Green Chemistry



PAPER View view





Cite this: *Green Chem.*, 2021, **23**, 8519

Received 27th July 2021, Accepted 29th September 2021 DOI: 10.1039/d1gc02693c

rsc.li/greenchem

Ultrasound-assisted extraction of metals from Lithium-ion batteries using natural organic acids†

Xiong Xiao, 🕩 Billy W. Hoogendoorn, Yiqian Ma, Suchithra Ashoka Sahadevan, James M. Gardner, 🕩 Kerstin Forsberg 🕩 and Richard T. Olsson 🕩 Xa

An ultrasound-assisted extraction (leaching) method of valuable metals from discarded lithium-ion batteries (LiBs) is reported. Mild organic citric or acetic acids were used as leaching agents for a more environmentally-friendly recovery of the lithium, nickel, cobalt, and manganese from the discharged and crushed lithium nickel-manganese-cobalt oxide (NMC) LiBs. The extraction was performed with the presence/absence of continuous ultrasound (US) energy supplied by a 110 W ultrasonic bath. The effect of temperature (30-70 °C), reducing agent concentration (H_2O_2 : 0-2.0 vol%), as well as choice of specific acid on the metal dissolution were investigated. The US leaching decreased the leaching time by more than 50% and improved the leached percentage of Li, Mn, Co, and Ni due to the local heat and improved mass transfer between solid and liquid interfaces in the process. The X-ray diffraction results of residues from the US leaching further confirmed an improved dissolution of the crushed layered NMC structure, resulting in the significant improvement of the leached amounts of the valuable metals. Furthermore, it is demonstrated that using citric acid eliminated the need of additional reducing agents and suppressed the dissolution of copper (Cu) due to its inhibitor effect on the Cu surface, i.e. compared with using acetic acid as leaching reagent. Overall, it is shown that recovery of the battery metals can be facilitated and carried out in a more energy-efficient manner at low temperatures (50 °C) using ultrasound to improve metal ions mass transportation in the residue layers of the NMC during the organic acid leaching.

1 Introduction

Lithium-ion batteries (LiBs) are increasing in demand due to their use in electronic vehicles (EVs). LiBs have been responsible for more than half of the energy output from all batteries since 2017 and the use of LiBs is expected to increase to 75% of the market by 2025. At the same time, the most frequently used functional metal elements, *i.e.* lithium, nickel, and cobalt, are limited in supply and global geological exploration predicts that a shortage of the elements may occur within 25 years. This calls for intense efforts to develop strategies to reuse, recover, and recycle different battery types to avoid the build-up of valuable materials in landfills. Resource efficient

metals recycling, allowing for continuous raw material supply,

is here a cornerstone in a future sustainable battery market. The recycling of valuable metals requires facile processing with high selectivity and demands as low energy input as possible, with minimal environmental impact for the extraction of the metal ions from their host materials. This ultimately demands that the energy and emissions (volatile compounds and toxic greenhouse gases) associated with the strategies to recover the metals are overviewed in complete lifecycle analysis (LCA) studies. 1,4 Hydrometallurgy represents a promising recycling method since it requires lower energy input and generates lower levels of toxic emissions than pyrometallurgy. 5,6 In hydrometallurgy recycling, battery shredding is frequently used to obtain the process raw material consisting of a mixture of iron, aluminium, copper, in addition to the active battery metals, and graphite, liquid electrolytes, plastics, and binders.⁷ The polymers are sieved off in a flotation process,⁶ leaving behind a mixture of materials that are processed with strong and harsh acids to dissolve the metals for further separation by leaching, i.e. before precipitation, 8 solvent extraction, 1 and/or electrowinning of the valuable metals.9 The drawbacks of the processing include occasionally time-consuming steps

^aDepartment of Fiber and Polymer Technology, KTH Royal Institute of Technology, Teknikringen 56, 11428 Stockholm, Sweden. E-mail: rols@kth.se

^bDepartment of Chemical Engineering, KTH Royal Institute of Technology, Teknikringen 42, 11428 Stockholm, Sweden

^cDepartment of Chemistry, KTH Royal Institute of Technology, Teknikringen 30, 11428 Stockholm, Sweden

 $[\]dagger\,\text{Electronic}$ supplementary information (ESI) available. See DOI: 10.1039/d1gc02693c

Paper

including calcination (ca.450-700 °C), organic solution treatment, and NaOH treatment for removal of battery binders and Al, with reported losses of valuable metals, e.g. lithium, etc. and emission of toxic greenhouse gases. $^{10-14}$

Previously, ultrasound was demonstrated as an efficient and green technology that can be used in the process of extraction from biological samples. Most ultrasonic baths send mechanical pressure waves with frequencies greater than 20 kHz through the liquid medium. By sending the ultrasonic waves through an aqueous medium, microbubbles are formed. 15 The collapse of the bubbles generates local temperatures of 5000 K along with the production of highly reactive free radicals.¹⁶ The events will cause physical effects, such as turbulence, shear forces, shock waves, and microjets, resulting in increased mass transfer while degassing the solution of the dissolved gases. Ultrasound may however also allow for improving mass transfer in the extraction of valuable battery metals to an extent that more environment-friendly organic acids can be used in the recovery of the metals. So far, investigations on the extent of the improved efficiency of metal extraction from used LiBs using ultrasound are scarce. 12,13,17,18 In these studies, the ultrasound extraction of metals from LiBs focuses on the dissolution of LiCoO2, LiNiO2, and LiMn2O4, and occasionally reports on the leaching kinetics of these materials with the complexity of added reducing agents. The more environment-friendly acids, so far reported, include formic acid, 19 lactic acid, 14 acetic acid, 20 and citric acid. 21 In the absence of ultrasound, these weaker organic acids would normally be regarded as insufficiently strong for efficient extractions and require the use of additional chemicals or pre-treatments steps before the actual extraction experiments (leaching

bly of batteries.22 In this work, the ultrasound-assisted extraction of the metals from lithium nickel-manganese-cobalt oxide (NMC) batteries (the most common lithium battery type) is reported for the first time, and the kinetics of the extraction is described in detail. The difference between leaching with, and without ultrasound, using weak acetic or citric acids is shown. Citric acid was used as a natural, mildly reducing ($E_0 = -0.18 \text{ V}$ vs. NHE), weak triprotic acid that is commercially produced by fermentation.^{18,23} It is 'green', abundant, comparatively inexpensive (to other organic acids), and has been widely used by the food and beverages industry, and as a non-toxic household detergent for years.24 The citric acid was compared with acetic acid due to its inexpensive nature at high purity, and high availability, although not as acidic as the citric acid i.e. being the sole organic acid with a p K_a value of 4.76 ²⁵ (see Table S2†). The batteries used were shredded 24 kW h Volvo C30 NMC batteries. The leaching processing was however carried out as a universal approach, making the demonstrated methods applicable to work with other battery types. It is shown that ultrasound significantly increases the leaching rate of the battery metals during the processing. The leaching time

process) to reach recovery levels >95%, see Table S1.† In LiBs

recycling, low and high-power ultrasound vibration was used to assist the electrode delamination process during disassemwas reduced by more than 50% when using a standard laboratory ultrasonic bath, as opposed to when carrying out traditional extraction in a stirred reactor. Almost complete metal ion extraction is demonstrated, using the acids in combination with the ultrasound. The citric acid further improved the leached percentage of lithium, nickel, cobalt, and manganese while suppressing the dissolution of Cu, when compared with acetic acid. The results were confirmed with multiple measurements and characterizations of all battery residues before, and after, the leaching process.

2 Experimental

2.1 Materials

Discarded NMC LiBs were delivered as a shredded 'blackmass' powder from Volvo Cars AB (Sweden), Fig. 1a. The 'blackmass' was prepared by crushing followed by removing a magnetic fraction via magnetic separation. Plastics, copper, aluminium foils were removed by air separation and sieving, and the binder was kept in contrast to previous works adapting ultrasound to remove the binder.22 The use of calcination at >600 °C was avoided to reduce the energy demands of the entire recycling/recovery of metals procedure, eliminating the unnecessary generation of greenhouse gases.26 The citric acid $(C_6H_8O_7, \geq 99.5\%)$, acetic acid $(C_2H_4O_2, \geq 99.7\%)$, 37% HCl (analytical grade), $\geq 65\%$ HNO₃ (analytical grade), and hydrogen peroxide (H2O2, 30 wt%) were purchased from Sigma-Aldrich AB, Sweden. Standard solutions of 1000 mg L⁻¹ ICP-OES for Li, Ni, Co, Mn, Cu, Al, and Fe in 2-5 w% HNO3 or HCl solutions for ICP-OES calibration were purchased from Alfa Aesar, Sweden. MiliQ water (>18.2 M) was used in all the experiments.

