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Introduction

LIBs are one of the most studied systems for clean energy storage due to their compact size, longer life cycle, high efficiency and high energy density.^{1,2} Their ever-increasing demand for the extended range of advanced applications, e.g., electric mobility requires high energy and high power densities.³⁻⁵ Since decades, intense research has been done on transition metal oxides such as CoO, MnO_2 , Fe₂O₃, and Co₃O₄ to replace graphitic anodes that have a limited theoretical capacity of 372 mA h g $^{-1}$ vs. Li $^{\text{+6-8}}$ Among them, Co $_3$ O $_4$ can be produced in large volumes due to its low cost, and it has been demonstrated as a potential high-energy material for application in LIBs with a theoretical capacity of around 890 mA h g^{-1} .⁹ However, the commercial application of $Co₃O₄$ has not been viable yet due to its low electrical conductivity, high volume expansion during charge and discharge cycles, capacity fading at high current density, and poor cyclic stability.^{10,11} Some new synthesis strategies have been demonstrated for producing nanostructures of Co3O4 with different morphologies having different pore distributions and a specific surface area, leading to the high capacity and improved stability of LIBs.¹²⁻¹⁹ The high surface areas and small dimensions of these nanostructures help to improve the interaction of the electrode–electrolyte surface and reduce the Li-ion diffusion length into the solid surface. However, the high volume expansion of $Co₃O₄$ results in the crystal deformation and agglomeration of anode materials, restricting the capacity gains.²⁰ It has been reported that preparing composites of metal oxide nanostructures with conducting carbon networks compensates for the volume expansion and the lacking conductivity, which also enhances the electrochemical performance. This strategy helps in remarkably reducing the volume expansion of $Co₃O₄$; moreover, the carbon matrix improves the conduction of electrons to the current collector.²¹ Different carbon forms such as nano-onions, nanotubes, nanofibers, graphene sheets and 3D porous carbon networks are widely used to prepare composites with $Co₃O₄$, demonstrating anodic capacity enhancement.²¹⁻²⁸ Among these carbon structures, spherical carbon has maximum packing density, low surface-to-volume ratio and high structural

Experimental and theoretical investigations of the effect of heteroatom-doped carbon microsphere supports on the stability and storage capacity of nano- $Co₃O₄$ conversion anodes for application in

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revealed the role of doped supports in controlling the volume expansion upon lithiation.

Conversion-type anode materials have been intensely studied for application in Li-ion batteries (LIBs) due to their potentially higher capacities than current graphite-based anodes. This work reports the development of a high-capacity and stable anode from a nanocomposite of N and S co-doped carbon spheres (NSCSs) with Co_3O_4 (NSCS–Co₃O₄). A hydrothermal reaction of saccharose with L-cysteine was carried out, followed by its carbonization. CSs when used as supports for conversion-type materials provide efficient electron/ion transfer channels, enhancing the overall electrochemical performance of the electrodes. Additionally, the heteroatoms doped in a carbon matrix alter the electronic properties, often increasing the reactivity of the carbon surface, and they are reported to be effective for anchoring metal oxide nanoparticles. Consequently, the NSCS–Co₃O₄ nanocomposites developed in this work exhibit enhanced and stable reversible specific capacity over several cycles. Stable cycling behavior was observed at 1 A g⁻¹ with 1285 mA h g⁻¹ of specific capacity retained after 350 cycles along with more than 99% of coulombic efficiency. This material shows excellent rate capability with a specific capacity of 745 mA h g⁻¹ retained even at a high current density of 5 A g⁻¹. Detailed DFT-based calculations

lithium-ion batteries†

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resistance.²⁹–³¹ However, there is a large scope for further improvements in the performance by introducing defects in carbon matrices, e.g., doping of heteroatoms such as N, B, and S.³²⁻³⁷ These dopant atoms on the surface of carbon materials improve the reactivity, which enhances the lithiation capacity. Additionally, the large atomic size of sulfur atoms can increase the interlayer spacing of the graphitized carbon, create micropores in the carbon matrix and hence improve the charge storage capacity of the carbon material. Moreover, lone-pair electrons of N and S atoms contribute to extending the conjugated electron clouds of the carbon matrix, resulting in increased conductivity that enhances the rate capacity and cycle life of the electrode material.

Taking into consideration the potential effects of the codoped carbon matrix for the stable performance of highenergy conversion-type anode materials like metal oxides, we synthesized a composite of nitrogen and sulphur co-doped carbon spheres and $Co₃O₄$ as an anode material for application in LIBs with enhanced reversible capacity and potential for large-scale production (Scheme 1). N and S co-doping in carbon spheres is found to efficiently inhibit the agglomeration of $Co₃O₄$ NPs and facilitate robust interactions between the electrode and Li-ions, which improves the kinetics of lithium diffusion and reduces the charge transfer resistance. The NSCS– $Co₃O₄$ nanocomposite exhibits excellent rate performance with a specific capacity of 745 mA h g^{-1} at a high current density of 5 A $\rm g^{-1}$. It retains a remarkable and stable discharge capacity of 1285 mA h g^{-1} at 1 A g^{-1} after 350 cycles with more than 99% coulombic efficiency. Paper

