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## Recovery of graphite from industrial lithium-ion battery black mass†

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The escalating production of commercial lithium-ion batteries (LIBs) is anticipated to result in a substantial accumulation of waste upon end-of-life disposal of LIBs, which however also represents a secondary source of raw materials. Among the components of LIBs, graphite anode is a critical material and its production via high-temperature carbonisation is highly energy- and cost-intensive. One of the major challenges regarding recycling of graphite materials from spent LIBs is the presence of residual metal and organic species that are difficult to eliminate, preventing direct reuse as anodes. Here, we propose a recycling workflow to eliminate the various impurities and regenerate the graphite materials from industrially sourced black mass, composed of mixed cathode materials, anode materials, aluminium and copper current collectors, Li salts, and polyvinylidene fluoride binders. After selective extraction of high-value transition metal ions, such as Li, Ni, and Co, from the black mass, the proposed workflow for graphite recovery involves a second step of acid leaching for the removal of Al, Cu, and other residual metal species, and mild-temperature pyrolysis for the removal of polyvinylidene fluoride (PVDF). The regenerated graphite (AG-2.0M-800) demonstrates an initial specific charge capacity of 387.44 mA h g<sup>-1</sup> at 0.1C (35 mA g<sup>-1</sup>) in lithium half cells, on par with commercial battery-grade graphite. This workflow provides a promising approach to the recycling of spent graphite that could be integrated with existing cathode materials' recycling processes developed in the industry.

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### Sustainability spotlight

The rise of electric vehicles has led to increased production of lithium-ion batteries (LIBs), presenting significant environmental challenges and raw material shortages due to end-of-life battery waste. Graphite recycling is often neglected because of the complexity and cost associated with impurity removal. Our study seeks to enhance the sustainability of the LIB supply chain by reducing the need for energy-intensive graphite production, which typically requires carbonising raw materials at over 2800 °C. We explored various recycling methods to recover graphite from industrial LIB black mass, with a focus on minimising energy costs. This comprehensive recycling strategy addresses a critical gap in the LIB recycling industry and offers a scalable solution adaptable to different compositions of black mass. Integrating this process with existing cathode recycling techniques would result in a comprehensive and eco-friendly solution. This work directly supports the UN Sustainable Development Goals: affordable and clean energy (SDG 7), industry, innovation, and infrastructure (SDG 9), and responsible consumption and production (SDG 12).

## Introduction

In the global transition to net-zero carbon emissions, the electric vehicle revolution is poised to transform the automotive industries,<sup>1</sup> driving the global lithium-ion battery (LIB) market to increase tenfold by 2030.<sup>2</sup> Consequently, the continuing accumulation of end-of-life LIBs poses a substantial safety and environmental risk arising from the flammable and hazardous

organic substances and the inevitable leaching of heavy metals upon improper disposal.<sup>3,4</sup> However, spent LIBs represent a valuable secondary source of strategic elements and critical materials, reducing the environmental footprint and improving the sustainability of the LIB supply chain.<sup>5,6</sup> Currently, the recycling of LIBs mainly focuses on cathode materials to recover high-value critical materials such as Co, Ni, and Li.<sup>7</sup> The recovery of graphite anodes, however, is overlooked and they are subjected to disposal after separating cathode materials from the black mass, mainly due to the difficulty and complexity in the complete removal of residual impurities.<sup>8</sup> Graphite anodes account for approximately 20% of the total weight in a battery pack, and the production of graphite by carbonising raw materials such as petroleum coke at temperatures above 2800 °C is energy intensive with the cost reaching up to \$15000 per ton.<sup>9</sup> Alongside economic benefits, regulatory policy and

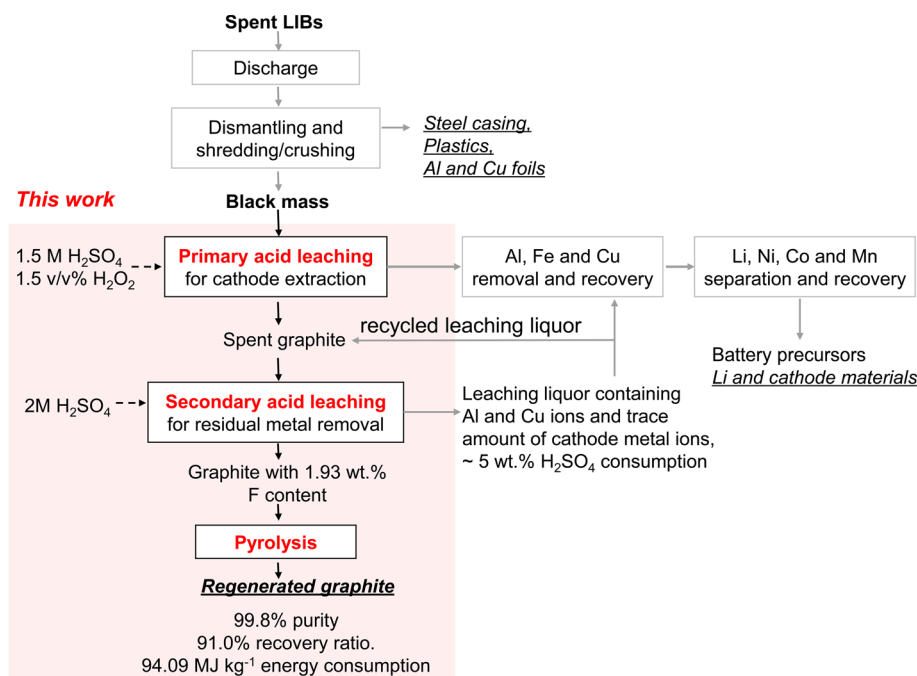
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**Scheme 1** Recycling of LIBs. The workflow for the graphite recovery proposed in this work is highlighted in the red-shaded area. Underscored are potential products to be recovered.

a strong push towards sustainability and circular economy are other drivers for developing economically viable recovery processes of spent graphite anodes.

Several recycling strategies have been reported to recover and regenerate graphite from spent LIBs. Yang *et al.* used citric acid as the extraction reagent to recover Li and regenerate spent graphite, demonstrating a discharge capacity of  $330 \text{ mA h g}^{-1}$  after 80 cycles at  $0.5\text{C}$  ( $1\text{C} = 372 \text{ mA g}^{-1}$ ).<sup>2</sup> Wang *et al.* developed a facile water-treatment method to remove residual Li and the solid electrolyte interphase (SEI) layer, utilising the hydrogen evolution reaction.<sup>10</sup> More recently, Chen *et al.* proposed a flash Joule heating process to decompose residual metal oxides and Li salts and regenerate spent graphite anodes.<sup>9</sup> Some alternative methods have also been proposed, such as the subcritical  $\text{CO}_2$ -assisted electrolyte extraction process<sup>11</sup> and the froth flotation method.<sup>12</sup> However, spent graphite anodes used in most of these studies were the anode coating layer that was manually disassembled and separated from spent LIBs with a surprisingly high initial purity of above 95 wt% before any treatment.<sup>2,10,13-15</sup>

In the LIB recycling industry, cathode and anode coating layers are recovered as a mixture, referred to as black mass, through the shredding/crushing process, which also brings other cell components such as copper and aluminium current collectors, PVDF binders, and Li salts to the black mass. It remains unclear whether the processes previously reported for graphite recycling would be effective for industrial black mass with a variety of impurities at high quantities. It is highly desirable to develop a graphite anode recycling process that can address this issue and that ideally can be integrated with the existing hydro-metallurgical process for cathode material recovery.<sup>16-18</sup> This process should also accommodate variations in the composition

of black mass, as different cell chemistries from different LIB manufacturers are often mixed together for recycling.

