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Sustainability spotlight

Reduced graphene oxide from dead Li-ion batteries with β -Co(OH)₂ as a potential electrode for enhanced charge storage capabilities[†]

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Supercapacitive studies were performed on two composites of reduced graphene oxide $(rGO)/\beta$ -Co(OH)₂ of the same weight composition, with the difference among them being the graphite source, one being the graphite recovered from spent Li-ion batteries (GC150) and the other being the pristine graphite (GPGC150). Results show that GC150 exhibits superior energy storage performances compared to GPGC150. However, the rate capability of the GPGC150 was found to be higher than that of GC150. Both the composites exhibited better cyclic stabilities up to 10 000 cycles at a current density of 10 A g⁻¹. GC150 exhibited zero deterioration while GPGC150 exhibited 9.23% of deterioration. The energy storage parameters, *viz.*, specific capacitance, specific capacity, specific energy, specific power, and coulombic efficiency, exhibited by GC150 at a current density of 1 A g⁻¹ are 36 F g⁻¹, 43 C g⁻¹, 7.16 W h kg⁻¹, 0.77 kW kg⁻¹, and 94.99%, respectively, in a symmetric two-electrode system. The rGO/ β -Co(OH)₂ composites synthesized using recovered graphite as the source for rGO (GC150) exhibit supercapacitance better than their analog that is synthesized using pristine graphite as the source for rGO by the synthetic route followed in the present study.

The growing world is also accompanied by the growth of waste piles all across the environment; inadvertently, the idea of waste to wealth is being taken up as one of the crucial steps to control these waste piles. These waste piles are of different natures; some are bio-degradable while some are not, and some are toxic to the environment to a greater extent. One such waste pile that is given the least importance in recycling is spent Li-ion batteries (LIBs), which are being piled up globally after their use in various gadgets. Our study focuses on recycling LIBs and using their components again in energy storage devices such as supercapacitors. The main aim was to recover the graphite from spent LIBs, use it as a base material for the synthesis of reduced graphene oxide (rGO), and integrate it with other faradaic materials in facile methods. In this work, one such rGO composited with cobalt hydroxide and its supercapacitive behavior is presented. These results highlight the importance of recycling spent LIBs and are highly relevant to UN SDGs 7 (Affordable and Clean Energy) and 13 (Climate Action).

1 Introduction

The production, utilization, and disposal of Li-ion batteries (LIBs) are at their maximum level, presently, due to their high energy characteristics. The most used anodic material in them is graphite due to its easy availability, inexpensiveness, electrical conductivity (band gap \sim 0.4 eV), chemical stability, mechanical stability, moderate theoretical capacity of 372 mA h g⁻¹, and reversible and effective intercalation

reactions that it undergoes with Li-ions.¹ Producing one million electric vehicles is estimated to require $\sim 75 \times 10^6$ kg of graphite.² LIBs produced using such a huge amount of graphite are just discarded as landfills once their purpose is served, and LIBs get to a stage where they are called spent. It is foreseen that by the year 2030, 11 million tonnes of LIBs will be discarded.³ Of this huge amount of LIBs discarded, only <5% is recycled due to the high cost involved in their recycling process.³ On the other hand, the present supply of raw materials for the production of LIBs is perceived to be inadequate, considering the growing demand for LIBs.³ At this juncture of pressing situations, knowing the properties of constituent materials of LIBs after their life span and considering their possible reuse is essential so that the anticipated inadequacy of raw materials for producing LIBs can effectively be dealt with.

Now, considering the structural aspects of graphite, which is the most common anode material in LIBs, its reuse will come in handy. As its structural aspects are not substantially disturbed

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[†] Electronic supplementary information (ESI) available: Recovery of graphite from spent Li-ion batteries and synthesis of graphite oxide (GO) by modified Hummers' method (S1), fabrication details (S2), characterizations and measurements techniques (S3), formulae involved in determining the crystallite size, number of graphene layers per domain or crystallite (eqn (S1)–(S3)), Fig. S1 (XPS studies of GPGC), Fig. S2 (FE-SEM of GC), Fig. S3, Tables S1, S2, Fig. S4, Fig. S5, eqn (S4) to (S10). See DOI: https://doi.org/10.1039/d4su00133h

or altered even after several cycles. The early studies conducted on the structural properties of graphite recovered from spent LIBs have revealed that its layered structures, electrical conductivity, and chemical and mechanical stabilities are intact without much alterations.⁴ This one aspect could be the driving force behind removing the graphite from spent LIBs and reusing it in new charge storage systems like supercapacitors. It is also imperative to consider that the graphite recovered from LIBs would possess residues of Li-ion containing electrolytes, binders like polyvinylidene fluoride (PVdF) and Nafion, and corrosion products of the current collector (Cu), which is possible in the long course of the use of LIBs. Such impurities should be removed in an attempt to reuse it. There are other reasons, such as the electrolyte (LiPF₆ in ethylene carbonate, propylene carbonate, etc.) and the heavy metals (Mn, Co, Ni, etc.) present in them, that cause severe water and soil pollution.³ The expanded d-spacing of recovered graphite⁴ facilitates its conversion to other derivatives (like graphite oxide, reduced graphene oxide (rGO), and graphene) and necessitates the reuse of the parts of LIBs.