2.2 Leaching/extraction methods

2.2.1 Aqua regia leaching digestion. LiBs powder (1 g) was dissolved in the 100 mL aqua regia solution $(\text{HNO}_3: \text{HCl} = 1:3, \text{v/v})$ at 343 K for 5 h to determine the contents of metals in the ground LiBs powder. The experiment was repeated thrice to ensure that the contents of the metals in the reference the sample was accurately determined.

2.2.2 Ultrasound (US) leaching. The leaching experiments were performed in 500 mL three-necked round bottom flasks equipped with condenser units for recirculation of any evaporated liquid. LiBs powder (5 g) was added to the round bottom flasks with 200 mL of 1.5 mol $\rm L^{-1}$ citric acid, or 200 mL of 4.5 mol $\rm L^{-1}$ acetic acid as aqueous solutions. 0 to 2.0 vol% $\rm H_2O_2$ was added as a reducing agent depending on the experiment. The acid concentrations and the solid to liquid (S/L) ratio was motivated from results based on screening experiments, see Fig. S1.† The round bottom flasks were placed in the same position in the ultrasonic bath (BRANSON 2800 CPX2800H-E, with a max sonics power of 110 W at a frequency of 40 kHz) in all ultrasound experiments. The bath temperature was maintained at 323 K (\pm 2 K). The extracted metal ions contents were determined from 2 ml aliquots taken in regular

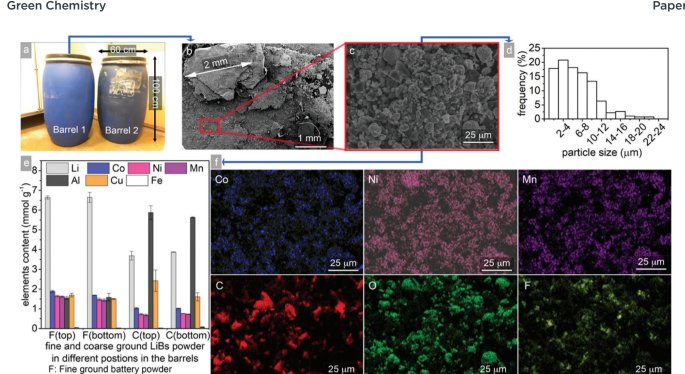


Fig. 1 (a) Barrels of discharged and ground batteries, (b) SEM images of fine ground LiBs powder, and (c) the fine ground powder in high magnification, (d) size distributions of fine ground powder derived from >50 micrographs using min, 600 particles for accurate statistics, (e) metal elements content variation (ICP-OES measurements) from top to bottom of the Barrel 1 and 2, respectively, (f) EDS maps taken from (c). Triple samples tested on each location in the barrels confirmed the nonuniform powder composition from the natural fractionation that had occurred in the coarse ground powder, demonstrated by the error bars in Fig. 1e.

intervals during the progress of extraction and separated from residues by centrifugation at 13 400 rpm for 2 min before the removal of the solid phase, dilution, and analysis by inductively coupled plasma-optical emission spectrometry (ICP-OES).

C: Coarse ground battery powder

2.2.3 Oil bath leaching. The leaching was carried out in the same round bottom flasks that were used in the US leaching experiments with the only difference being that the round bottom flasks were placed in an oil bath. Teflon-coated magnets sized 2 cm in length were put inside the flasks for continuous stirring at 500 rpm during the entire extraction process at different temperatures. With reference to the previous literature (Table S1†), 303 K to 343 K served as good points for comparisons. The aliquots were taken in the same manner as during US leaching experiments. After the leaching experiments, the residues were filtered, washed with MiliQ water, and dried at 343 K before further characterizations. All the leaching experiments were carried out under closed and sealed reactions conditions and well-ventilated areas to minimize exposure to the LiPF₆ electrolyte.

2.3 Characterization

2.3.1 Thermal characterization. Thermogravimetric analysis (TGA) was performed on a Mettler-Toledo TGA/SDTA851 instrument. The measurements were carried out under 50 mL min⁻¹ oxygen from 40 to 950 °C, with a heating rate of 10 °C min^{-1} .

2.4 Microscopy

The morphology and size distribution of the samples were observed in a field emission scanning electron microscope (FE-SEM, Hitachi S-4800) and SEM images were taken at an acceleration voltage of 15.0 kV, and a current of 10 mA. The size of the particles was obtained by manually measuring 600 particles using ImageJ (National Institute of Health, Maryland, USA). Sputtering of the samples was performed for 20 s using a Cressington 208HR, equipped with a Pt/Pd target. Energydispersive X-ray spectroscopy (EDS) was carried out at 15.0 kV to measure the distribution of the elements in the LiBs powder before and after leaching.

Characterization of elemental compositions

All solutions were diluted into metal concentration ranges useful in the elemental analysis. The concentrations of metal ions in the solutions were determined by ICP-OES (Thermo Fisher iCAP 7400, USA), using at least 3 suitable wavelengths that did not interfere with the wavelengths of other metal ions in the solution. The calculation of the leached percentage is represented as follows:

$$E(\%) = \frac{C_{\rm s}({\rm mmol}\,{\rm g}^{-1})}{C_{\rm 0}({\rm mmol}\,{\rm g}^{-1})} \times 100\% \tag{1}$$

where E (%) is the leached percentage, C_s (mmol g^{-1}) is the concentration of metals extracted to the solution per gram of

the added blackmass, and C_0 (mmol g^{-1}) is the corresponding total amount of leachable metals determined by digestion in aqua regia (mmol g^{-1}). C_s and C_0 are calculated from the raw

ICP-OES results, using the following eqn (2):

$$C(\text{mmol g}^{-1}) = \frac{c(\text{mg L}^{-1}) \times V(L)}{M(\text{g mol}^{-1}) \times m(\text{g})}$$
(2)

where C (mmol g⁻¹) represents C_s and C_0 , c (mg L⁻¹) is the raw data from ICP-OES results, V(L) is the volume of solution, M(g) mol^{-1}) is the mole weight of metals, and m(g) is the weight of the ground LiBs powder added in the solution.

2.6 Crystallography

Paper

The ground LiBs powder and all sample residues were characterized by X-ray diffraction (XRD), and the PANalytical X'pert3 software was used for peak identification. The XRD patterns were recorded on a powder diffractometer operating at 45 kV and 40 mA using Cu K α radiation ($\lambda = 0.15418$ nm).

2.7 Viscosity

Kinetic viscosities of the aqueous solutions were measured using an Ubbelohde capillary viscometer from Schott. The capillary used was a type II (Di = 1.13 mm, $K \approx 0.1$) and a type 0c (capillary constant = $0.0031476 \text{ mm}^2 \text{ s}^{-2}$). The capillary was placed in a Lauda iVisc (LAUDA, Germany), an automated infrared measuring device, which measures the time of the experiments. The iVisc with capillary was placed in a water bath (Lauda ET 15) at 298 K, and the temperature was regulated with an immersion heating thermostat (Lauda Eco Silver).

3 Results and discussion

Analysis of the ground LiBs powder provided

Fig. 1a shows the two barrels containing fine and coarse ground LiBs 'blackmass' powder. The barrels were 100 × 60 cm (height × diameter), containing ca. 100 kg ground powder. Fig. 1b and c shows that even the finest ground battery powder displayed significant particle size differences. The size distribution of the fine ground powder is displayed in Fig. 1d, showing that the average particle size was in the range of ca. 2–8 μ m (average: 5.9 \pm 0.3 μ m), although large sized flakes sized up to 2 mm could occasionally be observed Fig. 1b.