In this study, we employed industrially sourced black mass produced from a combination of three prevailing cell chemistries, namely, lithium manganese oxide (LMO), lithium nickel manganese cobalt oxide (NMC), and lithium iron phosphate (LFP). Graphite content in the black mass was ~40 wt%, while that of metal was ~40 wt% and phosphorus and fluorine ~20 wt%. We explored several methods including physical sieving, acid leaching, calcination, and pyrolysis, both individually and in combination, to provide a comprehensive analysis of their efficiency in removing various metal and organic impurities from spent graphite. Our findings indicated Al and F as primary sources affecting overall purification efficiency, while the remaining metal and phosphorus can be readily eliminated. By integrating acid-leaching with mild-temperature pyrolysis (Scheme 1), high-quality graphite was regenerated with a purity of 99.80% from ~40% initially. The regenerated graphite exhibited excellent rate performance and electrochemical stability, both comparable or superior to pristine battery-grade graphite.

## Experimental methods

### Pretreatment of industrial black mass *via* acid leaching

Black mass was provided in-kind by Alt lithium, UK. The black mass was leached with sulfuric acid ( $\text{H}_2\text{SO}_4$ ) or citric acid, with or without adding  $\text{H}_2\text{O}_2$ , to extract high-value metals (mainly those in cathode materials). After leaching, the solid residue was filtered, sieved, and dried. The resulting powder was designated as spent graphite. All leaching processes were conducted using a 1.5 M acid concentration with a solid-liquid



ratio of  $100 \text{ g L}^{-1}$  at  $90^\circ\text{C}$  for 40 minutes, with and without the addition of 1.5% (v/v)  $\text{H}_2\text{O}_2$  (30% w/w) as a reducing agent.

### Recycling of anode materials

For the combination of secondary acid leaching and air calcination, the obtained spent graphite was first subjected to calcination at  $500^\circ\text{C}$  for 1–3 hours in the air to oxidise Cu current collectors and remove binders. Subsequently, both calcined and non-calcined spent graphite samples were subjected to another step of leaching under identical conditions using 1–2 M  $\text{H}_2\text{SO}_4$ , with a solid-to-liquid ratio of  $50 \text{ g L}^{-1}$  at  $90^\circ\text{C}$  for 3 hours, to remove residual metal ions. The resulting acid-leached graphite was filtered, washed, and dried at  $120^\circ\text{C}$  for 12 hours.

For the combination of secondary acid leaching and  $\text{N}_2$  pyrolysis, acid leaching was performed before pyrolysis under the same conditions stated previously, to avoid the undesired graphite consumption due to reactions with the residual cathode at high temperatures. Pyrolysis duration was set at 6 hours for all experiments. After pyrolysis, the samples were washed with deionised water and dried at  $120^\circ\text{C}$  for 12 hours, yielding regenerated graphite.

### Structural characterisation

Graphite samples were subjected to microwave digestion (MARS 6, CEM) and digested solutions were analysed by inductively coupled plasma mass spectrometry (ICP-MS 7900, Agilent Technologies) to quantify the composition and percentage of metals in the samples. Thermogravimetry analysis (TGA) was conducted in a TGA800, PerkinElmer and programmed to heat the sample at a rate of  $10^\circ\text{C min}^{-1}$  up to  $1000^\circ\text{C}$  with a hold time of 45 min. The morphology of the samples was determined by scanning electron microscopy and energy dispersive spectrometry (SEM-EDS; JEOL 6010LA). X-ray diffraction analysis (XRD) was performed using an XRD Panalytical X-Pert with Cu  $\text{K}\alpha$  radiation. Raman spectroscopy was carried out using a Renishaw Raman spectrometer with a 532 nm laser. X-ray photoelectron spectroscopy (XPS) was performed using a ThermoFisher X-ray spectrometer with an Al Ka monochromated X-ray source.

### Electrochemical characterisation

Coin cell components (CR2032), and Li metal foil were purchased from Guangdong Canrd New Energy Technology Co., Ltd. The electrolyte 1.0 M  $\text{LiPF}_6$  in ethylene carbonate (EC) and dimethyl carbonate (DMC) electrolyte, sodium carboxymethyl cellulose binder (CMC,  $M_w \approx 250\,000$ ) and Whatman GF/A glass microfiber were purchased from Sigma Aldrich. All chemicals were used as received without further purification.

All the slurries including commercial graphite, spent graphite, acid-leached graphite, and regenerated graphite were prepared by mixing 90 wt% graphite powder with 10 wt% sodium carboxymethyl cellulose ( $M_w \approx 250\,000$ ) binder solution (5 wt% in deionized water). The slurries were well-mixed in a centrifugal mixer (Thinky, ARE-250 CE) for 10 min at 2000 rpm. The electrodes were then coated from slurries onto

battery-grade Cu foil (9  $\mu\text{m}$  in thickness). The coated electrodes were dried at room temperature for 6 hours followed by vacuum drying for 18 hours. The 1.0 cm circular electrodes were cut with mass loading ranging from 2.5 to 3.0  $\text{mg cm}^{-2}$ . Coin cells were then assembled inside a glovebox ( $\text{H}_2\text{O} < 0.5 \text{ ppm}$ ,  $\text{O}_2 < 0.5 \text{ ppm}$ ) against Li metal 12 mm in diameter. The Whatman GF/A glass microfiber separator was saturated with 100  $\mu\text{L}$  of 1 M  $\text{LiPF}_6$  (EC/DMC, 1:1 by vol). Three independent cells were tested to confirm the reproducibility.

## Results and discussion

### Separation of spent graphite from black mass

Due to the lack of material separation in the shredding/crushing process, black mass produced in the recycling industry normally contains significant amounts of Al and Cu current collectors and organic materials, alongside cathode and anode materials. We sourced black mass from the industry, and this black mass was produced mainly from LMO and NMC cell chemistries with a minor proportion of LFP. High-value metals, including Li, Co, Ni and Mn, constituted 26.7% of the total weight of the black mass, while Al and Cu current collectors contributed 12.8 wt%, as quantified by ICP-MS (Tables 1, S1 and S2†). The F content of 17.94%, as calculated from the XPS spectrum (Fig. S1†), can be attributed to PVDF binders and lithium hexafluorophosphate salt,  $\text{LiPF}_6$ .

Prior to graphite recovery, we conducted acid leaching to extract high-value metals from the black mass using  $\text{H}_2\text{SO}_4$  and organic citric acid (Fig. 1). This leaching process can be described as occurring in two distinct stages: an initial rapid phase, governed by solution pH and temperature, with limited influence from redox reactions, followed by a slower second stage driven by electrochemical processes and surface-controlled dissolution.<sup>19</sup> The dissolution of transition metal

Table 1 Composition of as-received black mass and spent graphite

	Composition (wt%)								
	Li	Co	Ni	Mn	Fe	P	Al	Cu	F
Black mass	2.53	1.78	6.43	15.95	0.57	1.34	4.79	8.08	17.94
Spent graphite	0.24	0.08	0.13	0.11	0.60	0.95	5.51	1.44	7.51

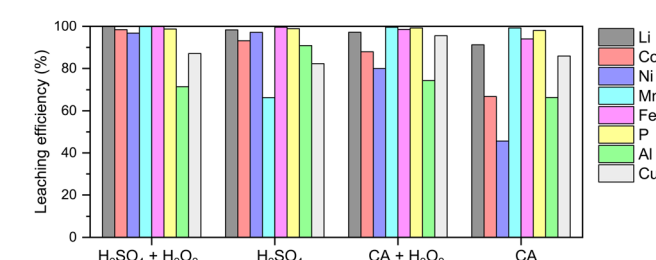


Fig. 1 Pretreatment of industrial black mass via acid leaching. This step was to extract high-value metals from the black mass and prepare spent graphite for subsequent recycling. Acid concentration: 1.5 M;  $\text{H}_2\text{O}_2$  concentration: 1.5 vol%. CA: citric acid.



