(g)), which illustrate how the phase angle changes with frequency, provide additional insight into

the behaviour of the system. It can be computed by using eqn (15).40

$$\phi = \tan^{-1}(Z'/Z'') \tag{15}$$

Within this equation, Z' (real) and Z'' (imaginary) impedance are functions of frequency. These plots help distinguish between resistive and capacitive traits. A phase angle of -90° typically corresponds to a purely capacitive system, signifying minimal resistive losses. The τ , representing the shortest time needed to release more than 50% of the stored energy, was computed using the relation $\tau = 1/(2\pi f)$, where f is the frequency at which the phase angle is 45°. This characteristic frequency indicates the point of equilibrium where the capacitive and resistive components of the impedance are equal; above this point, the behaviour of the device becomes more resistive. A shorter τ indicates faster ion/electron movement, usually linked to improved rate capability. Table 5 reports the τ values obtained for all samples. Another crucial factor that directly affects SCs' EC efficiency is σ , which governs the movement of ions between electrodes during CD cycles. Effective ion transport, promoted by strong σ , ensures faster kinetics and enhanced rate capability. The following eqn (16) can be used to compute it using the R_1 derived from EIS:

$$\sigma = L/(R_1 \times A) \tag{16}$$

In this equation, L and A represent the electrode thickness and SA, respectively.⁴⁷ Fig. 8(h) shows the σ plot of all samples, where CCO-II can be seen exhibiting the highest value (0.088 S cm⁻¹), indicating superior ion transport characteristics. Table 5 lists the corresponding σ values for each sample. In summary, the EIS results unequivocally show the electrodes' exceptional interfacial stability and electrical conductivity, supporting their potential for EC applications.

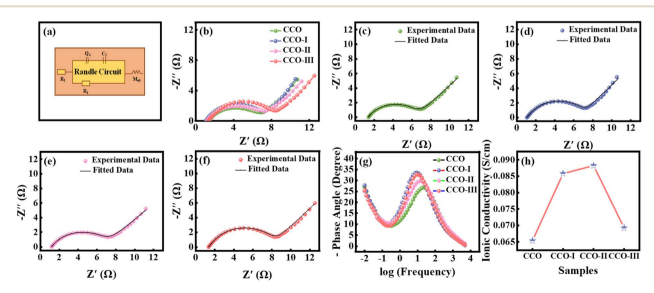


Fig. 8 (a) Resultant Randle circuit, (b) combined Nyquist plot of all samples, (c-f) Nyquist plots of CCO, CCO-I, CCO-II, and CCO-III, respectively, (g) Bode phase plot, and (h) samples vs. ionic conductivity plot.

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Table 5 Calculated values of relaxation time, conductivity, and diffusion coefficients for CCO, CCO-I, CCO-II, and CCO-III composites

| Sample | Frequency (Hz) | Relaxation time (s) | Conductivity (S cm ⁻¹) | Diffusion coefficients (cm ² s ⁻¹) |
|---------|----------------|----------------------|------------------------------------|---|
| CCO | 1.48 | 10.6×10^{-2} | 6.5×10^{-2} | 3.3×10^{-11} |
| CCO- I | 1.03 | $15.3 	imes 10^{-2}$ | 8.5×10^{-2} | 3.6×10^{-11} |
| CCO-II | 1.27 | 12.4×10^{-2} | 8.8×10^{-2} | 3.9×10^{-11} |
| CCO-III | 1.07 | 14.8×10^{-2} | 6.9×10^{-2} | 3.1×10^{-11} |

3.4.4 Galvanostatic intermittent titration technique analvsis. The ion diffusion behaviour of the as-synthesized electrode materials was evaluated using GITT, which calculates D_0 by analysing a material's voltage response to controlled current pulses. This technique involves applying a small, steady current for a short period to induce ion intercalation into the active material. The system then relaxes under open-circuit conditions, allowing it to approach near-equilibrium. 49,50 GITT measurements were conducted in 1 and 3 M KOH and NaOH electrolytes, respectively, within a potential window ranging from 0.4-1 V by subjecting the cell to a 10 min constant current pulse of 1.0 mA. During this period, ions move into and out of active sites, and polarization builds up, causing a characteristic change in potential. After each pulse, the τ of 10 min was employed, which enables the voltage to decrease toward its open-circuit value by stopping the current and allowing the ions within the electrode to redistribute. Each sample underwent four measurement cycles. These conditions provided sufficient time for ion movement and equilibrium, resulting in a more accurate assessment of diffusion properties. The following eqn (17) is used to calculate D_0 .