Fig. 1e shows the elemental presence in the top and bottom powder fractions of the barrels for the fine and coarse ground powder, as a comparison. The coarse ground powder contained a markedly higher content of aluminium and copper compared to the more valuable metals Co, Ni, and Mn (right hand side columns, Fig. 1e). The coarse ground powder also demonstrated a wider particle size distribution (Fig. S2b†), which explained a more extensive fractionation that had occurred during the barrel transportation. It was thus evident from the elemental analysis and microscopy that only the fine ground powder, with almost identical metal composition on the top and bottom (left hand side columns, Fig. 1e) was

useful as starting material for developing optimized extraction protocols, i.e. due to its more uniform composition. The fine ground powder was therefore further analysed and used in all the remaining experiments. The larger size particles in Fig. 1b (representing less than 5% of the particle fraction, although a considerable part of the metallic mass content) was however firstly filtered off and identified as composed of aluminium or copper (Fig. S2†). The larger sized particles were accordingly excluded from the extraction protocols focusing on the active battery materials.

Fig. 1c shows a magnified area in Fig. 1b, displaying the finer particles, with the dominant phase represented in particle sizes by the histogram shown in Fig. 1d. Fig. 1f shows the EDS mapping images of the same area displayed in Fig. 1c. The detected elements were Co, Ni, and Mn (top row), while the bottom row shows C, O, and F. The latter elements were remains from the conductive additives and binders used in the LiBs fabrication process. Lithium is not shown in Fig. 1f since it was not detected by the normal EDS probe analysis due to its low characteristic radiation energy.²⁷ For comparison, Fig. S3† shows the EDS mapping of the residue of the same LiBs powder after aqua regia leaching digestion, demonstrating that the aqua regia leaching digestion could effectively be used for complete extraction (100%) of the Ni, Co, to obtain the reference values synonymous with complete extraction of all active battery metals from the remaining carbon residuals.

Fig. 2 shows the TGA curves and XRD patterns of the fine ground LiBs powder before and after aqua regia leaching digestion. TGA was carried out under O2 flow to ensure that all organic materials in the fine ground LiBs powder were removed, and its relative content could be determined. The results show that before aqua regia leaching digestion about 61.8 wt% residue remained after heating to 950 °C, which was associated with the metal fraction of the battery powder. After aqua regia leaching digestion, less than 1 wt% material remained although a small amount of ash existed. The results were consistent with an almost complete oxidation of the organic materials (Fig. 2a), at a temperature above 500 °C. Moreover, XRD characterization was used to investigate the crystalline phases in the fine ground powder before and after the aqua regia leaching digestion.

The peaks in the diffractograms before leaching originated from the layered NMC phase (active material), Al, Cu, and graphite, respectively, Fig. 2b (top). After the aqua regia leaching digestion, the remaining material was graphite and polypropylene (PP). The PP came from the separator/battery casing, see Fig. 2b (bottom). The TGA and XRD of powders before and after aqua regia leaching agreed with the EDS mapping and confirmed that a full extraction of the metal elements had been carried out.

ICP-OES was used to carry out the quantitative analysis of the aqua regia leachate composition. The concentration (mmol g⁻¹) and standard deviation of all metals are summarized and shown in Table 1. Fe was detected in the fine ground LiBs powder at a concentration of 0.048 mmol (pristine powder) g⁻¹, which is not included in Table 1 due

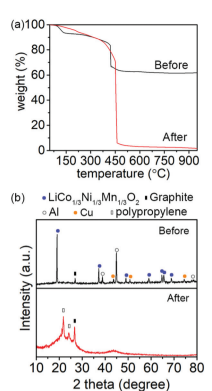


Fig. 2 (a) TGA with O_2 flow and (b) XRD of the fine ground LiBs powder before and after aqua regia leaching digestion. The peaks for related phases referred to ICDD database: (LiNi $_{1/3}$ Co $_{1/3}$ Mn $_{1/3}$)O $_2$ (NMC) (PDF 56-0147), graphite (PDF 08-0415), Al (PDF 01-1180), Cu (JCPDS 003-1018), and polypropylene (PDF 61-1416).

Table 1 ICP-OES results of the metal elements extracted from the fine ground LiBs powder by aqua regia

	Metal elements (mmol (pristine powder) g^{-1})						
ICP-OES result	Li	Mn	Со	Ni	Al	Cu	
F ^a STDEV	6.642 0.078	1.623 0.027	1.644 0.036	1.886 0.047	1.550 0.073	1.690 0.092	

^a F: the fine ground LiBs power from the barrel 1.

to its comparably small amount. The metal mole-to-mole ratio of the active material Li:Mn:Co:Ni was ca. 1.289:0.315:0.319:0.366, reflecting that the composition of active material (LiMn_{0.315}Co_{0.319}Ni_{0.366}O₂) and electrolyte in the hybrid EV battery was closed to LiNi_{1/3}Co_{1/3}Mn_{1/3}O₂. The phase composition was also in agreement with the diffraction pattern in Fig. 2b (top), although the additional 29% Li originated from the electrolyte (LiPF₆) that was present in the sample. The Al and Cu in Fig. 2b originated from the current collector and were not shown in the EDS mapping (Fig. 1f) due to their large particle sizes and coincidentally were found in similar amounts as the targeted metal ions (Ni, Co, and Mn). In total, the weight of Al, Cu, and the NMC phase occupied ca. 65.7 wt% of the fine ground powder, which was in good agree-

ment with the remaining residue percentage (61.8 wt%, Fig. 2a) obtained from the gravimetric measurements.

In summary, the characterization in terms of the natural fractionation and deviation of uniformity (coarse νs . fine ground powder), the phase characterization, and the chemical composition allowed only the fine ground powder to be identified as a useful reference material. The coarse ground powder was always displaying different powder characteristics at different locations in the barrel and was therefore discarded as a source for more experiments.

3.2 Effect of ultrasound on metal ion extraction using soft acids

Fig. 3 shows the effect of ultrasound on the metal ion extraction from the fine ground LiBs powder using 1.5 mol L^{-1} citric acid, in the absence of any H_2O_2 reducing agent, which commonly is used to improve leaching efficiency. The most extended leaching times were selected to 24 h (1440 min) on

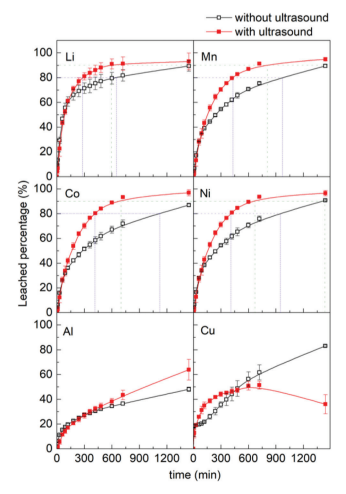


Fig. 3 Effect of leaching method (US vs. oil bath) on metal ions extraction from the fine ground LiBs powder. Error bars are standard errors from triplicate measurement of different batches. Conditions: citric acid, 1.5 mol L⁻¹; S/L, 25 g L⁻¹, H₂O₂, 0 vol%; ultrasonic bath temperature, 323 K; oil bath temperature, 323 K; stirring speed, 500 rpm. The lines between measurement points serve as the guide to the eye.

Paper Green Chemistry

the basis of possibly reaching full extraction of the NMC material within a reasonable time frame. In the first 150 min, the leaching behaviour of the Li, Mn, Co, Ni, and Al in the US leaching system and the traditional oil bath were similar and the extracted amount per time was almost identical, see overlapping curves in Fig. 3. After 150 min, the leached percentage of Li, Mn, Co, and Ni in the US leaching system was higher than that in the oil bath, reaching on average ca. 97 \pm 3% after 1440 min. Among Li, Mn, Co and Ni, the most effective leaching was found for the cobalt reaching 100% extraction, while Li and Mn were extracted to 94% and 96%, see Table 2. Also, the leached percentage of Al at 1440 min was significantly improved from 48% to 60% using the ultrasound in the leaching process. An increase in temperature from 323 to 343 K (Fig. S4†, Table 2) of the oil bath was insufficient to reach the same extraction efficiency as when solely using the ultrasound for the Li, Mn, Co, Ni metal ions, although Al leaching was sensitive to an increase temperature from 323 K to 343 K. More specifically, raising the temperature 20 K resulted in more aluminium extracted compared to when only ultrasound was used.