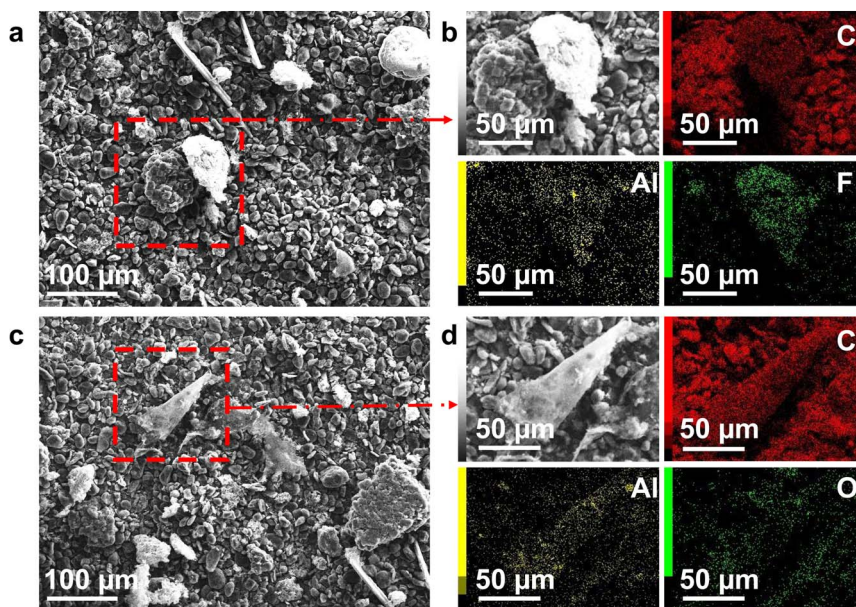


Fig. 2 SEM images of spent graphite anodes. (a and b) SEM-EDS mapping of Al foils detected in graphite. (c and d) SEM-EDS mapping of  $\text{Al}_2\text{O}_3$  particles detected in graphite.

oxides during the second stage is inherently a reduction reaction, and therefore requires the presence of a reductant. Fe, Cu and Al, which are already present in the raw material, could act as reductants due to their lower redox potential than transition metal oxides (Tables S3 and S4,† reaction mechanisms are described using eqn S1–S7†).<sup>20–23</sup> Nevertheless, excess dissolution of Cu and Al into the leach liquor significantly reduces the efficiency of subsequent steps for separating and recovering Co, Ni, and Mn (not explored in this work, Scheme 1).<sup>24</sup> Hence, we added a small volume of  $\text{H}_2\text{O}_2$  (1.5 vol%) as a competing reducing agent<sup>21,23</sup> (eqn S3†) and achieved the highest extraction efficiency of cathode materials, particularly Li, Co, and Ni, while decreasing the dissolution of Al and Cu, under the condition of 1.5 M  $\text{H}_2\text{SO}_4$  with 1.5 vol%  $\text{H}_2\text{O}_2$ .

Residual solids obtained after the optimised acid leaching were dried and sieved to remove large pieces of plastics and undissolved Al and Cu foils, yielding an underflow fraction of  $\leq 125\text{ }\mu\text{m}$ . The resulting solids, composed primarily of graphite, were designated as spent graphite for further experiments in this study. The composition and morphology of spent graphite were characterised using ICP-MS, XPS, and SEM. F, Al and Cu became the major impurities in spent graphite with a weight percentage of 7.51, 5.51, and 1.44 wt%, respectively, where Al existed in both metallic and oxide forms (Table 1 and Fig. 2). The weight percentage of Li, Co, Ni, and Mn was below 0.3 wt% (Table 1 and Fig. S2†), substantiating the efficacy of acid leaching regarding the extraction of cathode materials.

### Secondary acid leaching for residual metal removal

An additional step of acid leaching is required to remove residual metals from spent graphite. This serves a different purpose from the previous acid-leading step, which aims at the selective extraction of metals from cathode materials. Generally, acid

concentration, temperature, reaction time, and solid-to-liquid ratios are critical factors influencing the efficiency of metal removal. Among these parameters, we focused on the investigation of acid concentration, as it is the most important factor that has a direct impact on the dissolution rates.

Fig. 3 shows the weight percent of remaining impurities in spent graphite as a function of acid concentration, with samples designated as AG-X, where X represents  $\text{H}_2\text{SO}_4$  concentrations. Higher removal efficiencies were achieved for all impurities on increasing acid concentration, showing the lowest overall impurity content of 0.8 wt% in AG-2.0M, in contrast to 9.04 wt% before acid leaching. Metals from cathode materials (*i.e.*, Li, Co, Ni, and Mn) and Cu were eliminated nearly in full at a  $\text{H}_2\text{SO}_4$  concentration higher than 1.5 M, while Al, Fe and P required a concentration of 2 M. Al remained as a major impurity under all conditions with the predominance of  $\text{Al}_2\text{O}_3$  *versus* metallic Al as detected in SEM-EDS (Fig. S3†). The standard reduction potential ( $E_0$ ) of metallic Al ( $-1.66\text{ V}$  *versus* the standard hydrogen electrode) indicates a strong thermodynamic affinity for electron donation and oxidation and the redox reaction with protons readily produces  $\text{Al}^{3+}$  and hydrogen gas. In contrast, the chemical dissolution of  $\text{Al}_2\text{O}_3$  poses significant challenges due to the passivating layer that acts as a protective barrier, thereby hindering further leaching processes.<sup>25</sup> Therefore, the rate-limiting step of Al removal is not the electrochemical dissolution of Al to  $\text{Al}^{3+}$ , but rather the chemical dissolution of  $\text{Al}_2\text{O}_3$  to  $\text{Al}^{3+}$ . More aggressive leaching at higher acid concentrations may further reduce the content of metal impurities by breaking down the complex compounds. However, higher acid concentrations were not explored due to the safety considerations and higher costs associated with the procurement, storage, handling and disposal of concentrated acids, especially under the setting of potential industrial applications at large scale.



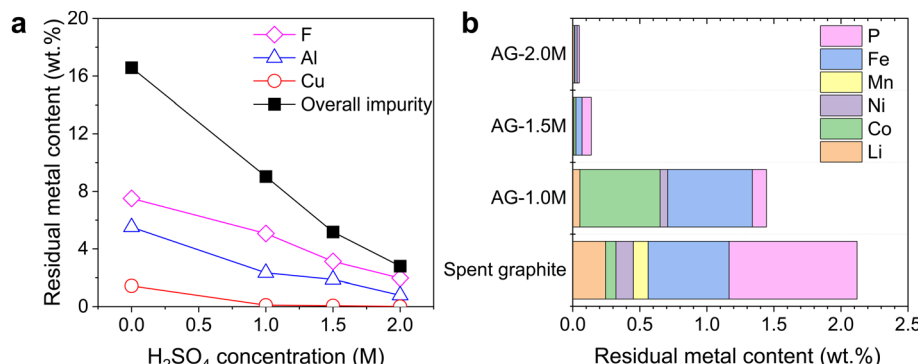


Fig. 3 Weight percentages of residual metals in spent graphite after secondary acid leaching. (a) Al, Cu, F and overall impurity contents as a function of acid concentration. (b) Other residual metal contents after leaching processes. AG-X samples, where X represents  $\text{H}_2\text{SO}_4$  concentration.