The use of graphene (contains no oxygen functionalities) and rGO (contains minimal oxygen functionalities, yet resembles the properties of graphene) in supercapacitors owing to their large surface area, electrical conductivity, mechanical stability, chemical stability, high charge-carrying ability, etc., is well established.⁵ Co(OH)₂ is a well-known faradaic material known for its use in batteries and supercapacitors due to the facile synthesis and electrochemical processes that it undergoes.6 It exhibits good electrical conductivity7 and possesses a theoretical capacitance of 3460 F g⁻¹.^{8,9} There are two polymorphs of $Co(OH)_2$, which are in the α and β forms. The α -form possesses a hexagonal hydrotalcite structure, and the β-form possesses a hexagonal brucite structure, and their electronic band gaps are calculated to be 2.45 and 2.85 eV, respectively.10 In addition, α -Co(OH)₂ exhibits larger interlayer spacing than β -Co(OH)₂ (ref. 11) to exhibit faster intercalations of electrolytic ions in between the layer structures. However, β -Co(OH)₂ is facile in its synthesis; therefore, its electrochemical energy storage properties should also be explored. A few of the Co(OH)₂ containing supercapacitors and their specific capacitances are as follows: $Co(OH)_2/CNT/Ni$ (61 F g⁻¹ at a current density of 0.15 A g⁻¹),⁷ $Co(OH)_2/CoSe_2$ nano-sheet (349 C g⁻¹ at a current density of 1 A $g^{-1})\text{,}^{12}$ Co(OH)_2/Co_3S_4 nano-sheet (231 F g^{-1} at a current density of 1 A g⁻¹),¹² CoOOH/Co(OH)₂ (104 F g⁻¹ at a current density of 1 A g⁻¹),¹³ NiCoP/Co(OH)₂ (118 F g⁻¹ at a current density of 1 A g^{-1}),¹⁴ NiCo₂S₄/Co(OH)₂ (101 F g^{-1} at a current density of 0.5 A g^{-1}),¹⁵ β -Co(OH)₂ nano-flower (416 F g^{-1} at a current density of 1 A g^{-1}),¹⁶ and sheet-like Co(OH)₂ (417 F g^{-1}).¹⁷ These early studies indicate the fruitfulness of using $Co(OH)_2$ as a supercapacitor electrode material.

Therefore, in this study, an attempt has been made to synthesize $rGO/Co(OH)_2$ composite electrode materials for supercapacitors using the graphite recovered from LIBs as the source for rGO at different weight compositions of both rGO and $Co(OH)_2$ (named as GC composites) and to compare the supercapacitance of the weight composition that provided maximum performance with the $rGO/Co(OH)_2$ composite of

similar weight composition synthesized using pristine graphite as a source for rGO (named as the GPGC composite). The supercapacitance of both GC and GPGC composites was evaluated using two electrodes containing a symmetric supercapacitor cell setup and 1 M H_2SO_4 as the electrolyte. The findings and the discussions are detailed in the forthcoming sections.

2 Experimental

2.1. Synthesis of rGO and Co(OH)₂ composite using recovered graphite (GC)

In the synthesis of GC, in a 250 mL round-bottomed (RB) flask, 35 mg of GO that was synthesized using the recovered graphite (RG) (Section S1[†]) and 5.34 mL of 0.1 M CoSO₄. 7 H₂O and about 75 mL of distilled water were taken and sonicated for 45 minutes. After that, the pH of the reaction mixture was enhanced to 14 using 6 M KOH and an adequate amount of N₂H₄. H₂O was added. The reduction reaction was conducted for 2 hours with constant stirring at 90 °C. Then, the end product was cooled, washed multiple times with distilled water (DW) and ethanol, extracted with ethanol, and dried at room temperature to get GC. The same method was followed with different amounts of CoSO₄. 7H₂O by keeping the amount of GO constant to obtain GC composites with varying weights of Co(OH)₂.