Table 3 show over time that the active battery metals could be extracted much faster using ultrasound. The required times for reaching 20%, 50%, 80%, and 90% of the total amount of extractable metal ions (Li, Ni, Co, and Mn) are shown in Table 3.

Among the metals, the fastest extraction was observed for the lithium, showing that 80% of the total amount of extractable Li $(t_{80\%})$ could be reached within 270 min. The same extent

Table 2 The leached percentage of the metal ions at 1440 min using citric acid as reagent in an ultrasound (US) leaching system and an oil bath (323 K, 343 K) system

	Metal elements (mmol (pristine powder) g^{-1})						
ICP-OES result	Li	Mn	Со	Ni	Al	Cu	
Oil bath, 323 K	89%	89%	87%	91%	48%	83%	
US, 323 K	94%	96%	100%	99%	60%	48%	
Oil bath, 343 K	90%	91%	93%	92%	78%	54%	

Comment: The data for each ion was acquired as repeated twice with separate ICP-OES measurements. The deviation was maximum ±3% with an exception of Cu (maximum $\pm 7\%$).

Table 3 Required times (min) for reaching different leached percentages using citric acid and ultrasound. Leaching times in absence of ultrasound in parentheses

Times (t, min)	Li	Mn	Со	Ni
t _{20%} t _{50%} t _{80%} t _{90%}	20 (20)	44 (40)	43 (45)	47 (41)
	75 (75)	140 (250)	160 (290)	150 (240)
	270 (650)	430 (975)	420 (1125)	420 (940)
	600 (>1440)	790 (>1440)	685 (>1440)	675 (1440)

Comment: The data in the table with parentheses represent the required time in an oil bath. Data from Fig. 3, see dashed lines for 80 and 90%.

of extraction, using traditional stirring, required over 650 min to be completed. At the same time, 420 min was needed to extract 80% of the cobalt ($t_{80\%}$) using the ultrasound in comparison to 1125 min using the traditional extraction, see Table 3 and Fig. 3.

The X-ray diffraction patterns of residues, as dependent on the leaching time with/without ultrasound, are shown in Fig. S5[†] and Fig. 4. From 0 to 720 min, a shift of the 19.05° peak (representing the NMC interlayer direction), occurred from a higher to a lower angle (Δ 0.8°, see Fig. 4a). The shift was caused by the proton exchange accompanied with water molecules²⁸ intercalated within the NMC layered space, resulting in an expansion of the interlayer distance, see Fig. 4b. With an additional increase in the leaching time to 960 min, the peak shifted back to a higher angle due to the destruction of the layered NMC phase and the formation of a spinel-type LiMn₂O₄. The spinel-type LiMn₂O₄ can retain its structure even when exposed to harsh HCl acid solutions.²⁹ After 1440 min leaching, the diffractogram peaks associated with the LiMn₂O₄ spinel structure disappeared in the residues from the US leaching method while they remained in the residues from the oil bath leaching at 323 K. The results suggested that the ultrasound not only assisted in the diffusion of the valuable metal ions into the solution, but also had an impact on the dissolution of the more stable LiMn₂O₄ spinel structure.

The only exception from the general observation that ultrasound assisted leaching outperformed the traditional oil bath

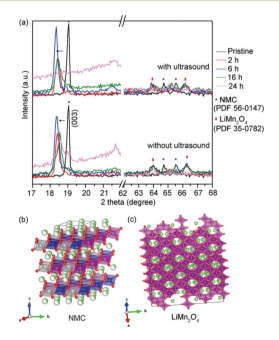


Fig. 4 (a) X-ray diffraction patterns of residues in the leaching system with/without ultrasound depending on leaching time at 323 K. (b) Crystal structures of $LiCo_{1/3}Ni_{1/3}Mn_{1/3}O_2$ (NMC) and (c) spinel-type $LiMn_2O_4$ (draw by VESTA³⁰), Li (green ball/octahedral), Ni/NiO₆ (grey ball/octahedral), Co (blue ball/octahedral), and Mn (purple ball/octahedral), vacancy (white ball).

Green Chemistry Paper

leaching was the Cu element. The leached percentage of Cu at 1440 min was supressed during the leaching, see Table 2. This was related to the depletion of dissolved O_2 in the solution (caused by the intense ultrasound), which have been reported to interact with Cu and citric acid in conditions without any oxidizing agents.³¹ For this reason, ultrasound has traditionally also been used for degassing and release of gases from solutions.^{32,33} A lower O_2 concentration consequently interfered with the Cu oxidation and dissolution mechanisms as previously reported.³¹ This would be in line with the results showing that extraction carried out at higher temperature in the oil bath, see Table 2 (343 K) also suppressed Cu leaching since a higher temperature results in a lower percentage of naturally dissolved O_2 .³⁴

3.3 Kinetics study with/without ultrasound on metal ions extraction

To further understand how ultrasound influenced the leaching process, the shrinking core model of noncatalytic fluid-solid reactions 20,35 was used to assign the contribution of different mass transfer phenomena to the observed leaching behaviour. The schematic diagram of the shrinking core model was shown in Scheme S1 \dagger and the kinetics of it is described by the following three equations representing different leaching rate contributions to the model:

(1) Liquid boundary layer mass transfer contribution (eqn (3)):

$$t = \frac{\chi R \rho_{\rm P}}{3M k_{\rm f} C_{\rm A}} X \tag{3}$$

(2) Surface chemical reaction contribution (eqn (4)):

$$t = \frac{\chi R \rho_{\rm P}}{M k_{\rm R} C_{\rm A}} [1 - (1 - X)^{1/3}] \tag{4}$$

(3) Residue layer diffusion contribution (eqn (5)):

$$t = \frac{\chi R \rho_{\rm P}}{6Mk_{\rm L}C_{\rm A}} [1 - 3(1 - X)^{2/3} + 2(1 - X)]$$
 (5)

where t represents the leaching time (min), χ is the electron transfer number in reactions, R is the radius (m) of the particle at t=0, ρ_P is the density (g m⁻³) of the ground LiBs powder, M is the molar weight (g mol⁻¹) of the ground LiBs powder, C_A is the concentration (mol m⁻³) of the acid solution at t=0, X represents the fraction (%) of metals leached, k_f , and k_L are the mass transfer coefficient (m min⁻¹) in the liquid boundary layer, and in the residue layer respectively, and k_R is the reaction rate constant (m min⁻¹). For simplification, the χ , R, ρ_P , M, and C_A were used as fixed values in this study since the same fine ground powder was always used, as well as the leaching conditions $C_A=1.5$ mol L⁻¹ (323 K). Therefore, the leaching rates contributions could be simplified to the following eqns:

$$X = k_1 t \tag{6}$$

$$k_1 = \frac{3Mk_f C_A}{\chi R \rho_P}$$

$$1 - (1 - X)^{1/3} = k_2 t$$

$$k_2 = \frac{M k_R C_A}{\chi R \rho_P}$$
(7)

$$1 - 3(1 - X)^{2/3} + 2(1 - X) = k_3 t$$

$$k_3 = \frac{6Mk_{\rm L}C_{\rm A}}{\gamma R \rho_{\rm P}}$$
(8)

where, k_1 , k_2 , and k_3 (min⁻¹) are the slopes of the fitted lines of X, $(1-X)^{1/3}$ and $1-3(1-X)^{2/3}+2(1-X)$ vs. t (leaching time), respectively. The different contributions were accordingly fitted to the leaching data from the curves shown in Fig. 3. Fig. 5 shows the fitting of the data to the residue layer diffusion contribution factor $(1-3(1-X)^{2/3}+2(1-X)$ vs. t), whereas Fig. S6† shows the same data fitted to the contributions refer as liquid boundary layer mass transfer (X vs. t) and surface chemical reaction $((1-X)^{1/3} \text{ vs. } t)$.

The results show that for both systems (with/without ultrasound), the fitting lines of Li, Mn, Co, Ni, and Al were in agreement with the residue layer diffusion contribution to the shrinking core model. A linear relationship (k_3) with a higher regression coefficient (R^2) value (≥ 0.9855) could be established, whereas no relationships could be established for any of the other contributions to the shrinking core model (Fig. S6†).