To underscore the necessity of employing a two-step leaching process, with primary leaching for cathode extraction and secondary leaching for residual metal removal, a control experiment of one-step leaching was conducted. Using a 2 M acid concentration, a solid-liquid ratio of  $50 \text{ g L}^{-1}$  at  $90^\circ\text{C}$ , and 1.5% (v/v)  $\text{H}_2\text{O}_2$  (30% w/w), the results from ICP-MS (Table S5†) revealed a residual impurity content of  $\sim 5$  wt% in all solids. This confirms that one-step leaching is inadequate for complete graphite purification. With an acid consumption of only  $\sim 5\%$  per cycle, the leaching liquor from the secondary acid leaching stage could be potentially recycled until the concentrations of metal ions reach sufficient levels (Scheme 1), enabling higher metal recovery and lower energy cost.

### Calcination and pyrolysis for PVDF removal

Previous studies on graphite recovery often overlook the challenge of removing PVDF binders, which are inevitably introduced into the black mass during the shredding and crushing process. While acid leaching is effective for metal removal, it proves inadequate for PVDF removal, leaving a fluorine content of 2.25 and 1.97 wt% in AG-1.5M and AG-2.0M, respectively (Fig. S4†). Air calcination is widely used in LIB cathode recycling to thermally decompose PVDF, which has an onset decomposition temperature of around  $380^\circ\text{C}$ .<sup>26–28</sup> As temperatures greater than  $600^\circ\text{C}$  lead to rapid combustion of graphite,<sup>15,28</sup> we chose an air calcination temperature of  $500^\circ\text{C}$ . After 1 h of air calcination,

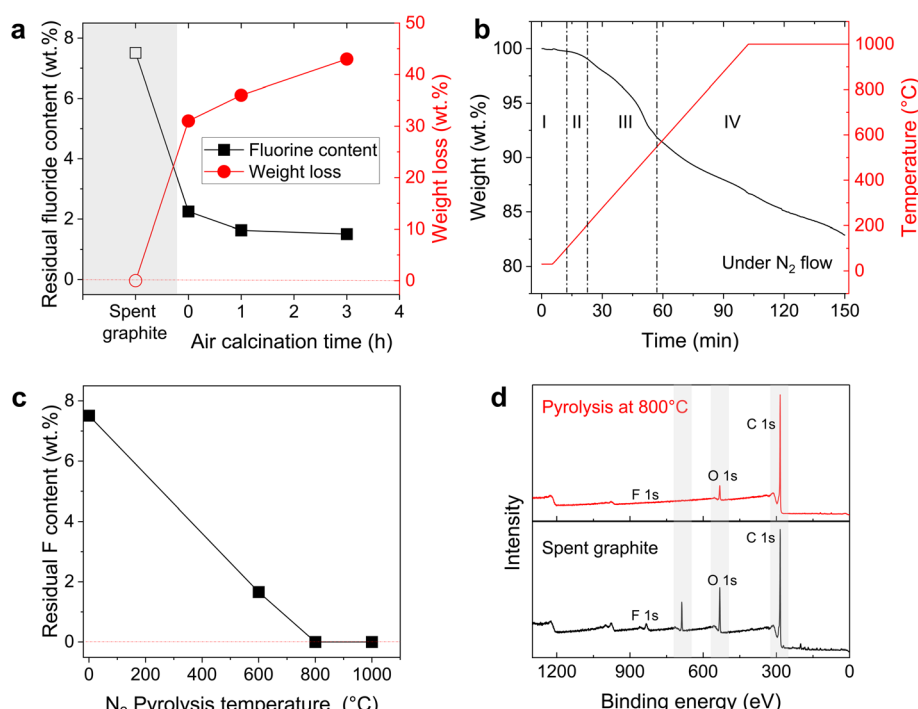


Fig. 4 Air calcination and  $\text{N}_2$  pyrolysis for fluorine removal. (a) Residual fluorine content and sample weight loss as a function of calcination time. The F content was measured after air calcination and secondary acid leaching, with the only variable being the calcination time, while data of the original spent graphite are added in the shaded area for comparison. (b) TGA curves of spent graphite under  $\text{N}_2$ . (c) Residual fluorine content as a function of  $\text{N}_2$  pyrolysis temperature. (d) XPS spectra of spent graphite sample and spent graphite after sintering at  $800^\circ\text{C}$  under  $\text{N}_2$ .



followed by secondary acid leaching using 1.5 M  $\text{H}_2\text{SO}_4$ , F content decreased to 1.51 wt% (Fig. 4a). Prolonging the calcination time to 3 h resulted in a further decreased F content of 1.38 wt% accompanied by an additional weight loss of 7 wt%. We attribute the weight loss to the incomplete combustion of graphite, which commences at temperatures above 400 °C and produces CO and  $\text{CO}_2$ .<sup>29</sup> While air calcination can decompose a certain amount of PVDF, the undesired graphite oxidation and combustion render this method less practical for graphite recovery (Fig. 4a). Furthermore, Fig. S5 and S6<sup>†</sup> indicate that calcination has limited efficacy on the removal of other impurities.

Pyrolysis, akin to calcination but under inert gas, can concurrently fulfil the requirements of thermal decomposition of PVDF and preservation of the structural integrity of graphite, while also providing opportunities to repair structural defects in spent graphite. Thermal gravimetric analysis (TGA) of black mass reveals distinct degradation stages during pyrolysis in Fig. 4b.<sup>30</sup> During the initial stages (I and II) below 200 °C, moisture evaporation and the decomposition of organic species occur. In stage III, PVDF undergoes rapid thermal decomposition between 450 and 550 °C, with maximum weight loss at 500 °C. Above 500 °C, decomposition releases various fluorinated compounds,<sup>31</sup> and also involves hydrogen fluoride release

and polymer chain scission.<sup>32</sup> At stage IV, the weight loss can be attributed to the reduction reaction of residue metal oxides with graphite, Al and F, producing simpler metal oxides, metals and LiF.<sup>33</sup> We note that pyrolysis in the presence of reducing agents (such as Al and graphite) is widely deployed in the pyrometallurgical process for LIB recycling to extract Cu, Ni, and Co as metal alloys.<sup>34</sup> These findings emphasise the necessity to eliminate residual metal oxides from the black mass prior to the thermal treatment, in order to prevent graphite loss and ensure the complete removal of fluorine.

To evaluate the efficiency of F removal, we performed pyrolysis at 600, 800, and 1000 °C under a  $\text{N}_2$  atmosphere (Fig. 4c, d, S7 and S8<sup>†</sup>). Complete F removal requires a pyrolysis temperature of at least 800 °C, while extending the temperature further doesn't necessarily improve the purity of graphite. Thus, 800 °C was identified as the optimal temperature for the pyrolysis process to eliminate F compounds and repair degraded graphite by eliminating oxygen groups from the graphite surface. After pyrolysis, graphite samples were washed, ground, and sieved, yielding an overall 91.0% recovery, accounting for both the secondary acid leaching and pyrolysis steps. The weight loss for AG-1.5M-800 and AG-2.0M-800 samples before and after sieving was negligible (~2 wt%).