2.2. Synthesis of rGO and Co(OH)₂ composite using pristine graphite (GPGC)

For the synthesis of GPGC, the same synthetic procedure as that of GC with the exception of usage of GO synthesized using pristine graphite instead of GO synthesized using RG was followed. The fabrication of synthesized GC and GPGC into symmetric supercapacitors is detailed in the ESI (Section S2†). The structural and electrochemical characterization details of GC and GPGC are provided in ESI (Section S3).†

3 Results and discussion

3.1. Structural characterizations

3.1.1 XRD analyses. The XRD patterns of the GC and GPGC (Fig. 1a) possess the characteristic peaks of Co(OH)₂ and rGO, confirming the successful synthesis of rGOs and Co(OH)2 composites. The observed peaks at 19.04°, 32.49°, 37.94°, 51.39°, 57.98°, 61.62°, 68.11°, 69.54°, and 71.43° are corresponding to the (001), (100), (101), (102), (110), (111), (200), (103) and (201) planes of $Co(OH)_2$, respectively (JCPDS file no.: 30-0443 and 01-0357). The peaks at 61.62°, 68.11°, and 69.54° correspond to Co(OOH) (JCPDS file no.: 07-0169), indicating the presence of a trace amount of impurity in the GC composite. Further, it possesses reflections at 26.63° and 43.01° associated with rGO (JCPDS file no.: 01-075-2078). The GPGC exhibits peak positions at 19.00° and 36.68° corresponding to the Co(OH)₂ phase. It also possesses a broader peak covering the region corresponding to rGO at $\sim 26.6^{\circ}$ and another peak at 44.30° corresponding to the (101) plane. Both GC and GPGC do not

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possess peaks corresponding to the planes of (003) and (006), indicating the absence of an α -Co(OH)₂ phase. But both possess peaks corresponding to the (001) plane, and the *d*-spacing corresponding to this plane is 4.6 Å, affirming the presence of β -Co(OH)₂.¹⁸ The broad peaks of rGO located at ~26.6° in both the GC and GPGC XRD patterns indicate the loose packing of rGO layers and effective Co(OH)₂ intercalation between them.¹⁹ The crystallite size of Co(OH)₂ in GC calculated using the most intense peak corresponding to Co(OH)₂ is 25.65 nm (eqn (S1)†), and the crystallite size of GPGC was left undetermined as the peaks are convoluted and broader. The number of rGO layers per domain (eqn (S2)†) in GC is determined to be 44, indicating the multilayered nature of rGO in it.

3.1.2 FT-IR spectra. The FT-IR spectra of both GC and GPGC (Fig. 1b) possess the characteristic peaks of $Co(OH)_2$ and rGO, indicating the successful synthesis of GC and GPGC. The transmittance peaks exhibited by GC between 409 and 490 cm⁻¹ and at 1228 cm⁻¹ correspond to the vibration bands of the Co-O bond in Co(OH)₂.^{20,21} The peaks at 1365, 1568, and 1737 cm⁻¹ correspond to the deformations of -OH/C-O-C,²¹ C=C bonds of sp² hybridized carbon,²² and the C=O vibrations of the carboxylic acid groups²¹ present on GC, respectively. The peak at 3630 cm⁻¹ pertains to the β -form of Co(OH)₂.²³ GPGC possesses a series of transmittances from 421 to 668 $\rm cm^{-1}$ corresponding to the different vibrational modes of bonds that exist in Co(OH)₂. In that range of transmittances, the peaks at 513, 571-583, 634, and 668 cm⁻¹ correspond to Co-OH tensile/Co-O vibrations,^{22,24} vibrations of Co-O bond of β-Co(OH)₂,²⁰ Co-OH flexural vibrations,25 and vibrations of Co-O-H bonds,26 respectively. The transmittances at 1254, 1351, and 1570 cm⁻¹ are due to the vibrations of C-O stretching,²¹ O-H,²⁵ and sp² hybridized C=C bonds,²² respectively.