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Experimental section

Synthesis of NS co-doped carbon spheres

For the synthesis of NS co-doped carbon spheres, saccharose and L-cysteine in 5 : 1 ratio were used as precursors. In a typical experiment, 10 g of saccharose was first dissolved in 120 ml of de-ionized water followed by the addition of 2 g of L-cysteine

Scheme 1 Schematic of the synthesis of the NSCS– $Co₃O₄$ nanocomposite.

under stirring. The resultant solution was then treated hydrothermally at 180 \degree C for 24 h. After cooling down to room temperature, the obtained product was washed several times with de-ionized water and ethanol by vacuum filtration followed by overnight drying at 80 °C. After drying, the as-prepared material was annealed at 800 °C in an inert atmosphere for 1 h. For the synthesis of non-doped CSs, only saccharose was treated hydrothermally.

Synthesis of a $Co₃O₄$ NP composite with NS co-doped carbon spheres

To synthesize an NSCS– $Co₃O₄$ nanocomposite, 50 mg of NSCS was dispersed in 20 ml of ethanol by ultra-sonication and a separate solution of 0.29 g of $Co(NO₃)₂·6H₂O$ in 20 ml of ethanol was prepared by stirring. Both solutions were mixed under stirring for 30 minutes followed by the addition of ammonia solution (NH₃ \cdot H₂O, 25%). The obtained solution was then transferred into a Teflon-lined stainless steel autoclave and heated at 190 \degree C for 24 h. Finally, the reaction mixture was washed with DI water and ethanol by vacuum filtration followed by overnight drying at 80 °C. For comparison, pure $Co₃O₄$ was synthesized without the addition of NSCSs and CS–Co₃O₄ with non-doped CSs.

Material characterization

Powder X-ray diffraction (p-XRD) patterns were recorded using a Phillips PAN analytical diffractometer with CuK α radiation (λ $= 1.5406$ Å). Transmission electron microscopy (TEM) was carried out using a Tecnai F30 FEG machine operating at an accelerating voltage of 300 kV. The morphology and chemical composition of the ternary hybrid composite were examined using a Quanta 200 3D, FEI scanning electron microscope (SEM). X-ray photoelectron spectroscopic (XPS) measurements were carried out using a VG Micro Tech ESCA 3000 instrument. Thermogravimetric analysis (TGA) was performed using an SDT model Q600 of TA instrument at a heating rate of 10 $^{\circ}$ C min⁻¹ in an air atmosphere.

Electrochemical characterization

The electrochemical properties of the as-prepared electrode materials were examined in a two-electrode coin cell (CR2032) configuration. The $Co₃O₄$ nanostructures, non-doped CSs and $NSCS-Co₃O₄$ were tested as anode materials by preparing their slurry with a conducting carbon additive (super P) and a binder (polyvinylidene difluoride, PVDF) in an NMP solvent. Further, the slurry was coated onto a Cu foil, which serves as the current collector, and subsequently dried at 90 \degree C overnight before assembling the cell in an Ar-filled glove box. The average mass loading of the active material in various electrodes was nearly 1 mg. The coin cells were assembled using a lithium metal foil as the counter electrode, a quartz microfiber paper (Whatman) as the separator and 1 M LiPF₆ in ethylene carbonate (EC)– diethyl carbonate (DEC) $(1:1$ by volume) as the electrolyte. Cyclic voltammetry (CV) was performed using a SP-300 EC Biologic potentiostat at a scan rate of 0.25 mV s^{-1} in the potential window of 0.01–3.0 V. The galvanostatic discharge–

charge cycling of the cells was carried out at different current densities between the potential windows of 0.01 V and 3.0 V using an MTI battery analyser. The electrochemical impedance spectroscopy (EIS) spectra were used to measure the impedance in the frequency range from 100 kHz to 50 mHz with an AC amplitude of 10 mV.

Computational details

All the calculations were carried out using the Kohn–Sham formulation of DFT. The projector augmented wave potential³⁸ is used with the Perdew–Burke–Ernzerhof (PBE) approximation for the exchange-correlation³⁹ and generalized gradient approximation, as implemented in a plane-wave, pseudopotential-based code, VASP.⁴⁰⁻⁴² Furthermore, for the correct representation of 3d orbitals of Co, spin-polarized calculations with the Hubbard U correction (GGA + U) were included. The U parameter (3.32 eV) was selected from the material project⁴³ and further verified for its transferability. The Monkhorst-Pack (MP) formulation was used for the grid of $6 \times 6 \times 6$, for bulk Co₃O₄, which resulted in 108 k-points in the irreducible Brillouin zone (IBZ) and yielded an energy convergence of about 0.005 eV. While, for the composites of $Co₃O₄$ with doped and undoped graphite, a 2D slab was created with a vacuum of 15 \AA , which increased appropriately as lithiation progressed. A gammacentered grid of 6 \times 6 \times 1 was employed for all the surface calculations. This choice of MP grid resulted in 20 k-points in the IBZ. As Li-ions are alloying with $Co₃O₄$ and its composites, the enthalpy of alloy formation ΔE_{mix} is calculated (see eqn (1)) to estimate the stability of the systems:⁴⁴ Nanoscale Advances