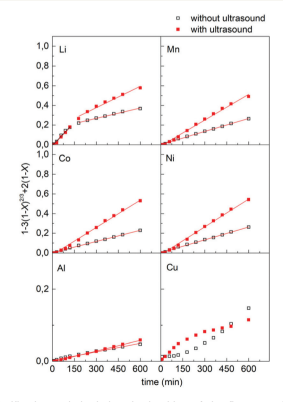


Fig. 5 Kinetics analysis during the leaching of the fine ground LiBs powder with/without ultrasound, adapting leaching time data to the residue layer diffusion contribution (eqn (5)) of the shrinking core model. Conditions: citric acid, 1.5 mol L^{-1} ; S/L, 25 g L^{-1} , H₂O₂, 0 vol%; temperature, 323 K; stirring speed, 500 rpm.

Li (2nd stage) Li (1st stage) Ni Αl Mn Co $k_3 \times 10^{-4}$ (\min^{-1}) (\min^{-1}) (\min^{-1}) (\min^{-1}) (min^{-1}) (min^{-1}) R^2 Without 15.6 0.9916 3.52 0.9917 0.9980 0.9981 0.9981 0.9855 4.41 3.81 4.36 0.79 US With US 0.9982 0.9814 7.28 0.9763 8.75 0.9944 9.30 0.9945 0.9956 0.99 15.3 9.47

were assessed from the residue layer diffusion contribution to the shrinking core model

Table 4 Kinetic parameters during the leaching process of the metals from the fine ground LiBs powder without/with uhltrasound. The values

Comment: Copper was excluded in the table because of its low fitting R^2 value from the US leaching system, regardless of the control models described in the shrinking core model. Conditions: citric acid, 1.5 mol L⁻¹; S/L, 25 g L⁻¹, temperature, 323 K; stirring speed, 500 rpm.

The specific R^2 values are shown in Table 4 for the different ions. The only exception was the leaching of Cu that instead reasonably well correlated with the reaction control contribution, showing a R^2 value of 0.989 when the extraction was done in absence of ultrasound (Fig. S6†). This result again suggested that having oxygen dissolved in the solution contributed to a uniformly progressing Cu dissolution reaction (see section 3.2). Overall, it was evident that the rate limiting contribution to the US leaching behavior could be assigned to the residue layer diffusion (eqn (5) and (8)), while the ultrasound favoring the residue layer diffusion may also be used to suppress the dissolution of copper, compare Table 2.

Table 4 shows the k_3 values for Li, Mn, Co, Ni, and Al, respectively, as well as their related R^2 values from the fitted slopes based on residue layer diffusion contribution (Fig. 5). The k_3 values represent the release rate of the metal ions as the steepness of the linear slope in Fig. 5. It is shown that the Li extraction that required a two-stage fitting of the residue layer diffusion contribution in the 1st stage leaching demonstrated k_3 values that were independent of the leaching method (oil bath or ultrasound), and ca. 2-4 times higher than other metal ions in the NMC. This was caused by some of the lithium originating from the electrolyte, making it easier to

release to the acid solution. During the 2nd stage, the release of the lithium was dominantly from the solid phase and therefore occurred at a slower rate, with k_3 value similar to the Mn, Co, and Ni metal ions.

Notably, although the k_3 values for Mn, Co, and Ni were similar (Fig. 5 and Table 4), ultrasound always yielded higher release rates (regardless of the specific metal ion) suggesting that this release was caused by ultrasound impact on the solid phase. This observation was supported by the observation of almost identical k_3 values for Li during the 1st stage leaching, compare 15.6×10^{-4} to 15.3×10^{-4} min⁻¹ (without and with US).

Fig. 6 shows the SEM micrographs of the blackmass residue after the leaching had been carried out, with or without ultrasound, for the leaching times; 1.5 h, 3 h, and 6 h. The corresponding average values of extracted metal ions (Mn, Co and Ni) are included as marked 'Ext. xx%' in the upper righthand corners of the micrographs. The morphology of the NMC particles was initially made up of larger ca. 10 µm spherical secondary particles shown in Fig. 6a (white arrows), which consisted of smaller aggregated primary particles of irregular sizes (Fig. 6b). The primary particles ranged in size from ca. 0.5-3.0 µm in diameter, and the boundaries between these

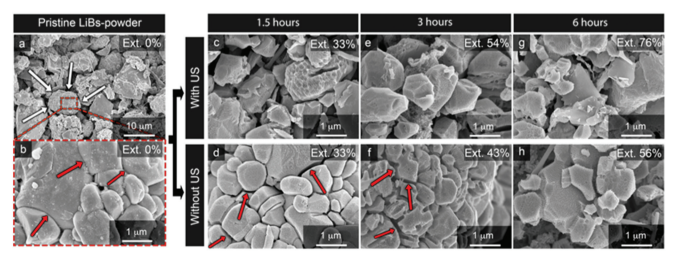


Fig. 6 Scanning electron micrographs of the pristine LiBs powder before the ultrasound extraction/leaching of battery metals (a and b), and after (c-h) citric acid leaching. The top row shows ultrasound assisted leaching while bottom row shows the oil bath leaching for different times, where c and d is 1.5 h, e and f 3 h, and g and h 6 h. The average values (displayed as inserts in each micrograph) of the extracted metal ions (excluding Li) were derived from Fig. 3.

Green Chemistry

particles are distinguishable as fused surfaces (red arrows) in Fig. 6b. The surface of the primary particles was smooth although covered by significantly smaller particles with approximate size of 40–100 nm (Fig. 6b). With ultrasound treatment during the leaching, the secondary particles comple-

although covered by significantly smaller particles with treatment during the leaching, the secondary particles completely fell apart into the primary particles after 1.5 h and all the adhesion between primary particles were lost (Fig. S7c†). Without ultrasound, the secondary particles still held together for longer time than 3 h, i.e. the positioning of the primary particles remained almost the same in the larger secondary particles, see Fig. 6d, f and Fig. S7d.† Accordingly, a significant portion of the NMC surfaces were inaccessible to the acid during the early hours of the leaching when performed in absence of US. The extended particle size decrease in the US system consequently resulted in enhanced residue layer diffusion for the ultrasound system, see eqn (8). It is noteworthy that the smaller 40-100 nm particles disappeared completely in both cases. It is suggested that the disappearance of these particles followed a similar dissolution behavior as the aluminum, see Fig. 5, which showed the best fitting to a linear dissolution behavior from 0 to 1440 min ($R^2 = 0.9982$, Table 4). Another observation was that the morphology of the particles exposed to ultrasound had developed more faceted edges, compare Fig. 6c and d. These facetted edges of the samples became most pronounced with longer extraction times (>6 h), while appearing only after 3 h with for the traditional extraction without ultrasound. Fig. 7a and b show high-resolution images of typical NMC primary particles with the layered grains after 3 h of extraction. The stacked sheet appearance was suggested to exist as a consequence of a more intense metal ion drainage in the surface of the particles, see arrow in Fig. 7b.

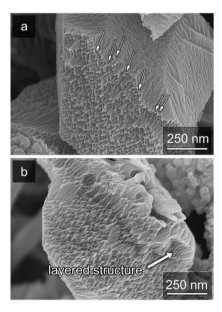


Fig. 7 Scanning electron micrographs of the primary particle of the layered NMC grains after 3 hours of extraction. Note the visible channel formations along different facets (a and b).

It should be noted that lithium-proton ion exchange dominated in the earlier stages of extraction. Thus H^+/H_3O^+ replaced Li^+ sites, leaving primarily a Ni/Mn/CoO₆ built layered structure (Fig. 7b) with less lithium present than represented by the crystal structure in Fig. 4b. This intercalation of H_3O^+ into the interlayers to some extent resulted in the shift of (003) to the lower 2 theta angle (Fig. 4a). It is suggested that although the morphological differences shown in Fig. 6 and 7 primarily related to initially breaking up the secondary particles, a more efficient metal ion drainage of the primary particles was always present when the ultrasound was used. This was evidenced by k_3 values remaining constant for leaching times >3 h up to 24 h (1440 min) as shown in Fig. 3.