3.1.3 XPS analyses. The elements present in GC and GPGC were confirmed to be C, O, and Co via XPS studies (Fig. 1c). The deconvolution of the core level spectra of GC is as follows. The deconvolution of C 1s core level spectra (Fig. 1d) provided three peaks corresponding to the C=C, C-OH/C-O,19 and C=O27 bonds at 284.60, 286.69, and 289.08 eV, respectively. The deconvolution of O 1s (Fig. 1e) provided four peaks at 531.21, 532.94, 529.82, and 535.61 eV corresponding to Co-OH,²⁸ C= O,²⁹ OH of Co(OOH)³⁰ and OH of Co(OH)₂,²⁸ respectively. The Co 2p (Fig. 1f) spectra provided four peaks corresponding to the spin-orbit coupling of 2p_{3/2} and two peaks corresponding to $2p_{1/2}$. The peaks at 780.97 eV are associated with Co $p_{3/2}$, and the peaks observed at 787.17 and 790.18 eV are corresponding to the satellite peaks.³¹ The peak at 784.41 eV indicates the presence of Co(OOH) along with Co(OH)2. The peaks corresponding to Co $2p_{1/2}$ and its satellite peak are at 797.59 and 801.70 eV, respectively.32,33 The doublet peak separation by 16.62 eV confirms the presence of Co(OH)₂.³⁴ Co 2p_{3/2} has two satellite peaks, one with a difference of 6.2 eV and another with 9.21 eV,^{25,35} indicating the presence of both Co^{2+} (Co(OH)₂) and Co^{3+} (Co(OOH) in GC), respectively. GPGC also exhibited similar peaks upon deconvolution, and its deconvoluted core level spectra of C 1s, O 1s, and Co 2p are shown in Fig. S1.†

3.1.4 FE-SEM analyses. The FE-SEM images of GC at different magnifications (Fig. 2 and S2[†]) clearly depict that the

flake-like structured Co(OH)₂ are dispersed completely across the bulk of the rGO layers. The nano-pores present in the GC are distinctly seen in Fig. 2c and d. These nano-pores facilitate the permeation of electrolytes deep into the GC composite to bring about the maximum occurrence of electrochemical reactions (both faradaic and non-faradaic) and in turn, maximum energy storage. The molten rock particles-like deposition on the surface of GC (Fig. 2c and d) is NaOH, whose incorporation occurred during the synthesis of GC while enhancing the pH of the reaction content to 14 using 6 M NaOH, and this lower percentage of NaOH is the leftover after a series of washings of GC using DW and ethanol. This leftover NaOH on the GC surface is anticipated to work in unison with the aqueous electrolytes used to obtain higher energy storage. The elements of GC are duly confirmed by the energy dispersive spectrum (EDS) to be C, O, Co, and Na. The weight and atomic percentages of these elements are shown in Table S1.[†] The elemental maps of GC (Fig. S4[†]) signify the uniform distribution of its elements across the bulk.

3.1.5 Transmission electron microscopy (TEM) and scanning transmission electron microscopy (STEM) analyses. The TEM images further substantiate the results of FE-SEM by clearly depicting the Co(OH)2 dispersed rGO layers of GC (Fig. 3a and b). Co(OH)₂ possesses a plate-like shape with a hexagonal structure, which confirms the presence of β -Co(OH)₂ (hexagonal brucite structure)³⁶ (Fig. 3c and d). The STEM image (Fig. 3e) further substantiates the hexagonal structure of Co(OH)₂ present in the GC. The SAED (selected area electron diffraction) pattern of GC possesses concentric circles with highly intense bright and low intense dull spots, indicating the presence of both amorphous and polycrystalline phases in GC. The circles corresponding to different planes of β -Co(OH)₂ and rGO are determined from the SAED patterns and are shown in Fig. 3f. The elemental maps obtained from EDS attached with the TEM instrument further confirm the uniform dispersion of C, O, and Co across the bulk of the GC (Fig. S5[†]).

3.1.6 BET analysis. The N₂ adsorption isotherms of GC150 and GPGC150 (Fig. 4a) possess type IV isotherms with a hysteresis loop of the H3 kind, signifying that both of them are populated with meso-pores.³⁷ The BET surface area exhibited by GC150 and GPGC150 are 86.94 and 36.48 m² g⁻¹, respectively. The cumulative pore volume of GC150 and GPGC150 (Fig. 4b) is 0.181 and 0.076 m³ g⁻¹, respectively, and the cumulative surface area (Fig. 4c) exhibited by these pores is 27.75 and 19.26 m² g⁻¹, respectively. The higher surface area and the pore volume of GC150 bring about higher magnitudes of electrochemical reactions and higher energy storage in comparison with GPGC150.

3.2. Electrochemical characterization

The influence of $Co(OH)_2$ in enhancing the energy storage GC composite was studied by maintaining the concentration of rGO constant and varying the quantity of $Co(OH)_2$. There were four composites synthesized in the series, and they had a constant quantity of rGO as 25 mg and variable quantities of $Co(OH)_2$ as 50, 100, 150, and 200 mg. The composites were named as GC50,



Fig. 1 (a) X-ray diffraction (XRD) patterns, (b) FT-IR spectra of GC and GPGC, (c) survey spectra of GC and GPGC, core level spectra of (d) C 1s, (e) O 1s, and (f) Co 2p of GC.