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$$
\Delta E_{\text{mix}} = E_{\text{sys}} - \text{Ratio}_{\text{sys}} \times E_{\text{Li}} - (1 - \text{Ratio}_{\text{sys}}) \times E_{\text{bare_sys}} \quad (1)
$$

where $E_{\text{bare_sys}}$, E_{Li} and E_{sys} are the energy of the bare systems $(Co₃O₄, CS-Co₃O₄$, and NSCS–Co₃O₄), Li atom, and Li alloyed systems respectively. The Ratio_{sys} is taken as the ratio of the number of alloyed Li atoms to the total number of the atoms in the system. We modelled the charging of the electrode via the lithiation process by varying the number of Li atoms from 0 to 64 in steps of 8. To model CS– $Co₃O₄$ and NSCS– $Co₃O₄$ systems, the periodic 2D slab was used by layering $Co₃O₄$ on undoped and doped graphite. Sufficient vacuum was added to minimize the interaction between the subsequent periodic images. Furthermore, the bottom two layers of graphite were fixed to represent the bulk of the system and all other layers were allowed to reconstruct. These structures were then optimized using the aforementioned parameters until the forces on the atoms were reduced to 0.01 eV $\rm \AA^{-1}.$

Result and discussion

The crystallographic structures and phase purity of the asprepared materials were investigated by p-XRD. The p-XRD spectra of the synthesized materials are given in Fig. 1a. The p-XRD spectra of CSs and NSCS samples showed the presence of two broad peaks appearing at 24.2 $^{\circ}$ and 43.6 $^{\circ}$ of 2 θ corresponding to the (002) and (101) planes of graphitized carbon respectively. The p-XRD spectra of $Co₃O₄$, CS– $Co₃O₄$ and NSCS–

Fig. 1 (a) p-XRD pattern of CS, NSCS, $Co₃O₄$, CS- $Co₃O₄$ and NSCS- $Co₃O₄$. (b) TGA analysis of $Co₃O₄$, CS– $Co₃O₄$ and NSCS– $Co₃O₄$.

 $Co₃O₄$ consist of peaks at 19.0°, 31.29°, 36.94°, 44.86°, 59.29° and 65.27 $^{\circ}$ of 2 θ corresponding to the (111), (220), (400), (511) and (440) planes respectively, indicating the $Co₃O₄$ spinel structure with a face-centred cubic lattice (JCPDS card no. 42- 1467). Peaks of the CSs and NSCSs are not detected in the composite samples as the carbon spheres are completely covered by $Co₃O₄$ NPs. No residual/impurity peaks could be seen in the p-XRD spectra, indicating the good quality of the samples.

TGA was performed for all five samples in an air atmosphere at a rate of 10 °C min⁻¹ up to 900 °C to quantify the weight percentage of individual components in the composites. In Fig. 1b, the weight loss below 150 \degree C can be attributed to the release of adsorbed moisture and gases from the surface of the samples.²⁵ CSs and NSCSs show complete weight loss after heating at 600 °C due to the formation of $CO₂$ in an oxygen atmosphere. In case of the $CS-Co₃O₄$ and NSCS–Co₃O₄ nanocomposites, weight loss occurred from 400 $^{\circ}$ C to 500 $^{\circ}$ C and then showed a stable TGA profile, indicating the removal of carbon from the composites, and the residual peak corresponded to the weight percentage of $Co₃O₄$. Hence, $Co₃O₄$ and carbon contents in the composites are approximately 60% and 40% respectively.

X-ray photoelectron spectroscopic (XPS) analysis was carried out to find out the oxidation states of cobalt and elemental composition in the NSCS– $Co₃O₄$ composite. Fig. 2a shows the XPS survey spectra of the composite, which displays the presence of Co 2p, C 1s, N 1s and S 2p peaks. In Fig. 2b–e, core level peaks are further deconvoluted to understand the surface composition of the constituting elements. As shown in Fig. 2b, a combination of tetrahedral Co^{2+} and octahedral Co^{3+} contributes to the doublet 2p spectral profile of $Co₃O₄$ separated in high- and low-energy components due to spin–orbit coupling corresponding to $2p_{1/2}$ to $2p_{3/2}$ with a separation of 15.2 eV.^{45,46} The deconvolution of the Co 2p peak at 779.3 eV and 780.6 eV is assigned to $Co^{3+} 2p_{3/2}$ and $Co^{2+} 2p_{3/2}$, respectively. The other spin–orbit components, $Co^{3+} 2p_{1/2}$ and $Co^{2+} 2p_{1/2}$, have been detected at 794.5 and 795.8 eV, respectively. The peaks at 782.1 eV and 797.3 eV may occur due to the chemical shift of the main spin–orbit components, resulting from the chemical interaction of Co cations with surface hydroxyl groups. In addition to these peaks, four small peaks are also observed corresponding to the shake-up satellite peaks of $Co₃O₄$ in the high-binding-energy side of $2p_{3/2}$ and $2p_{1/2}$ transitions,

Fig. 2 (a) XPS survey scan of NSCS–Co₃O₄. Deconvoluted spectra of (b) Co 2p (c) C 1s (d) S 2p and (e) N 1s.