3.4 The effect of temperature on the release order of metal ions

Fig. 8 shows the effect of temperature on the metal ion release from the LiBs powder for times from 0 to 1440 min in absence of ultrasound. The curves in Fig. 8 show that the leaching rate

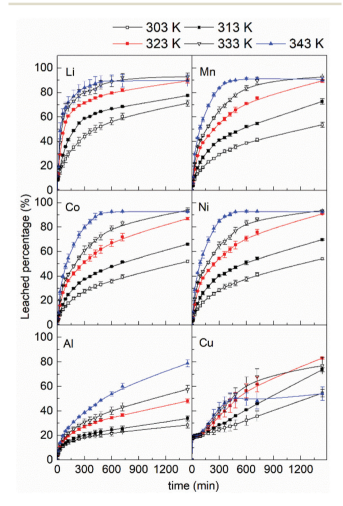


Fig. 8 Effect of temperature on leaching behaviour of metal ions (from the fine ground LiBs powder) depending on leaching time. The errors (error bars) were derived from triplicate leaching experiments. Conditions: citric acid, 1.5 mol L^{-1} ; S/L, 25 g L^{-1} , stirring speed, 500 rpm. The lines between measurement points serve as the guide to the eye.

Paper Green Chemistry

of all metals steadily increased with higher temperature (from 303 K to 343 K) during the first ca. 300 min and thereafter changed more moderately for all systems. The increase in leaching rate with increasing temperature was a consequence of the increasing dissociation constant of the citric acid,³⁶ since all metal ions showed the same behavior. The first dissociation constant of citric acid increased from $7.7 \times 10^{-4} \text{ s}^{-1}$ to $8.1 \times 10^{-4} \text{ s}^{-1}$ when the temperature was increased from 303 K to 323 K,36 leading to the formation of more available H⁺ ions and a faster chemical reaction. Moreover, a high temperature also increased the rate of mass diffusion for the released metal ions in the residue layer, which also contributed to the increased leaching rate, see section 3.3.

Table 5 summarize the leached percentages of all the metals at 1440 min. At 303 K only 71% Li, 53% Mn, 52% Co, and 54% Ni were released, while at 333 K ca. 20% more Li were released and about 40% more of the divalent Mn, Co, and Ni.

A further increase in temperature to 343 K resulted in more leached Al at 1440 min, whereas that of Li, Mn, Co, and Ni remained essentially the same and showed no additional leached amount of metal ions. The only exception from the general behavior that an increase in temperature favored more dissolved metal ions was for the copper (Cu). Here the leached percentage reached a maximum of 83% at 323 K and then gradually decreased to 54% when temperature was increased to 343 K. The explanation was argued related to the depletion of the dissolved O2 causing a supressed dissolution of Cu as described in section 3.2.

The activation energies (E_a) were calculated according to egn (9) to investigate if there was a correlation between the minimum energy required to release of Li, Ni, Co, and Mn and the total amount of leached metal ions, as well as rate of the release of metal ions in the NMC grain structure. The activation energies were derived for the oil bath system since the $E_{\rm a}$ values were required to be calculated for uniform temperature conditions, which could not be established for the US system. 15,16

$$K = Ae^{\frac{-E_a}{RT}} \tag{9}$$

Table 5 The leached percentage of the metal ions at 1440 min using citric acid as reagent the oil bath system depending on temperature

	Metal elements						
Leached percentage	Li	Mn	Со	Ni	Al	Cu	
303 K	71%	53%	52%	54%	28%	54%	
313 K	77%	72%	66%	70%	34%	73%	
323 K	89%	89%	87%	91%	48%	83%	
333 K	92%	92%	93%	93%	58%	77%	
343 K	90%	91%	93%	92%	78%	54%	

Comment: The data for each ion was acquired as repeated more than twice. The deviation was maximum ±3% with an exception of Cu (maximum ±7%).

where, K represents the kinetic constant (min-1) obtained from fitting slopes in Fig. S8,† A is the pre-exponential factor (\min^{-1}) , E_a is the apparent activation energy (J \mod^{-1}), R is the universal gas constant $(I \text{ mol}^{-1} \text{ K}^{-1})$, and T is the absolute temperature (K). By plotting $\ln K vs. 1/T$, the E_a values for leaching of Li (1st stage), Li (2nd stage), Mn, Co, and Ni were determined from the slopes of the corresponding fitted lines in Fig. 9.

Fig. 9 shows that the following E_a values could be established for the residue layer diffusion of the different ions: 67.9 (Li 1st stage), 39.6 (Li 2nd stage), 55.7 (Mn), 56.6 (Co), and 57.2 (Ni) KJ mol⁻¹, respectively. The activation energy for the 1st stage Li leaching was in fact significantly higher than other metal elements implying that a slower leaching rate should have existed although the opposite was observed. It should be highlighted here on the basis of the Li ion location in the NMC phase that the proton exchange process likely has a different reaction/diffusion path as compared to the other metal ions due to its location within the interlayer space of the NMC. This is in agreement with the structural data, wherein direct bonding holds Co/Mn/Ni in an oxide-rich coordination sphere, as shown in Fig. 4b. The 1st stage Li leaching was also reflecting contributions from both the electrolyte Li ion release process and the proton exchange process within the Ni/Co/MnO₆ layers (see Fig. 4b and Fig. 7). It was calculated that approximately 50% of the Li leaching connected to the 1st stage leaching was related to the electrolyte releasing process, see further section 3.1. It is noteworthy that while the electrolyte release process may have had a small contribution to the derived E_a value for the 1st Li leaching part, the residue layer diffusion for the first E_a value may have required an even higher activation energy due to the intense electrostatic attractions in the interlayer space of the NMC structure. On the contrary, during the 2nd stage of Li leaching, Li showed a diffusion rate similar to the Mn, Co and Ni metals ions (Table S3†), although the estimated E_a was lower, i.e. 39.6 kJ mol⁻¹. A

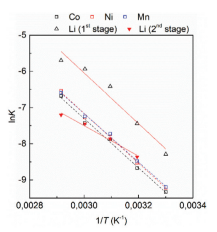


Fig. 9 Arrhenius plots for Li, Co, Ni, and Mn leaching using the k_3 values obtained from fitting slopes of residue layer diffusion control model. Conditions: citric acid, 1.5 mol L⁻¹; S/L, 25 g L⁻¹; stirring speed, 500 rpm.

Green Chemistry Paper

suggested explanation would be that the layered structure of the NMC started to dissociate at this stage as it was converted into the spinel structure (see Fig. 4c), thereby facilitating the release of Li. At this stage, the activation energy based on the surface residue layer diffusion was on the order of Ni > Co > Mn > Li (2nd stage), which shows that the binding between Nicitrate is the strongest while that of Li-citrate is the weakest among all the metal ions in NMC. A prevailing composition of Ni-citrate may thus exist in the residue layer interface surrounding the dissolving NMC grains.

The R^2 values for the Arrhenius plots of Cu and Al, shown in Fig. S10 and Table S4,† are significantly lower than the other elements. Since the Arrhenius equation assumes that the activation energy is independent on temperature, 37 this observation suggests that the leaching mechanisms of Al and Cu were temperature dependent to a larger degree than for the NMC-metals. The deviation from the Arrhenius equation can especially be expected for Cu, since the temperature directly affects the concentration of dissolved oxygen in the solution.

3.5 The impact of ultrasound depending on acid species

Table 6 summarize the total amounts of extracted metal ions for the two different acids over 24 h (1440 min).