GC100, GC150, and GC200, respectively, in accordance with their $Co(OH)_2$ concentration. The energy storage performance evaluation was conducted using 1 M H₂SO₄ as the electrolyte and at the potential range of 0–1.2 V in a symmetric two-electrode configuration. The results revealed that there was an increase in the energy storage of GC composites with an

increase in the quantity of $Co(OH)_2$ until 150 mg, beyond which there was a decrease in the energy storage. Therefore, a composite with a weight ratio similar to that of GC150 was synthesized using pristine graphite as the source of GO and in turn rGO, and named as GPGC150. It was observed that the supercapacitance of GC150 was higher than that of the



Fig. 2 (a-d) The FE-SEM images of GC at different magnifications.

GPGC150, indicating the fruitfulness of using graphite recovered from the spent Li-ion batteries (LIBs) as a source for synthesizing rGO, specifically for the synthesis of $rGO/Co(OH)_2$ composites, following the synthetic route mentioned in Section 2.1.

3.2.1 CV studies. The CV curves of all the GC composites (Fig. 5a) are quasi-rectangular, indicating the simultaneous roles of both the faradaic and non-faradaic energy storage processes. The non-faradaic energy storage process involves the formation of electrical double layers (EDLs) at the interfaces of the 1 M H₂SO₄ (electrolyte) and GC (electrode), electrolyte and separator (Whatman filter paper) and electrolyte and current collector (stainless steel plates) while applying the potential. Among these interfaces, the electrode and electrolyte interface cause higher EDL capacitance as the surface area available on the surface of the electrode is larger than that available on the separator and current collector. The faradaic energy storage processes are the possible redox reactions that $Co(OH)_2$ and Co(OOH) undergo with electrolytes and the intercalation of electrolytic ions in and out of the layered structures of the GC. The area under the CV curves is the measure of energy stored, which increases with an increase in the concentration of $Co(OH)_2$ until 150 mg, beyond which the CV area decreases, indicating a reduction in energy storage and the optimum Co(OH)₂ load on 25 mg of rGO as 150 mg. The area inside the CV curves of GC50, GC100, GC150, and GC200 were calculated to be

2.35, 2.58, 3.77, and 2.3 mA V, respectively, indicating the maximum energy stored by GC150. The initial increase in energy is because of the added Co(OH)₂ disperses itself on the rGO, uniformly increasing the space for the EDLC, redox, and intercalation phenomena, and these phenomena bring about more energy storage. The later decrease in energy storage is due to the reason that the excessive addition of $Co(OH)_2$ causes thick aggregates of $Co(OH)_2$ on the surface of rGO, hampering the electrolytic access into the entire bulk of the GC, increasing the internal resistance of the GC, and in turn decreasing the overall energy storage.19 The CV curve of GC150 is different in shape, which is highly deviated from the rectangular behaviour because of the occurrence of faradaic processes causing energy storage. The CV curves of GC150 at different scan rates are seen to have good reversibility up to a scan rate of 100 mV s^{-1} (Fig. 5b), indicating the promising reversibility of the electrode reactions even at high potential loads.

3.2.2 GCD studies. The quasi-triangular shape of the GCD curves (Fig. 5c) is due to the simultaneous action of both faradaic and non-faradaic energy storage reactions. The quantity of energy stored is proportional to the discharge time (t_d), which is 19, 22, 33, and 20 seconds for GC50, GC100, GC150, and GC200, respectively. The high t_d of GC150 indicates its higher energy storage capability. The energy storage parameters like specific capacitance (eqn (S4)†), specific capacity (eqn (S5)†), specific energy (eqn (S6)†), specific power (eqn (S7)†) and coulombic





Fig. 3 (a-d) TEM images of GC at different magnifications, (e) STEM image, and (f) SAED pattern of GC.

efficiency (eqn (S8)†) of GC composites are presented in Table 1. The *IR* drops observed at a current density of 1 A g⁻¹ are 0.075, 0.023, 0.022, and 0.035 V for GC50, GC100, GC150, and GC200, respectively. The decrease in the *IR* drop with the increase in Co(OH)₂ content indicates that adding Co(OH)₂ up to 150 mg increases the energy storage, and the least *IR* drop of GC150 is reflected in its higher C_s . The discussion on the increase in C_s with the increase in concentration of Co(OH)₂ up to 150 mg and the decrease in C_s beyond 150 mg is presented in Section 3.2.1. The maximum C_s , Q, E, P, and η exhibited by GC150 at a current density of 1 A g^{-1} are 36 F g^{-1} , 43 C g^{-1} , 7.16 W h k g^{-1} , 0.77 kW k g^{-1} and 94.99%, respectively.