indicating the co-existence of $Co(\Pi)$ and $Co(\Pi)$ on the surface of the material. Fig. 2c shows four deconvolution peaks in the C 1s spectra at 284.6, 285.6, 286.5 and 288.5 eV, which correspond to the sp² hybridized carbon, sp³ hybridized carbon, C–O/C–N and $C=O$ bonds respectively.⁴⁷⁻⁴⁹ The high-resolution S 2p spectra (Fig. 2d) show two peaks at 163.9 eV and 165.2 eV, which may correspond to the presence of C – S – C and C = S respectively and weak peaks at around 168 eV can be related to the oxidized sulfur.50,51 The high-resolution N 1s spectrum (Fig. 2e) with four deconvoluted peaks at 398.5 eV, 399.7 eV, 400.7 eV and 401.6 eV could be attributed to the presence of pyridinic N, pyrrolic N, quaternary-graphitic N and oxidized species respectively.⁵²

The surface and structural morphology of the as-prepared materials was observed by scanning electron microscopy (SEM), and the images are shown in Fig. 3. Fig. 3a represents the micronsized particles of NSCSs. Fig. 3b shows $Co₃O₄$ NPs decorated on the surface of CSs. Here, $Co₃O₄$ NPs are distributed unevenly on the surface of CSs, leading to clustering, whereas in NSCS– $Co₃O₄$ (Fig. 3c), $Co₃O₄$ NPs are well distributed and homogeneously decorated on the NSCS surface forming an extended nanosheetlike structure, which uniformly covers the surface of NSCSs. It is also observable from the images that N and S doping is playing a vital role for the uniform nucleation and growth of $Co₃O₄$ nanoparticles by providing well-dispersed nucleation sites in NSCSs. Morphological control on the uniform growth of $Co₃O₄$ NPs through N and S doping can be highlighted using atomic distribution of different elements over the surface of NSCS– $Co₃O₄$. SEM elemental maps are shown in Fig. S1a-f,[†] which indicate the presence of all five elements $(Co, C, N, S$ and O) that are homogeneously distributed throughout the sample.

Transmission electron microscopic (TEM) characterization was performed on all samples to provide an insight into the inner architecture and crystallographic structure of the asprepared materials. Fig. 3d represents the TEM image of welldispersed N- and S-doped carbon spheres with a diameter ranging between 2 and 5 micrometres. Fig. 3e shows the $Co₃O₄$ nanoparticles with 10-20 nm size. The higher magnification image in Fig. 3f clearly verifies the cubic structure of $Co₃O₄$ NPs with interplanar spacings of 0.28 nm and 0.24 nm, corresponding to the (220) and (311) planes of the face-centred cubic phase of $Co₃O₄$, respectively.⁵³ Fig. 3g shows that N and S codoped carbon spheres are closely and densely surrounded by $Co₃O₄$ NPs, which are extended into sheet-like structures due to uniform coverage over the NSCS surface. The highmagnification image (Fig. 3h) of NSCS– $Co₃O₄$ also reveals the surface boundary of NSCSs and $Co₃O₄$ NPs aggregated like sheets, which indicates the homogeneous distribution of $Co₃O₄$ NPs on the surface of NSCSs. Fig. 3i illustrates the typical selected-area electron diffraction (SAED) pattern of the NSCS– $Co₃O₄$ nanocomposites. The diffraction rings correspond to (220), (311), (400), (511), and (440) planes of polycrystalline $Co_3O_4.^{54}$

Lithium storage properties of $Co₃O₄$, CS– $Co₃O₄$ and NSCS– $Co₃O₄$ composites were examined in a coin cell configuration $versus$ Li/Li⁺ under identical conditions, and the results are shown in Fig. 4a–e. The cyclic voltammetry (CV) measurements for $Co₃O₄$, CS- $Co₃O₄$ and NSCS- $Co₃O₄$ composite electrodes were performed at a scan rate of 0.25 mV s^{-1} in the potential window of 0.01 V to 3.0 V. A distinct and irreversible reduction peak appeared at 0.77 V in the first cathodic scan of the $Co₃O₄$

Fig. 3 ESEM images of (a) NSCS, (b) CS–Co₃O₄ and (c) NSCS–Co₃O₄, TEM images of (d) NSCS, (e) low- and (f) high-resolution images of Co₃O₄ nanoparticles, (g) low-resolution image of NSCS–Co₃O₄, (h) high-resolution image of NSCS–Co₃O₄, and (i) SAED pattern of NS–Co₃O₄.