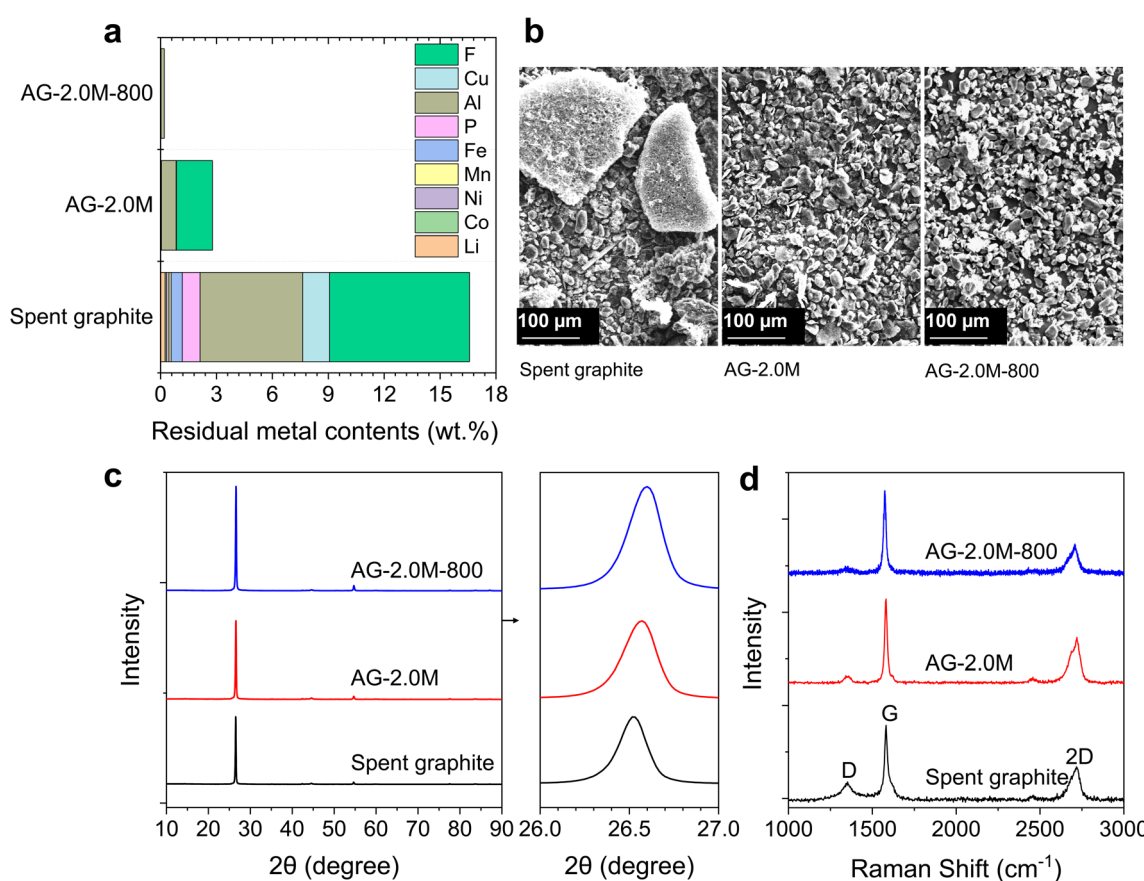


Fig. 5 Characterisation of recovered graphite. (a) Residual metal contents after optimized recycling processes. (b) SEM images. (c) XRD spectra. (d) Raman spectra. AG-2.0M: graphite recovered via 2 M  $\text{H}_2\text{SO}_4$  leaching. AG-2.0M-800: graphite recovered via 2 M  $\text{H}_2\text{SO}_4$  leaching followed by 800 °C pyrolysis.



## Characterisation of regenerated graphite

The regenerated samples (AG-1.5M-800 and AG-2.0M-800), acid-leached spent graphite (AG-1.5M and AG-2.0M) and the original spent graphite were thoroughly characterised to compare their composition, morphology, and microstructure. Chemical compositions are summarised in Fig. 5a and S8.† AG-2.0M-800 demonstrates a high graphite purity of 99.80 wt%. Despite Al being the predominant impurity across all samples, its concentration has markedly decreased from 5.51 wt% in spent graphite to less than 0.17 wt% in AG-2.0M-800. Large aggregates were found in spent graphite due to the presence of residual binders (Fig. 5b). The degree of agglomeration is significantly

reduced in AG-2.0M and AG-2.0M-800 (Fig. 5b and S9†), showing an average particle size below 30  $\mu\text{m}$  with a certain distribution. This explains the negligible loss of graphite after sieving (sieve size 45  $\mu\text{m}$ ). Graphite particle size is a critical factor in the processes of ink preparation and anode electrode manufacturing, with a preferred size in the range of 8 to 30  $\mu\text{m}$ , where our regenerated graphite falls within.<sup>35</sup>

The microstructure of these graphite samples was analysed using XRD, Raman spectroscopy and XPS as shown in Fig. S9–S12,† 5c, d and 6. All graphite samples, including spent graphite, exhibited a characteristic diffraction peak at  $2\theta = 26.5^\circ$ , corresponding to the (002) crystal plane, without any noticeable impurity peak. This suggests the structural stability

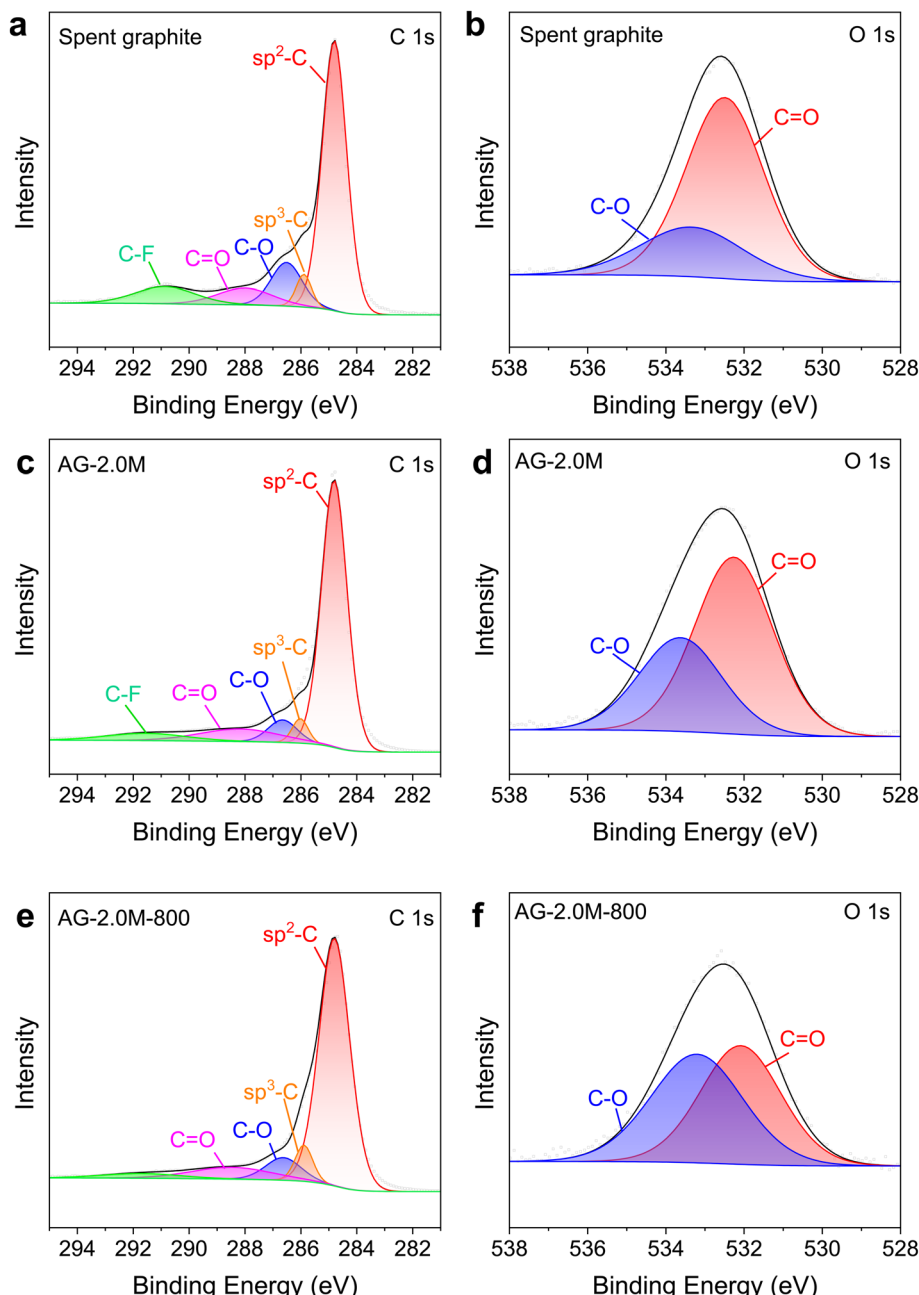


Fig. 6 High-magnification XPS spectra for C 1s and O 1s. (a and b) Spent graphite. (c and d) AG-2.0M. (e and f) AG-2.0M-800.