The effect of the strength of the organic acid on the leaching behavior was investigated using a weaker organic acid, namely the monocarboxylic acetic acid (p $K_a = 4.76$). ²⁵ 4.5 mol L⁻¹ acetic acid was used to normalize the concentration for the reduced number of carboxylic groups. Fig. 10 shows the leaching behavior when using 4.5 mol L⁻¹ acetic acid and 1.5 mol L⁻¹ citric acid at 323 K. During the first 150 min, the leaching behaviors of the acids were almost similar, thereafter the leached percentage of Li, and especially Mn, Co, and Ni showed to be significantly higher for the citric acid, although the higher number of carboxylic groups had been compensated for with 3 times higher concentration of acetic acid. Thus, without any additional reducing agent, 1.5 M citric acid (pH \approx 1.14) showed better transition metal (Ni, Mn, Co) leaching performance compared to the 4.5 M acetic acid (pH \approx 1.74). The leached percentage of Cu in 1.5 mol L^{-1} citric acid was, however, lower than in 4.5 mol L⁻¹ acetic acid during the entire course of the reaction. This may be explained as a result from that citric acid has been reported as an efficient chelating agent to suppress the Cu dissolution in acid.³⁸

The leaching behavior with/without ultrasound using acetic acid was also investigated. As shown in Fig. S11,† the leached

Table 6 The leached percentage of the metal elements at 1440 min using 1.5 mol L⁻¹ citric acid and 4.5 mol L⁻¹ acetic acid as leaching agent in oil bath at 323 K

	Metal elements					
Leached percentage	Li	Ni	Со	Mn	Al	Cu
1.5 mol L^{-1} citric acid 4.5 mol L^{-1} acetic acid	89% 83%	89% 75%	87% 72%	91% 73%	48% 53%	83% 86%

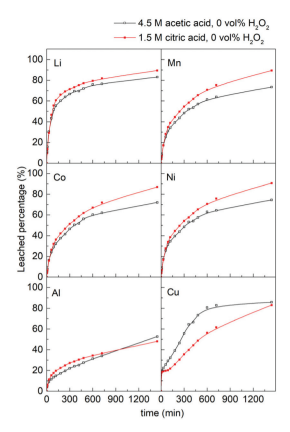


Fig. 10 Effect of organic acid species on extraction of metal ions from the fine ground LiBs powder in oil bath at 323 K. Conditions: citric acid, 1.5 mol L^{-1} ; acetic acid, 4.5 mol L^{-1} ; 0 vol% H_2O_2 ; S/L, 25 g L^{-1} ; stirring speed, 500 rpm. The lines between measurement points serve as the guide to the eye.

percentage of the targeted metal ions were significantly improved with ultrasound also for the acetic acid, reaching ca. 96% \pm 2% (Table 7). The result further highlighted the impact of ultrasound in improving the extraction of metals from the blackmass, irrespectively of the acid species. This might be due to the formation of hydroxyl radicals in the US leaching system, possibly allowing continuous formation of hydrogen peroxides.39 Fig. S12† shows the overlapping nature of the citric and the acetic acid during the entire leaching cycles, suggesting that the primary difference between the acids was their leaching rate (speed) under the influence of the ultrasound.

Table 7 The leached percentage of the metal ions at 1440 min using 4.5 mol L⁻¹ acetic acid as leaching agent in ultrasound (US) leaching system and oil bath at 323 K

	Metal elements					
Leached percentage	Li	Ni	Со	Mn	Al	Cu
Oil bath, 323 K US, 323 K	83% 94%	75% 97%	72% 97%	73% 95%	53% 60%	86% 73%

Paper

The E_a values for leaching of Li (1st stage), Li (2nd stage), Mn, Co, and Ni were 70.1, 41.5, 44.4, 45.1, and 44.1 kJ mol⁻¹, respectively (see Fig. S13, S14, S15, and Tables S5, S6† for their derivation). The activation energies for the citric and the acetic acid were thus similar for the Li (1st stage) and Li (2nd stage), while for Mn, Co, and Ni the E_a values were more than 10 kJ mol⁻¹ lower for the acetic acid as compared to the citric acid. The reason to the lower activation energy for the divalent metal ions when using acetic acid is presently unknown. It was speculated being a consequence of a less restricted diffusion through the residue layers on the surface of the NMC grains. This was based on the presumption that the complex formation of the extracted divalent metal ions (Mn, Co, and Ni) and the citric acid likely formed stronger complexes by directing 2 oxygen rich carboxylic groups of the acid towards the positively charged metal ions, while leaving one carboxyl group facing outwards towards the bulk acid solution. This configuration stands in contrast to the nonpolar methyl group facing outwards in the case of the acetic acid when its carboxyl

The viscosities of the aqueous solutions of 4.5 mol L^{-1} acetic acid and 1.5 mol L^{-1} citric acid, before and after the 24 h leaching process, were taken as an indicative measure of this possible complex binding and coordination of the divalent metal ions (Table 8). The viscosity of the aqueous solution of the 1.5 mol L^{-1} citric acid was significantly higher than that of the 4.5 mol L^{-1} acetic acid. The differences in viscosity before and after (Δ) leaching of the 1.5 mol L^{-1} citric acid solution was: $\Delta = 0.33$ mm² s⁻¹, which in turn also was higher than that of 4.5 mol L^{-1} acetic acid solution: $\Delta = 0.17$ mm² s⁻¹. The higher viscosity of the citric acid system was suggested to reflect the stronger metal–organic ion complex binding. This would be in agreement with the higher activation energy required for the diffusion of transition metal ions through the residue layer in the 1.5 mol L^{-1} citric acid.

3.6 The effect of H₂O₂ concentration

group in turned towards the metal ions.

Hydrogen peroxide is frequently used to reduce high-valence value metal ions ($\mathrm{Co^{3^+}}$, $\mathrm{Mn^{3^+/4^+}}$) to lower valence values ($\mathrm{Co^{2^+}}$, $\mathrm{Mn^{2^+}}$), making them more soluble and facilitating their release from the NMC structure during the leaching. ⁴⁰ The $\mathrm{H_2O_2}$ concentration effect was therefore evaluated. Fig. S16† shows the leached percentage of metal ions over the 1440 min leaching

Table 8 The viscosity of the 1.5 mol L^{-1} citric acid and 4.5 mol L^{-1} acetic acid before and after 1440 min leaching treatment without adding reducing agents

	1.5 mol L	⁻¹ citric	4.5 mol L	4.5 mol L ⁻¹ acetic acid		
Acid species	Before	After	Before	After		
Viscosity (mm ² s ⁻¹)	1.67	2.00	1.33	1.50		

Comment: The data for each viscosity was acquired as repeated five times measurements at 298 K. The deviation was maximum $\pm 0.03\%$.

for different H₂O₂ concentrations from 0 to 2.0 vol%. The leaching rate of all metals were increased with an increase in the H₂O₂ concentration during the first 150 min, except for the Al ions. After 150 min, the leaching rate changed more moderately with increasing H₂O₂ concentration. Fig. S16† shows that no improvement in total amount of leached metals occurred when increasing the H₂O₂ concentration from 0 to 1.0 vol%. More than 1.0 vol% H₂O₂ resulted in a decreased in the leached percentages for all metals at 1440 min, with only ca. 80% being leached for 2.0 vol% of H₂O₂. This was caused by the precipitation of the copper ions, occurring with extensive addition of H2O2. The X-ray diffraction of the residues after 1440 min leaching with different H₂O₂ concentrations from 0 to 2.0 vol% confirmed that the peaks attributed to NMC disappeared when the H₂O₂ concentration reached 1.5 vol%, while the peaks attributed to Cu(OH)2 phase (ca. 22.8°) became apparent when the H₂O₂ concentration exceeded 1.5 vol%, see Fig. S17.† The explanation to the decrease in total amount of leached metal ions (for 2.0 vol% of H₂O₂) was therefore suggested to have occurred as a result of the Cu(OH)2 precipitating along with possible coprecipitates of Li, Mn, Ni, and Co. The formation of the Cu(OH)₂ (s) might be due to the fact that H2O2 functions as an oxidizing agent, improving the dissolution of Cu in acid solutions. 41 The quick dissolution of solid-phase Cu/CuO_x to Cu²⁺ is accompanied by a large consumption of protons, resulting in an increased concentration of OH at the solid/liquid interface. The sudden increase in pH at the interface eventually leads to Cu²⁺ precipitating in the form of Cu(OH)2. Cu(OH)2 has the lowest solubility constant $(K_{\rm sp})$, $\log(K_{\rm sp}) = -19.32$, ⁴² apart from the aluminium hydroxide with $log(K_{sp}) = -33.5$, ⁴² however, citric acid has been reported to prevent precipitation of Al(OH)3 when the citric acid/aluminium ratio is higher than 0.843 (which was the case in our work).