CV curve (Fig. 4a) due to the reduction of $Co₃O₄$ to metallic Co and the formation of amorphous $Li₂O$ by a conversion reaction. Simultaneous decomposition of electrolytes takes place to form a solid electrolyte interphase (SEI) layer on the surface of the electrode.^{26,27} From the second cycle onwards, cathodic peak split into two distinguishable peaks appearing nearly at 1.01 V and 1.23 V with a lesser intensity, which is due to a multistep reaction during lithiation and the presence of some irreversible transformation that gives structural modification in the first cycle.^{26,27,34} Moreover, two broad peaks are observed during an anodic scan at potentials of 1.35 V and 2.1 V, which could be attributed to the oxidation of metallic Co. The electrochemical reaction mechanism of $Co₃O₄$ involves the formation and decomposition of $Li₂O$, accompanied by the reduction of $Co³⁺$ to Co^{2+} and Co^{2+} to Co and the oxidation of Co to Co^{2+} and Co^{2+}

to $Co³⁺$ through an intermediate of CoO. This multiple electron process can be represented by eqn (2) and (3) .⁵⁵

 $Co_3O_4 + 2Li^+ + 2e^- \leftrightarrow Li_2O + 3CoO$ (2)

$$
3CoO + 6Li^{+} + 6e^{-} \leftrightarrow 3Li_{2}O + 3Co
$$
 (3)

In case of the $CS-Co₃O₄$ electrode, as shown in Fig. 4b, during the first cathodic scan, a reduction peak was observed at 0.71 V and subsequent cathodic peaks (2nd cycle onwards) were observed at 0.87 V and 1.29 V vs. Li/Li^{+} , whereas the anodic peaks were obtained at approximately 1.37 V and 2.14 V. Fig. 4c represents the CV curve of the NSCS– $Co₃O₄$ composite, where the first cathodic peak appeared at 0.85 V and subsequent

Fig. 4 Cyclic voltammetry curves of (a) $Co₃O₄$ nanoparticles, (c) CS -Co₃O₄ and (e) NSCS–Co₃O₄ composite electrodes over 6 cycles at a scan rate of 0.25 mV s⁻¹ and potential range of 0.01–3.0 V vs. (Li/Li⁺). Galvanostatic charge–discharge profiles of (b) Co₃O₄ nanoparticles, (d) CS– $\rm{Co_{3}O_{4}}$ and (f) NSCS– $\rm{Co_{3}O_{4}}$ composite electrodes at a current density of 100 mA g $^{-1}$.

cathodic peaks appeared at 0.96 and 1.34 V, whereas anodic peaks were observed at 1.3 V and 2.1 V. In the NSCS– $Co₃O₄$ composite, the subsequent cycles represent the proper overlapping of anodic and cathodic peaks, an indication of the good electrochemical reversibility of the electrode material.

The galvanostatic charge–discharge (GCD) curves are shown in Fig. 4d, e and f for $Co₃O₄$, CS– $Co₃O₄$ and NSCS– $Co₃O₄$ respectively within a voltage window of $0.01-3.00$ V (vs. Li/Li⁺). The GCD curves were recorded at a current density of 100 mA g^{-1} , and the obtained results are in well accordance with the CV results. In the charge-discharge profiles of $Co₃O₄$, $CS-Co₃O₄$ and NSCS–Co₃O₄, an extended voltage plateau is observed at around 1.1 V, resulting from the lithiation of Co_3O_4 and the formation of Co and $Li₂O$. In the charge curves, the slope region from 1.2 to 2.5 V corresponds to the reversible oxidation of $Co₃O₄$. Discharge capacity values calculated from the second cycle are around 1106 mA h $\rm{g^{-1}}$, 836 mA h $\rm{g^{-1}}$ and 1075 mA h g^{-1} for Co₃O₄, CS–Co₃O₄ and NSCS–Co₃O₄ respectively. A lower specific capacity value of $CS-Co₃O₄$ compared to bare $Co₃O₄$ is probably due to the addition of CSs, but in case of NSCS– $Co₃O₄$, reversible capacity increases due to N and S codoping in CSs, which may provide a large number of edge defects with enhanced electronic conductivity and improved lithium-ion accessibility.³⁴ Another reason could be the homogeneous distribution of $Co₃O₄$ nanoparticles on the surface of NSCSs, which may allow better charge transfer.

Impedance measurements of $Co₃O₄$, CS- $Co₃O₄$ and NSCS- $Co₃O₄$ electrodes were carried out in the frequency range of 100 kHz to 0.05 Hz at room temperature. As shown in Fig. 5a, Nyquist plots of all three electrodes consist of a depressed semicircle at the high-frequency region and an inclined line at

Fig. 5 (a) Impedance spectra of $Co₃O₄$, CS–Co₃O₄ and NSCS–Co₃O₄. (b) Cyclic stability of NSCS-Co₃O₄ at 1 A g^{-1} . (c) Rate performance of $Co₃O₄$, CS–Co₃O₄ and NSCS–Co₃O₄. (d) Relationship between voltage and x mole of Li insertion in the second discharge curve of $Co₃O₄$, CS–Co₃O₄ and NSCS–Co₃O₄. (e) Relationship between real resistance and frequency of $Co₃O₄$, CS–Co₃O₄ and NSCS–Co₃O₄.