and resilience of graphite towards long-term cycling and recycling processes, laying the foundation of graphite recycling and reuse. The (002) peak shifted slightly from  $26.52^\circ$  for spent graphite to  $26.55^\circ$  for AG-2.0M and further  $26.56^\circ$  for AG-2.0M-800, equivalent to an interlayer distance ( $d_{002}$ ) of 0.3359, 0.3355, and 0.3350 nm, respectively, based on the Bragg equation  $2d \sin \theta = n\lambda$ . The larger  $d_{002}$  in spent graphite originates from the structural changes (and potential degradation) due to the repeated intercalation/deintercalation of Li ions during battery cycling.<sup>36</sup> The reduced  $d_{002}$  in AG-2.0M-800 highlights the added benefit of the pyrolysis step in restoring the interlayer spacing of spent graphite, in addition to F removal.

Three distinct characteristic peaks were observed in the Raman spectra of all samples at 1350, 1580, and 2720  $\text{cm}^{-1}$ , corresponding to D, G and 2D bands, respectively. The D band

is associated with defects and disorder in the graphite lattice, while the G band corresponds to the degree of graphitisation and crystallinity. Therefore, the ratio of the intensity of the D peak ( $I_D$ ) to that of the G peak ( $I_G$ ) represents the defect density in the graphite structure.<sup>9</sup> As expected, spent graphite exhibited the highest  $I_D/I_G$  ratio of 0.24 and AG-2.0M-800 the lowest of 0.09, in line with the results of XRD.

XPS spectra in Fig. S1 and S3<sup>†</sup> show that C and O are the primary components in all graphite samples, with characteristic peaks around 285 and 532 eV, respectively. According to the high-resolution XPS spectra (Fig. 6), the C 1s peaks can be deconvoluted into five components:  $\text{sp}^2$  C (284.8 eV),  $\text{sp}^3$  C (285.89 eV), C–O (286.51 eV), C=O (288.01 eV), and C–F or  $\pi-\pi^*$  transition (290.89 eV).<sup>37</sup> Of note, the component at 290.89 eV can be attributed to both C–F and electron excitation from the  $\pi$

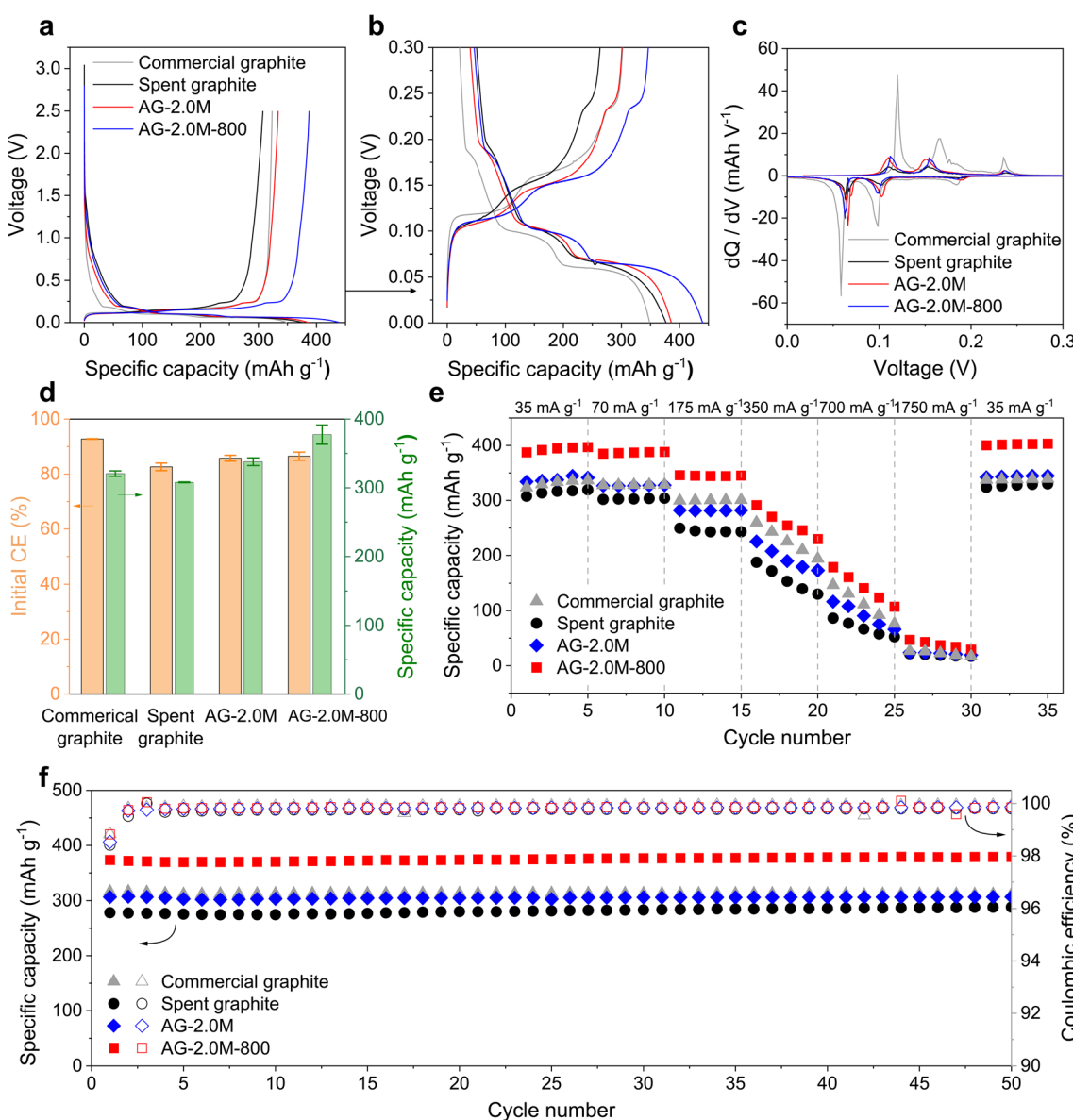


Fig. 7 Electrochemical performance of recovered graphite. (a and b) Charge–discharge profiles of the first cycle at 0.1C. (c)  $dQ/dV$  profiles. (d) Summary of the initial CEs and charge capacities. Error bars represent standard deviation based on three independent cells. (e) Rate performance. (f) Cycling performance in a half-cell configuration at 0.5C (175  $\text{mA g}^{-1}$ ).



bonding orbital to the  $\pi^*$  antibonding orbital, as documented in the literature and observed previously in pristine graphite.<sup>38–40</sup> The substantial reduction in the components of C–O, C=O, and C–F for AG-2.0M-800, relative to spent graphite and AG-2.0M, is in agreement with results from XRD and Raman spectra. The components at 532.49 and 533.33 eV in O 1s spectra correspond to C=O and C–O bonds, respectively. Peak areas for C–O and C=O are the highest in spent graphite among all the samples, particularly for the C=O peak, indicating a high concentration of oxygen-containing functional groups in spent graphite. Analysis of C 1s and O 1s in XPS spectra confirms that pyrolysis under N<sub>2</sub> can effectively remove F and oxygen elements from the graphite materials.<sup>41,42</sup>