The sustainability of higher applied current, storage, and delivery of high energy at these maximum applied currents are seen in the form of the rate capabilities of GC composites. The maximum tolerable current density exhibited by GC50, GC100, GC150, and GC200 are found to be 10, 15, 21, and 13 A g^{-1} , respectively. The high tolerable current density of GC150 indicates its robustness and faster mobility of electrolytic ions into the bulk of the GC150, which is reversible even at a high current



Fig. 4 (a) N₂ adsorption and desorption plots, (b) plots of cumulative pore volume vs. pore radius, and (c) cumulative surface area vs. pore radius of GC150 and GCPG150.

density (Fig. 5d, e, and 6a). The higher rate capability of GC150 in comparison with other GC composites indicates the faster kinetic of electrolytic ions and electrochemical reactions that are taking place in it even at higher applied currents in comparison with other GC composites.

The maximum E and P exhibited by GC50, GC100, GC150, and GC200 are 4.13 W h $\rm kg^{-1}$ and 0.77 kW $\rm kg^{-1};$ 4.30 W h $\rm kg^{-1}$ and 0.70 kW kg⁻¹; 7.16 W h kg⁻¹ and 0.77 kW kg⁻¹; and 4.20 W h kg⁻¹ and 0.75 kW kg⁻¹, respectively (Fig. 6b). The maintenance E with respect to increase in current density follows the same orders as that of their C_s order. All the GC composites exhibit the η close to 100% at all current densities, indicating the reversibility of their electrochemical reactions at all current densities (Fig. 6c). Considering the significance of the *b*-values with regard to the energy storage processes taking place in an electrode and its magnitude (eqn (S9)[†]), it is clear that all the GC composites other than GC150 (Fig. 6d) are involved in the intercalation mechanism at lower and higher potentials and majorly EDL formation at mid potential ranges. Meanwhile, GC150 involves the simultaneous operations of both mechanisms in the entire potential range, giving rise to higher energy storage than that of other GC composites. In order to comprehend the inference mentioned above, it is imperative to consider that the diffusion-controlled intercalation processes bring about higher energy storage in comparison with surface phenomena like EDL formation and redox reactions. In addition, this higher occurrence of intercalation

processes has produced a slightly different CV curve in GC150 in relation to other GC composites, as seen in Fig. 5a. Since GC150 has exhibited superior energy storage performance among the GC composites, a composite with a similar weight composition as that of GC150 was synthesized using pristine graphite as a source for GO and, in turn, rGO, whose energy storage performance was evaluated and compared with that of GC150. The CV area (Fig. 6e) and the t_d (Fig. 6f) exhibited by GPGC150 is lower than that of GC150, indicating its lower energy storage parameters of GPGC150 are presented in Table 2, indicating its low energy characteristics in comparison with GC150. The maximum C_s , Q, E, P, and η exhibited by GPGC150 are 15 F g⁻¹, 18 C g⁻¹, 3.06 W h kg⁻¹, 0.70 kW kg⁻¹, and 97.51%, respectively.

The maximum tolerable current density of GPGC150 is 16 A g⁻¹, at which it maintains 53.33% (8 F g⁻¹) of its initial C_s (15 F g⁻¹) (Fig. 6g). It should be noted that the rate capability exhibited by GPGC150 is superior to GC150, indicating the occurrence of an electrochemical reaction in a similar fashion as that of lower current densities even at higher current densities, which is not the case with GC150. The retention of *E* with an increase in current density follows a similar order to that of $C_s vs.$ current density. The plots of *E vs. P* depict the comparison between *E* and *P* of GC150 and GPGC150 (Fig. 6h). The η of GPGC150 is slightly higher than GC150 at lower current densities and is closer to 100% at its mid and higher current densities, indicating the higher reversibility of electrochemical



Fig. 5 Cyclic voltammetry (CV) curves of (a) different GC composites at a scan rate of 5 mV s⁻¹, (b) GC150 at different scan rates; galvanostatic charge and discharge (GCD) curves of (c) GC composites, (d) GC150 from the current density of 1 to 10 A g^{-1} , and (e) GC150 from the current density of 11 to 21 A g^{-1} .