the low-frequency region. The high-frequency semicircle is attributed to the SEI film formation, contact resistance, and charge transfer resistance on the electrode/electrolyte interface, whereas the inclined line in the low-frequency region corresponds to the lithium-diffusion process within electrodes.⁵⁶ The charge transfer resistance is calculated by the tting circuit model. It is observed to be lower for NSCS–Co₃O₄ (65.72 Ω) as compared to bare Co₃O₄ (152.5 Ω) and CS–Co₃O₄ (106.1 Ω), implying more conducting nature and faster $Li⁺$ diffusion rate of NSCS–Co₃O₄ nanocomposites. Reduced charge transfer resistance of NSCS– $Co₃O₄$ electrodes is possibly due to the carbon matrix, which reduces the mechanical strain and avoids volume expansion during the charge–discharge process. Additionally, N and S co-doping enhances the electronic conductivity, providing continuous and rapid electron transport, which may accelerate the Li ion storage capacity of the electrode material. Fig. 5b shows the cyclability of NSCS–Co₃O₄ composites at a current density of 1 A g^{-1} for 350 cycles. It can be observed that the as-prepared NSCS– $Co₃O₄$ electrode exhibits long cycle life over 350 cycles with higher specific capacity value and nearly 100% of coulombic efficiency. Over the cycling, more reactive sites of NSCS– $Co₃O₄$ are activated and contributing in lithium-ion storage, which is reflected in the gradually increasing specific capacity up to 919 mA h g^{-1} for initial 25 cycles. Additionally, heteroatom-doped carbon spheres provide more reactive sites, leading to the increased lithium-ion storage capacity while cycling.⁵⁶⁻⁵⁸ Afterwards, for 70 cycles, the capacity value decreases to 720 mA h $\rm{g}^{-1},$ which may be due to the deterioration of the electrode material on the continuous charge– discharge process, resulting in increased charge transfer resistance which is an inherent characteristic of the TMO electrodes.^{35,59} A further increase in capacity value was observed up to 250 cycles (1331 mA h $\rm g^{-1}$), which remained stable over 350 cycles with 96.5% capacity retention as well as >99% coulombic efficiency. This increasing trend in the discharge capacity is probably due to the enhanced kinetics of Li ion diffusion by the gradual activation process and the formation of the polymeric gel-like film with better electrolyte infiltration during cycling.⁶⁰ Moreover, heteroatomdoped carbon spheres reduce the volume expansion during cycling and provide structural stability to $Co₃O₄$ NPs. NSCSs increase the electronic and ionic conductivities of the composite electrode, leading to fast Li⁺ insertion kinetics in Co₃O₄.³⁴ A similar trend was observed in another composite anode $CS-Co₃O₄$ though with lesser specific capacity than NSCS– $Co₃O₄$. Therefore, the rate performance of these materials was analysed after 250 cycles. Fig. 5c illustrates the comparative rate performance of the $Co₃O₄$, $CS-Co₃O₄$ and NSCS–Co₃O₄ electrodes after 251 continuous charge–discharge cycles at 1 A g^{-1} and varying current densities between 0.1 A $\rm g^{-1}$ and 5 A $\rm g^{-1}$. Among all, NSCS–Co₃O₄ electrodes exhibit excellent rate performance with reversible discharge capacities of 1807, 1730, 1616, 1419, and 1151 mA h g^{-1} at 0.1, 0.25, 0.5, 1 and 2 A $\rm g^{-1}$, respectively. Even at a high current density of 5 A g^{-1} , the NSCS–Co₃O₄ electrode delivers a high specific capacity of 745 mA h g^{-1} . However, bare Co₃O₄ electrodes exhibit a specific capacity of 1060 mA h g⁻¹ at 0.1 A g⁻¹ and 384 mA h g⁻¹ at 5 A $\rm g^{-1}$, while CS–Co₃O₄ composite electrodes deliver a specific capacity of 1237 mA h $\rm{g^{-1}}$ at 0.1 A $\rm{g^{-1}}$ and 437 mA h $\rm{g^{-1}}$ at 5 A $\rm{g^{-1}}$. When current density reverted to 0.1 A $\rm g^{-1} , \, NSCS\text{-}Co_3O_4$ gives Nanoscale Advances

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1889 mA h g^{-1} of specific capacity after 292 cycles, whereas Co₃O₄ and CS–Co₃O₄ deliver 1007 and 1356 mA h g^{-1} of specific capacity. The enhanced rate performance of the NSCS– $Co₃O₄$ nanocomposites could be ascribed to the improved electron transfer due to nanostructured cobalt oxides and the increased electronic conductivity provided by the conducting carbon network, fast kinetics, and reduced volume expansion at a higher current density of the NSCS in the nanocomposites.³⁴ Hence, from Fig. 5c, it can be clearly observed that in comparison to $Co₃O₄$ and CS– $Co₃O₄$, NSCS– $Co₃O₄$ shows excellent rate performance at high current density, which is the result of doped nitrogen and sulphur in carbon.

Furthermore, the Li ion diffusion coefficient (D_{Li^+}) of the cell was calculated using the following eqn (4) :⁶¹

$$
D_{\mathrm{Li}^+} = \frac{1}{2} \{ V_{\mathrm{m}} / AF \sigma_{\mathrm{w}} \times \partial E / \partial x \}^2 \tag{4}
$$

where V_m is the molar volume, A is the area of the electrode surface (1.13 cm²), F is the Faraday constant (96 500 coulomb mol⁻¹), σ_w is the Warburg coefficient and $\partial E/\partial x$ is the slope of potential vs. Li concentration plot as shown in Fig. 5d. σ is obtained from the extrapolation of the straight line in the lower frequency region from the semicircle to real axis.⁶²