**Electrochemical performance.** To evaluate the performance of regenerated graphite at the device level, we assembled half-cells with graphite as the cathode and Li metal as the anode with voltage cutoffs of 0 and 2.5 V *vs.* Li<sup>+</sup>/Li. Due to the difficulty in obtaining the original graphite in the black mass and the fact that there might be a mixture of different anode manufacturers, we selected one commercially available, battery-grade graphite as a control for performance comparison. XRD and Raman spectra of this commercial graphite are provided in Fig. S12.†

In the first charge–discharge cycles (Fig. 7a and S13a†), the initial voltage drops from the open circuit voltage to approximately 1.5 V are consistent across all samples, indicating similar initial reactions including electric double layer, polarisation, and charge accumulation in the batteries. The most notable difference lies in the sloping region from approximately 1.5 V to about 0.2 V, where Li ions are adsorbed mainly by the defects, functional groups, and porous surfaces of graphite.<sup>43</sup> Commercial graphite exhibits the smallest sloping capacity, in agreement with the lower  $I_D/I_G$  ratio in the Raman spectrum. By contrast, all recycled graphite samples show higher sloping capacities due to their more defective structures. This increased sloping capacity results in higher irreversibility as solid electrolyte interphase (SEI) forms during the first cycles, thereby lowering the initial coulombic efficiency (CE).<sup>44</sup>

To elucidate the lithiation process, differential capacity plots ( $dQ/dV$ ) were derived for the plateau region between 0 and 0.3 V (Fig. 7b and c). Commercial graphite exhibited three prominent peaks at 0.18, 0.1, and 0.06 V, corresponding to the formation of different lithium intercalation compounds (LiC<sub>x</sub>,  $x = 1\text{--}6$ ), respectively.<sup>45</sup> These peaks showed lower intensities for cells assembled with regenerated graphite and became faint for that with spent graphite. The reduced intensities are associated with the presence of defects and mixed surface chemistry, while pristine graphite exhibits a highly crystalline structure with a fixed *d*-spacing that provides consistent energy to Li ions.

In rate tests, AG-2.0M-800 exhibited the highest specific capacity among all samples and an initial discharge capacity of 387.44 mA h g<sup>−1</sup> at a current of 0.1C (35 mA g<sup>−1</sup>), slightly higher than the theoretical capacity of graphite (372 mA h g<sup>−1</sup>). We attribute this to the high purity and presence of defects in AG-2.0M-800, allowing additional Li ion storage, as supported by results previously reported in the literature.<sup>4,15,36</sup> In long cycling tests performed following the rate test for the same cells, AG-2.0M-800 demonstrated an initial specific charge capacity of

373.45 mA h g<sup>−1</sup> at 0.5C (175 mA g<sup>−1</sup>) and no apparent capacity decay (Fig. 7f). Overall, these electrochemical analyses indicate that the rate capability and cycling stability of AG-2.0M-800 are comparable or superior to those of commercial graphite, confirming its viability for reuse as an anode material in LIBs.

## Conclusion

In summary, this study presents a streamlined process for recycling spent graphite from industrial black mass. The optimised recycling workflow comprises a primary acid leaching step, a secondary acid leaching step, and final mild-temperature pyrolysis, under the following operating conditions:

- (1) Initial acid leaching: a mixture of H<sub>2</sub>SO<sub>4</sub> (1.5 M) and H<sub>2</sub>O<sub>2</sub> (1.5 v/v%), a temperature of 90 °C, a solid–liquid ratio of 100 g L<sup>−1</sup>, and a leaching duration of 40 min.
- (2) Secondary acid leaching: H<sub>2</sub>SO<sub>4</sub> (2 M), a temperature of 90 °C, solid–liquid ratio of 50 g L<sup>−1</sup>, and leaching duration of 3 hours.
- (3) Pyrolysis: temperature of 800 °C under N<sub>2</sub> over 6 hours.

In the primary acid leaching stage, we optimised the leaching conditions by utilising self-contained Al, Cu, and LFP as reductants to maximise metal extraction from cathode materials. Following this, a secondary acid leaching process was conducted with a higher acid concentration and prolonged reaction time to remove residual impurities, particularly Al<sub>2</sub>O<sub>3</sub>. The acid leaching processes effectively minimised the levels of cathode materials and F content, preventing unnecessary graphite loss from reduction reactions of residual cathode materials during pyrolysis. Pyrolysis is beneficial not only for the complete removal of PVDF binders but also for repairing the structural defects in spent graphite, with the mild temperature minimising the operational costs. The overall purity of regenerated graphite reached 99.8% with a total recovery ratio of 91% as summarised in Scheme 1. The graphite loss is primarily due to size sieving and transferring at each step, with further optimisation being possible during the scale-up process.

Despite only partial recovery of structural crystallinity and slightly lower coulombic efficiency, graphite recycled from this workflow demonstrated high specific charge capacity and high capacity retention during long cycling, both superior to pristine battery-grade graphite. A preliminary economic analysis was conducted to compare the proposed recycling method with the conventional coal-based synthetic method for producing 1 kg of battery-grade graphite. We estimated the total inputs and outputs, including energy, water, and materials, for each stage of the proposed process, as outlined in Table S6 and Fig. S14.†<sup>9</sup> The proposed recycling method resulted in significant reductions in energy and water consumption, achieving 75% and 81% savings, respectively, compared to graphite production from coal.<sup>9</sup> These improvements are primarily attributed to the elimination of the mining and refining steps, responsible for approximately 81% of the total energy and water usage in conventional methods. Importantly, the workflow can integrate with existing facilities and cathode recycling processes, offering opportunities for the recycling of all critical materials from spent LIBs with minimal adjustments.



## Data availability

The authors confirm that the data supporting the findings of this study are available within the article and its ESI.† Raw data that support the findings of this study are available in figshare at <https://figshare.com/s/90994df100eb2c7aaf7f>.

## Author contributions

XW: conceptualisation, methodology, data curation, validation, writing and review. ZG: battery tests, battery data analysis, XPS and Raman tests, and review. YZ: battery tests and battery data analysis. YS: ICP-MS measurement. AH: supervision, resources and review. MT: supervision, resources, funding acquisition and review.

## Conflicts of interest

The authors declare no conflict of interest.