Table 1	The energy storage parameters of GC composites at a current
density	of 1 A g ⁻¹

Composite	$C_{\rm s} \left({\rm F g}^{-1} \right)$	Q (C g ⁻¹)	E (W h kg ⁻¹)	P (kW kg ⁻¹)	η (%)
GC50	21	25	4.13	0.77	98.67
GC100	21	26	4.30	0.70	99.28
GC150	36	43	7.16	0.77	94.99
GC200	21	25	4.20	0.75	97.58

reactions occurring in GPGC150 in comparison with GC150 (Table 2 and Fig. 7a). The plots of *b*-value *vs.* potential (Fig. 7b)

indicate that GPGC150 stores energy majorly by surface phenomena and to a minimum extent by the intercalation phenomena and thus it exhibits lower energy storage and better rate capability than GC150 as the surface phenomenon like EDLC is highly reversible compared to the intercalation reactions. As seen from Fig. 7b, GC150 and GPGC150 store energy not only by surface capacitive processes but also by diffusioncontrolled intercalation processes. The kinetics of intercalation processes are lower than the surface energy storage processes like electrical double layer formation and pseudo surface reactions, and therefore the lower coulombic efficiencies in the range ~95% are attained. The cyclic stability study conducted at a current density of 10 A g⁻¹ for 10 000 continuous This article is licensed under a Creative Commons Attribution-NonCommercial 3.0 Unported Licence.

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Fig. 6 Plots of (a) specific capacitance vs. current density, (b) specific energy vs. specific power, (c) coulombic efficiency vs. current density, (d) *b*-values vs. potential, (e) CV curves at a scan rate of 5 mV s⁻¹, (f) CD curves at a current density of 1 A g⁻¹, (g) plots of specific capacitance vs. current densities, and (h) plots of specific energy vs. specific power of GC150 and GPGC150.

charge and discharge cycles (Fig. 7c) depicts that there is a deterioration in C_s up to 4800 cycles and beyond, where a constant increase in C_s up to 10 000 cycles is seen in the case of GC150. At the end of the 10 000 cycles, GC150 exhibits 12.75% higher C_s than that of the initial capacitance, whereas GPGC150 exhibits a gradual deterioration and maintains 90.77% of its initial $C_{\rm s}$. The observed improvement in the specific capacitance after 5000 cycles in the case of GC150 is because the continuous electrolytic insertion and extraction increase the extent of the intercalation processes, increasing the

Table 2 The energy storage parameters of GC150 and GPGC150 at a current density of 1 A $\rm g^{-1}$

Composite	$C_{\rm s} \left({\rm F g}^{-1} \right)$	Q (C g ⁻¹)	E (W h kg ⁻¹)	P (kW kg ⁻¹)	η (%)
GC150	36	43	7.16	0.77	94.99
GPGC150	15	18	3.06	0.70	97.51

specific capacitance after 5000 cycles. This is confirmed by the *b*-values obtained after 10 000 cycles of continuous charge discharge. In addition, it is clear that the peak created at the lower potential region is due to the intercalation process as the

b-value corresponds to peak potential closer to 0.5, as seen in Fig. S6.† The Nyquist plots obtained after 10 000 cycles for GC150 (Fig. 7d) and GPGC150 (Fig. 7e) depict that in both cases, there is an increase in solution resistance (R_s) ,³⁸ a decrease in charge transfer resistance $(R_{ct})^{38}$ and decrease in Warburg resistance.³⁹ The decrease in R_{ct} and the Warburg resistance (*W*) resulted in appreciable cyclic stability despite the increase in R_s . Therefore, the detrimental effect caused by the increase in R_s to GC150 and GPGC150 is possibly compensated by the decrease in R_{ct} and W, and hence, there are negligible and minimal performance deteriorations in the cases of GC150 and GPGC150, respectively, at the end of the 10 000 cycles. The



Fig. 7 Plots of (a) coulombic efficiency vs. current density, (b) *b*-values vs. potential, (c) %retention of C_s vs. number of cycles obtained at a current density of 10 A g⁻¹, (d) Nyquist plots of GC150 before and after 10 000 cycles, (e) Nyquist plots of GPGC150 before and after 10 000 cycles, and (f) coulombic efficiency vs. no. of cycles obtained at a current density of 10 A g⁻¹ of GC150 and GPGC150.



Fig. 8 (a) Nyquist plots, plots of (b) series resistance vs. log frequency, and (c) capacitance vs. log frequency of GC150 and GPGC150.

coulombic efficiency (Fig. 7f) of both GC150 and GPGC150 is closer to 100% during the cyclic stability study until 10 000 cycles, indicating the intact faster kinetics of the energy storage processes.