$$
Z_{\rm re} = R_{\rm e} + R_{\rm ct} + \sigma \omega^{-0.5} \tag{5}
$$

where σ is the slope for the plot of Z_{re} vs. the reciprocal root square of the lower angular frequencies ($\omega^{-0.5}$), as shown in Fig. 5e. The obtained values for σ are 141.422, 39.8 and 27.422 for $Co₃O₄$, CS– $Co₃O₄$ and NSCS– $Co₃O₄$, respectively. As shown in Table 1, the achieved Li ion diffusion coefficients of $Co₃O₄$, CS–Co₃O₄ and NSCS–Co₃O₄ are 1.23 \times 10⁻¹³ cm² s⁻¹, 0.56 \times
10⁻¹² and 1.72 \times 10⁻¹² am² s⁻¹ representively. Among them, the 10^{-12} and 1.73×10^{-12} cm² s⁻¹ respectively. Among them, the NSCS– $Co₃O₄$ composite is having a higher Li ion diffusion coefficient than other electrode materials, which agree with the other electrochemical performances. Heteroatom-doped carbon spheres could increase the kinetics of Li ions on the electrode surface by improving the conductivity of the electrode material, which helps to improve the electrochemical performance of the battery.

To understand the effect of charge–discharge cycling on the cell after 350 cycles, we have performed the impedance measurement (Fig. S4a†) and TEM study (Fig. S4b and c†), and the corresponding figures and discussion are given in ESI.[†]

Computational study

The computed crystal structure of $Co₃O₄$ with the Hubbard U correction was verified against the experimentally reported

Table 1 Values of σ_{w} , X, and D_{Li} for Co₃O₄, CS–Co₃O₄ and NSCS– $Co₃O₄$ as determined from EIS data

Electrode	$\sigma_{\rm w}$ (Ω s ^{-1/2})	$X \text{ (mol cm}^{-3})$	$D_{\rm Li}$ (cm ² s ⁻¹)
Co ₃ O ₄	135.96	0.1852	0.123×10^{-12}
CS -Co ₃ O ₄	61.24	0.1768	0.56×10^{-12}
$NSCS$ -Co ₃ O ₄	27.42	0.1394	1.73×10^{-12}

structure. The deviation between the computed and experimental lattice parameters is found to be less than 0.1%, con firming that the parameters and methodology utilized are appropriate. As stated, the Li-atoms were added to the structure, in steps of eight atoms. Hence, the initial structure has 8 Liatoms, while the fully lithiated $Co₃O₄$ structure has 64 Liatoms. The lithiation strategy was modeled in accordance with eqn (2) and (3) representing the chemical reaction. Some representative models are shown in Fig. 6a–d for $Co₃O₄$, Fig. 6e– h for $CS-Co₃O₄$ and Fig. 6i–l for NSCS–Co₃O₄.

A trend of increase in the volume of the crystal structure $(Co₃O₄)$ is observed with the increase in number of Li-atoms. The volume of the bulk system is 573.57 \AA^3 , while the volume of the fully lithiated structure is 1421.70 \AA ³. An increase of approximately 248% is observed. $CS-Co₃O₄$ and NSCS–Co₃O₄ structures were also lithiated to investigate the effect of the substrate as well as doping on the volume expansion and overall functioning of the electrode. Volume expansion upon lithiation, in $CS-Co₃O₄$ and NSCS–Co₃O₄ systems, is significantly lower than that of bulk $Co₃O₄$. The initial volume of CS– $Co₃O₄$ and NSCS–Co₃O₄ systems was 790.832 \AA ³ and 808.515 \AA ³ respectively, which increased to 1628.31 \AA^3 and 1618.36 \AA^3 , respectively upon lithiation. The percentage volume expansion was found to be 206% and 200% for $CS-Co₃O₄$ and NSCS–Co₃O₄ respectively. A considerable downshift in volume expansion upon lithiation underlines the role of support in restraining the volume as compared to unsupported $Co₃O₄$. A downshift in the volume expansion and the corresponding ΔE_{mix} value reflect the enhanced stability of the system. In Fig. 7a, the ΔE_{mix} value (on the left y-axis) and change in volume (on the right y-axis) have been plotted against the number of Li atoms added to the system. The change in volume is considerably less in the composite systems compared to $Co₃O₄$. The support provides additional strength to the lattice of $Co₃O₄$ by providing an anchoring point. The presence of the support for $Co₃O₄$ results in bonding between C and Co as well as between C and O, thus restricting the expansion of at least Co and

Fig. 6 (a–d) Bulk $Co₃O₄$ with 0, 16, 40, and 64 Li atoms respectively. The pink-coloured balls represent Co atoms, red-coloured balls represent oxygen and violet-coloured balls represent Li atoms. Comparison between (a) and (d) shows a complete reordering of the $Co₃O₄$ crystal lattice. (e–h) $CS-Co₃O₄$, with 0, 16, 40, and 64, Li atoms alloyed, respectively. Similarly, $(i-1)$ NSCS–Co₃O₄, with 0, 16, 40, and 64 Li atoms, respectively.