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## References

- 1 G. Harper, R. Sommerville, E. Kendrick, L. Driscoll, P. Slater, R. Stolkin, A. Walton, P. Christensen, O. Heidrich, S. Lambert, A. Abbott, K. Ryder, L. Gaines and P. Anderson, *Nature*, 2019, **575**, 75–86.
- 2 J. Yang, E. Fan, J. Lin, F. Arshad, X. Zhang, H. Wang, F. Wu, R. Chen and L. Li, *ACS Appl. Energy Mater.*, 2021, **4**, 6261–6268.
- 3 J. Wu, M. Zheng, T. Liu, Y. Wang, Y. Liu, J. Nai, L. Zhang, S. Zhang and X. Tao, *Energy Storage Mater.*, 2023, **54**, 120–134.
- 4 S. Ji, A. Zhang, W. Hua, S. Yan and X. Chen, *Battery Energy*, 2024, **3**, 20230067.
- 5 Lithium-ion Battery Market Size, Share & Growth Report, 2030, [https://www.grandviewresearch.com/industry-analysis/lithium-ion-battery-market?gad\\_source=1&gclid=Cj0KCQjww5u2BhDeARIIsALBuLnPchSsVO9vO-ti2KJtF2elpRgwhGtUo7jRVrs3VViF61ZddTDgOXkaAgEFEALw\\_wcB](https://www.grandviewresearch.com/industry-analysis/lithium-ion-battery-market?gad_source=1&gclid=Cj0KCQjww5u2BhDeARIIsALBuLnPchSsVO9vO-ti2KJtF2elpRgwhGtUo7jRVrs3VViF61ZddTDgOXkaAgEFEALw_wcB), (accessed 28 June 2024).
- 6 B. K. Sovacool, *Extr. Ind. Soc.*, 2019, **6**, 915–939.
- 7 M. Olutogun, A. Vanderbruggen, C. Frey, M. Rudolph, D. Bresser and S. Passerini, *Carbon Energy*, 2024, **6**, e483.
- 8 B. Markey, M. Zhang, I. Robb, P. Xu, H. Gao, D. Zhang, J. Holoubek, D. Xia, Y. Zhao, J. Guo, M. Cai, Y. S. Meng and Z. Chen, *J. Electrochem. Soc.*, 2020, **167**, 160511.
- 9 W. Chen, R. V. Salvatierra, J. T. Li, C. Kittrell, J. L. Beckham, K. M. Wyss, N. La, P. E. Savas, C. Ge, P. A. Advincula, P. Scotland, L. Eddy, B. Deng, Z. Yuan and J. M. Tour, *Adv. Mater.*, 2023, **35**, 2207303.
- 10 H. Wang, Y. Huang, C. Huang, X. Wang, K. Wang, H. Chen, S. Liu, Y. Wu, K. Xu and W. Li, *Electrochim. Acta*, 2019, **313**, 423–431.
- 11 S. Rothermel, M. Evertz, J. Kasnatscheew, X. Qi, M. Grützke, M. Winter and S. Nowak, *ChemSusChem*, 2016, **9**, 3473–3484.
- 12 J. Liu, H. Wang, T. Hu, X. Bai, S. Wang, W. Xie, J. Hao and Y. He, *Miner. Eng.*, 2020, **148**, 106223.
- 13 L. Barbosa, F. Luna-Lama, Y. González Peña and A. Caballero, *ChemSusChem*, 2020, **13**, 838–849.
- 14 J. Zhang, J. Hu, Y. Liu, Q. Jing, C. Yang, Y. Chen and C. Wang, *ACS Sustainable Chem. Eng.*, 2019, **7**, 5626–5631.
- 15 X. Ma, M. Chen, B. Chen, Z. Meng and Y. Wang, *ACS Sustainable Chem. Eng.*, 2019, **7**, 19732–19738.
- 16 I. Rey, C. Vallejo, G. Santiago, M. Iturrondobeitia and E. Lizundia, *ACS Sustainable Chem. Eng.*, 2021, **9**, 14488–14501.
- 17 A. Chernyaev, A. Kobets, K. Liivand, F. Tesfaye, P. M. Hannula, T. Kallio, L. Hupa and M. Lundström, *Miner. Eng.*, 2024, **208**, 108587.
- 18 S. Natarajan and V. Aravindan, *Adv. Energy Mater.*, 2020, **10**, 2002238.
- 19 M. Joulié, E. Billy, R. Laucournet and D. Meyer, *Hydrometallurgy*, 2017, **169**, 426–432.
- 20 A. Porvali, A. Chernyaev, S. Shukla and M. Lundström, *Sep. Purif. Technol.*, 2020, **236**, 116305.
- 21 A. Chernyaev, J. Partinen, L. Klemettinen, B. P. Wilson, A. Jokilaakso and M. Lundström, *Hydrometallurgy*, 2021, **203**, 105608.
- 22 A. Chernyaev, Y. Zou, B. P. Wilson and M. Lundström, *Sep. Purif. Technol.*, 2022, **281**, 119903.
- 23 Y. Zou, A. Chernyaev, M. Ossama, S. Seisko and M. Lundström, *Sci. Rep.*, 2024, **14**, 1–14.
- 24 R. Zhang, Z. Meng, X. Ma, M. Chen, B. Chen, Y. Zheng, Z. Yao, P. Vanaphuti, S. Bong, Z. Yang and Y. Wang, *Nano Energy*, 2020, **78**, 105214.
- 25 S. Shukla, A. Chernyaev, P. Halli, J. Aromaa and M. Lundström, *Metals*, 2023, **13**, 1118.
- 26 Y. Shin, S. Kim, S. Park, J. Lee, J. Bae, D. Kim, H. Joo, S. Ban, H. Lee, Y. Kim and K. Kwon, *Renewable Sustainable Energy Rev.*, 2023, **187**, 113693.
- 27 N. Zhang, Z. Xu, W. Deng and X. Wang, *Electrochim. Energy Rev.*, 2022, **5**, 1–38.
- 28 S. Kim, J. Bang, J. Yoo, Y. Shin, J. Bae, J. Jeong, K. Kim, P. Dong and K. Kwon, *J. Cleaner Prod.*, 2021, **294**, 126329.
- 29 X. Luo, J.-C. Robin and S. Yu, *Nucl. Eng. Des.*, 2004, **227**, 273–280.
- 30 L. A. Gomez-Moreno, A. Klemettinen and R. Serna-Guerrero, *iScience*, 2023, **26**, 107782.
- 31 S. S. Choi and Y. K. Kim, *J. Anal. Appl. Pyrolysis*, 2012, **96**, 16–23.
- 32 G. Zhang, Y. He, Y. Feng, H. Wang and X. Zhu, *ACS Sustainable Chem. Eng.*, 2018, **6**, 10896–10904.
- 33 S. Babanejad, H. Ahmed, C. Andersson, C. Samuelsson, A. Lennartsson, B. Hall and L. Arnerlöf, *J. Sustain. Met.*, 2022, **8**, 566–581.
- 34 M. Chen, X. Ma, B. Chen, R. Arsenault, P. Karlson, N. Simon and Y. Wang, *Joule*, 2019, **3**, 2622–2646.



35 J. Asenbauer, T. Eisenmann, M. Kuenzel, A. Kazzazi, Z. Chen and D. Bresser, *Sustainable Energy Fuels*, 2020, **4**, 5387–5416.

36 R. Li, S. Zeng, L. Wang, X. Yu, H. Zeng, W. Liu, D. Fu and X. Liu, *J. Alloys Compd.*, 2024, **993**, 174691.

37 J. Yu, M. Lin, Q. Tan and J. Li, *J. Hazard. Mater.*, 2021, **401**, 123715.

38 X. Li, J. Shen, C. Wu and K. Wu, *Small*, 2019, **15**, 1805567.

39 Q. Xu, Q. Wang, D. Chen, Y. Zhong, Z. Wu, Y. Song, G. Wang, Y. Liu, B. Zhong and X. Guo, *Green Chem.*, 2021, **23**, 4531–4539.

40 X. Yang, R. Li, J. Yang, H. Liu, T. Luo, X. Wang and L. Yang, *Carbon*, 2022, **199**, 215–223.

41 Y. H. Xiao, J. Li, W. G. Huang, L. H. Wang and J. Luo, *J. Mater. Sci.: Mater. Electron.*, 2022, **33**, 16740–16752.

42 S. Xu, C. Liu, F. Ye, Y. Guo and J. Wiezorek, *Colloids Surf., A*, 2017, **515**, 1–11.

43 L. Zhao, B. Ding, X.-Y. Qin, Z. Wang, W. Lv, Y.-B. He, Q.-H. Yang, F. Kang and L. Zhao, *Adv. Mater.*, 2022, **34**, 2106704.

44 T. Placke, V. Siozios, R. Schmitz, S. F. Lux, P. Bieker, C. Colle, H. W. Meyer, S. Passerini and M. Winter, *J. Power Sources*, 2012, **200**, 83–91.

45 H. Fujimoto, H. Kiuchi, S. Takagi, K. Shimoda, K. Okazaki, Z. Ogumi and T. Abe, *J. Electrochem. Soc.*, 2021, **168**, 040509.