The Nyquist plots of GC150 and GPGC150 possess a semicircle at a higher frequency and a linear Warburg portion at a lower frequency, indicating their capacitive nature (Fig. 8a). The various electrical elements involved in the energy storage processes of GC composites were deduced by fitting the obtained Nyquist plots to an equivalent circuit of $R_s(R_{ct}Q_1)((R_{leak}C_{dl})(Q_2))W$, and the obtained values are provided in Table S2 (Fig. S7).† The descriptions for electrical elements of equivalent circuits are the same as they are provided in ref. 19 and also provided in ESI (Section S4).† As it is seen in the table, the sums of R_s , R_{ct} , R_{leak} , and W are 2.59, 0.78, 0.66, and 8.26 Ω for GC50, GC100, GC150, and GC200, respectively. The lower value of GC150 indicates its higher energy storage character than the other GC composites. The cumulative of R_s , R_{ct} , and W, are equivalent to the real axis of the Nyquist plot, is called as series resistance (Fig. 8b). The series resistances exhibited by GC150 and GPGC150 are 14.54 Ω and 30.48 Ω , respectively; this low series resistance of GC150 resulted in its high capacitance of 0.24 F (eqn (S9)†) at a lower frequency of 0.01 Hz (Fig. 8c), which is higher than the lower capacitance (0.13 F) exhibited by GPGC150 due to its higher series resistance under similar circumstances. The Nyquist plots of all the GC composites

Electrode	E Electrolyte (V)	$C_{ m s}~({ m F~g^{-1}})$	E (W h kg ⁻¹)	P (kW kg ⁻¹)	Cyclic stability	Ref.
2-EL						
rGO/β-Co(OH) ₂ (GC150)	1 M H ₂ SO ₄ 1.2	36 at 1 A g^{-1}	7.16	0.77	No deterioration up to 10 000 cycles at 10 A g^{-1}	Present work
3-EL						
α-Co(OH) ₂ /rGO/carbon fibre paper	1 M NaOH 0.5	1093 at 1.8 A g^{-1}	—	—	2% deterioration up to 5000 cycles at 1.8 A g^{-1}	34
rGO/Co(OH) ₂	2 M KOH 0.5	1100 at 0.5 A g^{-1}	—	—	1.9% deterioration up to 2000 cycles at 5 A g^{-1}	41
rGO/Co(OH) ₂	2 M KOH 0.5	474 at 1 A g^{-1}	—	—	10% deterioration up to 1000 cycles at 3 A g^{-1}	26
rGO/Co(OH) ₂	2 M KOH 0.8	734 at 1 A g^{-1}	—	—	5% deterioration up to 1000 cycles at 1 A g^{-1}	23

Table 3 Comparison of the supercapacitance of rGO/Co(OH)₂ obtained in the two-electrode system with those of the reported ones

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possess semicircles in the higher frequency region and linear Warburg lines in the lower frequency region (Fig. S8a[†]), indicating their capacitive nature. The series resistance is proportional to the real axis of the Nyquist plots and decreases in its magnitude with an increase in $Co(OH)_2$ concentration until the loading of 150 mg and beyond, which it increases (Fig. S8b[†]). The GC corresponding to the weight of 150 mg of $Co(OH)_2$ is GC150, which possesses the lowest series resistance at the lowest frequency of 0.01 Hz. This lower series resistance of GC150 is responsible for its high capacitance of 0.24 F (eqn (S10)[†]) at 0.01 Hz (Fig. S8c[†]). The supercapacitance of GC150 obtained is compared with similar reported rGO/Co(OH)₂ composites, and the comparison is presented in Table 3. The rGOs presented in Table 3 were obtained from pristine graphite, except in the present study. The comparison between the two and three electrodes is possible, considering that the performance obtained by the three-electrode system is four-fold higher than that obtained by the two-electrode system.⁴⁰

4 Conclusions

The supercapacitance studies were carried out on two composites of rGO/ β -Co(OH)₂ with the same weight composition, with the difference among them being the graphite sources, one being the graphite recovered from spent Li-ion batteries (GC150 (rGO_{25mg}/β-Co(OH)_{2 150mg})) and the other being pristine graphite (GPGC150). The study revealed that GC150 exhibits superior energy storage performances compared to GPGC150. However, the rate capability of GPGC150 is found to be higher than that for GC150. Both the composites exhibited better cyclic stabilities up to 10,000 cycles at a current density of 10 A g⁻¹. GC150 exhibited a zero deterioration, while GPGC150 exhibited 9.23% of deterioration. GC150 provided higher tolerable current density up to 21 A g^{-1} . The energy storage parameters, viz., specific capacitance, specific capacity, specific energy, specific power, and coulombic efficiency, provided by GC150 at 1 A g^{-1} were 36 F g^{-1} , 43 C g^{-1} , 7.16 W h kg⁻¹, 0.77 kW kg⁻¹, and 94.99%, respectively. The rGO/β-Co(OH)₂ composites synthesized using recovered graphite as the source for rGO exhibit supercapacitance better than their analog that is synthesized using pristine graphite as the source for rGO by the synthetic route followed in the present study. Therefore, using graphite recovered from the spent Li-ion batteries to make electrodes for supercapacitors could be a wise notion in attaining sustainable energy materials for the future.

Conflicts of interest

There are no conflicts to declare.

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