O atoms in contact with the support upon lithiation. This does reflect the lesser volume expansion for supported systems. The support also affects the functioning of $Co₃O₄$ in terms of slowing down the reaction from $Co₃O₄$ to CoO-like phase. This could be understood from the analysis of Co–O bonds as a function of lithiation in these three systems. In Fig. 7b, we have shown the shortest Co–O bonds as a function of increasing Li contents in the system. In the case of $Co₃O₄$, a sudden shift in the number of short Co–O bonds is observed, while the number of Li atoms within the system changes from 8 to 16. There are about 103 short Co–O bonds in the $Co_3O_4 + 8$ Li system. This number drastically reduces to 55 in case of the $Co₃O₄ + 16$ Li system. Furthermore, the longrange order in the system is lost as evident from the continuous distribution of the bond lengths. This sudden shift marks the transition from $Co₃O₄$ to CoO-like phase. It is interesting to note the effect of the support on the Co–O bond lengths in $CS-Co₃O₄$. The bare (or unlithiated) CS– $Co₃O₄$ has less number of Co–O short bonds compared to unsupported $Co₃O₄$ because Co, as well as O, gets bonded with the support. However, upon lithiation, the transition is slowed down, as reflected from the Co-O bond lengths shown in Fig. 7c. For 16 Li atoms in $CS-Co₃O₄$, there are 64 short Co–O bonds, which were about 55 in case of unsupported $Co₃O₄$, indicating 10–15% reduction in Co–O short bonds. This marks the negative effect of support which although helps in controlling the volume expansion but only at the cost of Li holding capacity and therefore showing reduced performance. However, when we compare it with NSCS– $Co₃O₄$, the number of Co–O short bonds upon addition of 16 Li atoms are about 57, indicating that the behaviour of NSCS– $Co₃O₄$ is similar to that of unsupported $Co₃O₄$ (Fig. 7d). Thus, $Co₃O₄$ placed over N and S doped carbon reduces the volume expansion and does not compromise on the Li holding capacity as much as in the case of $CS-Co₃O₄$. **Paper**
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The specific capacity of the $Co₃O₄$ system can be related to its ability to undergo reactions in order to form CoO and Li₂O and finally CoO getting converted into metallic Co (as given by eqn (1) and (2)). As lithiation progresses, $Co₃O₄$ gets converted into CoO and finally into metallic Co, as discussed above. In terms of effective charge on Co, it changes from Co^{3+} to Co^{2+} in CoO and then to metallic $Co⁰$ upon complete lithiation. This transition also reflects the Bader charges as well as Co-O bond lengths as a function of lithiation.

Our detailed DFT calculations demonstrate that the presence of the support below the $Co₃O₄$ nanoparticle does play an important role, in controlling both the volume expansion and the specific capacity. The carbon support holds the Co and O atoms that not only restrain volume expansion but also decrease the interaction of Li atoms with cobalt and oxygen, which would have otherwise lead to the conversion of $Co₃O₄$ into CoO and further into metallic Co. However, the presence of dopants have helped in controlling the formation of C–Co bonds, such that Li atoms can interact in a better way with the Co and O atoms to form CoO and metallic Co without resisting its capability to reduce volume expansion.

In summary, the excellent electrochemical performance of NSCS– $Co₃O₄$ could be attributed to the synergetic effect of $Co₃O₄$ and N and S co-doped carbon spheres. Nanoparticles of $Co₃O₄$ effectively reduce the path length for lithium-ion

Fig. 7 (a) Enthalpy of alloy formation (ΔE_{mix}) and change in volume of Co₃O₄, CS–Co₃O₄ and NSCS–Co₃O₄ with respect to the number of Li atoms. The Co–O distance in (b) Co₃O₄, (c) CS–Co₃O₄, and (d) NSCS–Co₃O₄ as a function of lithiation. The Co–O bonds in crystalline Co₃O₄ and CoO are plotted for reference. In case of bulk-Co₃O₄ and NSCS–Co₃O₄, the CoO phase is more dominant from 16 Li onwards, whereas, in case of $CS-Co_3O_4$, the formation of the CoO-like phase is delayed and appears to dominate from 24 Li onwards.

migration during the charge/discharge process. Heteroatom doping offers suitable pathways for electron transfer and provides a good conductive matrix for $Co₃O₄$ nanoparticles. NSCSs also act as a buffer substrate to accommodate the volume changes of $Co₃O₄$ in the process of lithium-ion insertion/ extraction. Nitrogen and sulphur doping introduce defects, thus leading to the formation of disordered carbon structure, which tends to be electron accepting and provides anchor sites for the $Co₃O₄$ nanoparticles.

Conclusions

An NSCS– $Co₃O₄$ nanocomposite has been synthesized as an anode material with a significant potential for application in lithium-ion batteries as a high-energy material. Doped N and S atoms enhance the electronic conductivity and also provide binding sites for the facile deposition of a large number of $Co₃O₄$ nanoparticles. It also provides the channels for charge and ionic transport, which effectively alleviates the aggregation of $Co₃O₄$ in the Li⁺ insertion/extraction. As a result, NSCS– $Co₃O₄$ shows excellent rate performance and long cycle life. Among all

the three electrodes, NSCS– $Co₃O₄$ exhibits less charge transfer resistance with a higher Li ion diffusion coefficient value of 1.73×10^{-12} cm² s⁻¹.

Conflicts of interest

The authors declare no competing financial interest.

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