$$D_{\rm o} = (4/(3.142 \times \tau))(m \times V_{\rm m}/M \times A)^2 (\Delta E_{\rm s}/\Delta E_{\rm t})^2$$
 (17)

In this equation, m and $V_{\rm m}$ indicate the mass and volume of the active material, τ shows the current pulse duration, M indicates the molar mass, A denotes the effective SA of the electrode immersed in the electrolyte, ΔE_t shows the voltage

shift during the pulse excluding IR drop and ΔE_s is the voltage change observed after the system stabilizes following the pulse.51-54 Fig. 9(a-d) displays the corresponding GITT voltage profiles for each sample in 1 M KOH electrolyte. Each profile exhibits the characteristic GITT pattern, which includes a steady potential increase caused by ionic movement and charge storage, followed by a rapid spike in potential at the start of each current pulse due to the IR drop. When the current stops, the IR component is removed, leading to a sudden potential drop. This is followed by a gradual decay as the system relaxes toward the open-circuit potential (OCP) through ionic redistribution. These responses provide insights into diffusion kinetics. To determine D_0 , the values of ΔE_t and ΔE_s were extracted from the GITT graphs. The potential difference (E_1 - E_2), measured just before and after the relaxation phase, yields $\Delta E_{\rm s}$. This process was repeated for each cycle, and the average value was used for accuracy. The change in voltage (E_3-E_4) under a galvanostatic pulse, excluding IR drop, represents ΔE_t , where E_3 is the potential at the start and E_4 is the potential at the end of the pulse. The calculated D_0 for all the samples in 1 M KOH were $(3.3 \times 10^{-11} \text{ cm}^2 \text{ s}^{-1}, \text{CCO}), (3.6 \times 10^{-11} \text{ cm}^2 \text{ s}^{-1}, \text{CCO-I}),$ $(3.9 \times 10^{-11} \text{ cm}^2 \text{ s}^{-1}, \text{CCO-II}), \text{ and } (3.1 \times 10^{-11} \text{ cm}^2 \text{ s}^{-1}, \text{CCO-II})$ III). CCO-II exhibited the highest D_0 (3.9 \times 10⁻¹¹ cm² s⁻¹), based on the GITT curves and the calculated values, owing to its optimized porosity and balanced composition as shown in Fig. 9(e). The subsequent samples confirmed that adding CNTs improves ion transport and reduces resistance, following the

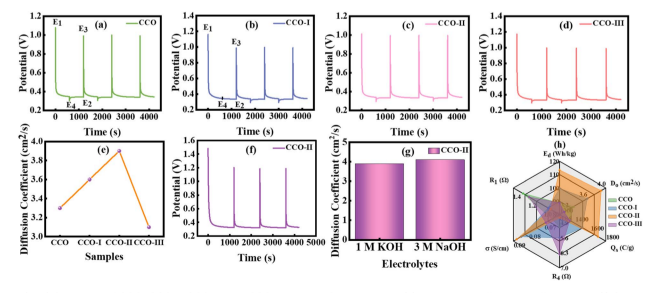


Fig. 9 (a-d) GITT plots of CCO, CCO-I, CCO-II, and CCO-III in 1 M KOH electrolyte, (e) samples vs. diffusion coefficient plot, (f) GITT plot of CCO-II in 3 M NaOH electrolyte, (q) diffusion coefficient of CCO-II in 1 M KOH and 3 M NaOH electrolyte, and (h) radar plot.

trend CCO-III > CCO-I > CCO. Fig. 9(f) shows the GITT plot of CCO-II in 3 M NaOH electrolyte, and it exhibited a $D_{\rm o}$ of 4.1 × 10⁻¹¹ cm² s⁻¹. Fig. 9(g) shows the $D_{\rm o}$ of CCO-II in both 1 M KOH and 3 M NaOH electrolytes. Table 5 contains the values of $D_{\rm o}$ for all synthesized samples in 1 M KOH electrolyte. In conclusion, GITT analysis demonstrates that incorporating CNTs into the electrode structure decreases internal resistance, stabilizes the EC interface, and significantly enhances ionic transport. Fig. 9(h) presents a radar plot that offers an integrated view of key EC metrics such as $Q_{\rm s}$, $E_{\rm d}$, $D_{\rm o}$, $R_{\rm 1}$, $R_{\rm 4}$, and σ for the optimized (CCO-II) electrode, underscoring its potential for use in next-generation SCs.

4 Conclusion

Paper

In conclusion, we synthesized CuCo₂O₄/CNT nanocomposites with varying CNT contents (5, 10, and 15%) to study their diffusion dynamics through GITT and evaluate their potential for energy storage devices. A two-step synthesis process, initial hydrothermal synthesis of CuCo₂O₄ followed by solvothermal incorporation of CNTs, was employed to prepare the nanocomposites with targeted morphologies, comprising nanorods and spherical particles. The combined impact of these structural features enabled excellent charge storage and ion transport, as evidenced by thorough EC experiments. A hybrid (capacitive-faradaic) charge storage mechanism was confirmed by CV in tandem with theoretical analysis (Dunn's model) and R^2 values. EC analysis revealed that the CCO-II electrode exhibited a remarkable $E_{\rm d}$ (113.46 Wh kg⁻¹), $Q_{\rm s}$ (1702.01 C g^{-1}), and P_d (1694.11 W kg⁻¹), demonstrating outstanding charge storage capacity and excellent rate performance. Additionally, it achieved a fast EC response and efficient ion diffusion, as indicated by a high σ (8.8 \times 10⁻² S cm⁻¹), a short τ (12.4 \times 10⁻² s), and a lower R_1 (1.02 Ω) and R_4 (4.65 Ω). The optimized sample displayed a $D_{\rm o}$ of 3.9 \times 10 $^{-11}$ cm 2 s $^{-1}$ in 1 M KOH and 4.1 \times 10 $^{-11}$ cm² s⁻¹ in 3 M NaOH electrolyte. These metrics highlight the effective integration of conductive carbon and redox-active oxide components, resulting in a highly efficient electrode architecture. Overall, the EC behaviour suggests that this optimized hybrid material is suitable for applications that require high capacity, fast kinetics, and good stability. Collectively, these results indicate that the CuCo₂O₄/CNTs composite is a promising and versatile candidate for next-generation high-performance energy storage systems.

Conflicts of interest

The authors declare that there are no financial or any other types of conflicts of interest to declare for this submission.

Abbreviations

| CCO | $\mathrm{CuCo_2O_4}$ |
|---------|--|
| CCO-I | CuCo ₂ O ₄ with 5% CNTs |
| CCO-II | CuCo ₂ O ₄ with 10% CNTs |
| CCO-III | CuCo ₂ O ₄ with 15% CNTs |
| ESDs | Energy storage devices |
| SCs | Supercapacitors |
| EC | Electrochemical |

| $P_{\rm d}$ | Power density |
|----------------------------|-----------------------|
| CD | Charge-discharge |
| $E_{ m d}$ | Energy density |
| SA | Surface area |
| EDLCsElectric double-laver | Electric double-laver |

capacitorsPCs capacitorsPCsPseudocapacitors

TMOsTransition metal oxidesHCsHybrid capacitors $C_{\rm sp}$ Specific capacitanceCNTsCarbon nanotubes $Q_{\rm s}$ Specific capacity $C_{\rm R}$ Capacitance retention

 $\begin{array}{ccc} \mathrm{NF} & & \mathrm{Nickel\ foam} \\ \sigma & & \mathrm{Ionic\ conductivity} \\ \tau & & \mathrm{Relaxation\ time} \\ D_{\mathrm{o}} & & \mathrm{Diffusion\ coefficient} \\ \mathrm{GITT} & & \mathrm{Galvanostatic\ intermittent} \end{array}$

titration technique

DI Deionized

AF Ammonium fluoride
RT Room temperature
WE Working electrode
PVDF Polyvinylidene fluoride
AC Activated carbon
XRD X-ray diffraction

FESEM Field emission scanning electron

microscopy

EIS Electrochemical impedance

spectroscopy

GCD Galvanostatic charge/discharge

CV Cyclic voltammetry

SR Scan rate

 $\begin{array}{ccc} {\rm CE} & {\rm Counter~electrode} \\ I_{\rm m} & {\rm Current~density} \end{array}$

 R_4 Charge transfer resistance C_2 Double-layer capacitor C_3 Warburg diffusion element

 R_1 Solution resistance Q_4 Constant phase element OCP Open-circuit potential

Data availability

Data will be made available on request.

Supplementary information (SI): detailed materials information, XRD patterns with refinement, electrochemical setup description and GCD mechanism, and literature review table (DOC). Electronic copy of CuCo₂O₄ structural data (.cif extension). Original XRD data of all samples (.txt extension). See DOI: https://doi.org/10.1039/d5na00916b.

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