Fig. S18† shows the curves in Fig. S16† as fitted to the residue layer diffusion contribution of the shrinking core model. The leaching processes of all NMC metal elements could be divided into two stages in the presence of $\rm H_2O_2$, with the corresponding k_3 values being summarized in Table S7.† When 1.5 vol% of $\rm H_2O_2$ was added, the k_3 values of the transition metal elements (Ni, Mn, and Co) were ca. 2–3 times higher during the 1st stage (initial 150 min) than when no $\rm H_2O_2$ was added. The increased rate constants were synonymous with a faster release of the structurally bonded metal ions. The increase of the k_3 value in the 1st stage was attributed to:

- The H_2O_2 reducing Co and Mn from high valence values to low valence ones, which improved the ability of acid to solvate the metal ions,
- The decomposition of the H_2O_2 resulting in the formation of bubbles, which improves the mixing of the solution, contributing to a facilitated mass transfer and a higher leaching rate.

After 150 min (the 2^{nd} stage), the k_3 values decreased and became similar, regardless of H_2O_2 concentration. It was suggested that H_2O_2 mainly worked during the beginning of the leaching process due to its depletion. Overall, it was con-

cluded that the addition of H_2O_2 enhanced the leaching rate of transition metal elements, whereas the total amount of metal ions decreased at excessively high H_2O_2 concentrations due to the formation of intermediate hydroxide phases (e.g., $Cu(OH)_2$).

4 Conclusions

Extraction of lithium-ion battery metals from blackmass (leaching) was carried out using mild organic acids (citric and acetic) in the presence of ultrasound. The ultrasound significantly impacted the leaching, resulting in, on average 97% ± 3% metal ion recovery, which was a substantially higher amount of recovered metal ions than for the same conditions when only mechanical stirring was used (89% ± 2%). The highest recoveries were achieved for cobalt and nickel, reaching >99%, while lithium and manganese were recovered at 94-96% efficiency. In addition, a facilitated dissolution of the spinel LiMn₂O₄ (formed during the leaching) was observed. The effect of temperature and leaching time was investigated (correlated with ICP-OES measurements) for the Li, Mn, Co, and Ni metal ions. It was shown that when ultrasound was used only half of the leaching time (≤600 min) was required to reach the same percentage of target metals (ca. 90%) as the mechanically stirred leaching procedure reached in 1440 min (24 h). The time and temperature data for different metal ions were fitted to the shrinking core model. The results revealed that the leaching of Li, Ni, Mn, Co, and Al (from the fine ground LiBs powder) was controlled by residue layer diffusion, and could be associated with the residue layer diffusion contribution of the shrinking core model. The ultrasound contributing to an improved residue layer mass diffusion. The immediate effect of the ultrasound was to facilitate the liberation of primary NMC particles (within 90 min) from larger-sized secondary particles, as demonstrated by microscopy. However, although the ultrasound provided better acid access to the surface of the aggregated, smaller primary NMC particles, the leaching rate remained markedly higher over the entire 24 h (1440 min) extraction process. The only exception could be established for Cu, which was controlled by the surface chemical reaction contribution of the model. This was suggested to be caused by the limited amount of dissolved O₂ in the solution, which resulted from the ultrasonic degassing of the acidic extraction solutions. Using citric acid, as compared to acetic acid, further suppressed the dissolution of Cu due to the chelating effect of citrate ions with the surface of Cu. The addition of H₂O₂ enhanced the leaching rate of Ni, Mn, Co, and Li, especially in the first 150 min, but showed less improvement or even decreased the maximum leached percentage of metal ions due to the co-precipitation with Cu²⁺ ions. The leached percentage using an even weaker organic acid (acetic acid, pK_a : 4.76) demonstrated that also for this acid ultrasound had a marked effect with extraction values reaching 96% \pm 2% with ultrasound, while the value was 77% \pm 5% without ultrasound. It could also be concluded that the ultrasound had a greater impact on the leaching results than that of the use of either citric or acetic acid. Overall, the technique shortened leaching time, eliminated reducing agents, and omitted reported pre-treatment procedures (e.g., binder remover chemicals, and pre-calcination above 600 °C). It is therefore suggested that ultrasound provides a promising route towards energy-efficient and more environment-friendly extraction of targeted metals in discarded lithium-ion batteries. Finally, it was concluded that the blackmass composition strongly depended on the method of powder preparation, highlighting the importance of using fine ground blackmass to avoid challenges in the interpretation of leaching data due to natural fractionation from transport and storage.

Conflicts of interest

There are no conflicts to declare.

Acknowledgements

This article was carried out within the PERLI (Processes for Efficient Recycling of Lithium-ion Batteries) project 48228-1 granted by the Swedish Energy Agency. Assoc. Prof. M. Petranikova at Chalmers University of Technology is acknowledged for establishing the contact with Volvo Cars and providing the Volvo C30 blackmass powder under energy authority grant no: 48204-1.

Notes and references

- 1 T. Or, S. W. D. Gourley, K. Kaliyappan, A. Yu and Z. Chen, *Carbon Energy*, 2020, 2, 6–43.
- 2 J. C. Kelly, Q. Dai and M. Wang, *Mitigation Adapt. Strategies Global Change*, 2019, 371–396.
- 3 L. Dahllöf, M. Romare and A. Wu, *Mapping of lithium-ion batteries for vehicles*, Nordic Council of Ministers/Publication Unit, Denmark, 2019.
- 4 D. L. Thompson, J. M. Hartley, S. M. Lambert, M. Shiref, G. D. J. Harper, E. Kendrick, P. Anderson, K. S. Ryder, L. Gaines and A. P. Abbott, *Green Chem.*, 2020, 22, 7585–7603.
- 5 E. Melin, State-of-the-art in reuse and recycling of lithium-ion batteries A research review, The Swedish Energy Agency, Eskilstuna, 2019.
- 6 X. Zheng, Z. Zhu, X. Lin, Y. Zhang, Y. He, H. Cao and Z. Sun, *Engineering*, 2018, 4, 361–370.
- 7 J. Diekmann, C. Hanisch, L. Froböse, G. Schälicke, T. Loellhoeffel, A.-S. Fölster and A. Kwade, *J. Electrochem. Soc.*, 2017, 164, A6184–A6191.
- 8 Y. Ma, M. Svärd, X. Xiao, J. M. Gardner, R. T. Olsson and K. Forsberg, *Metals*, 2020, **10**, 1–16.
- 9 C. Lupi and M. Pasquali, *Miner. Eng.*, 2003, **16**, 537–542.
- 10 J. Zhang, J. Hu, W. Zhang, Y. Chen and C. Wang, J. Cleaner Prod., 2018, 204, 437–446.

Paper

- 11 P. Liu, L. Xiao, Y. Chen, Y. Tang, J. Wu and H. Chen, *J. Alloys Compd.*, 2019, **783**, 743–752.
- 12 R. Golmohammadzadeh, F. Rashchi and E. Vahidi, *Waste Manage.*, 2017, **64**, 244–254.
- 13 L. Li, L. Zhai, X. Zhang, J. Lu, R. Chen, F. Wu and K. Amine, J. Power Sources, 2014, 262, 380–385.
- 14 L. Li, E. Fan, Y. Guan, X. Zhang, Q. Xue, L. Wei, F. Wu and R. Chen, ACS Sustainable Chem. Eng., 2017, 5, 5224–5233.
- 15 C. E. Brennen, Cavitation and bubble dynamics, 2013.
- 16 E. Ciawi, J. Rae, M. Ashokkumar and F. Grieser, J. Phys. Chem. B, 2006, 110, 13656–13660.
- 17 W. Y. Wang, C. H. Yen and J. K. Hsu, Sep. Sci. Technol., 2020, 55, 3028-3035.
- 18 Y. J. Shih, S. K. Chien, S. R. Jhang and Y. C. Lin, J. Taiwan Inst. Chem. Eng., 2019, 100, 151–159.
- 19 W. Gao, X. Zhang, X. Zheng, X. Lin, H. Cao, Y. Zhang and Z. Sun, *Environ. Sci. Technol.*, 2017, 51, 1662–1669.
- 20 W. Gao, J. Song, H. Cao, X. Lin, X. Zhang, X. Zheng, Y. Zhang and Z. Sun, *J. Cleaner Prod.*, 2018, **178**, 833–845.
- 21 X. Chen, B. Fan, L. Xu, T. Zhou and J. Kong, J. Cleaner Prod., 2016, 112, 3562–3570.
- 22 C. Lei, I. Aldous, J. M. Hartley, D. L. Thompson, S. Scott, R. Hanson, P. A. Anderson, E. Kendrick, R. Sommerville, K. S. Ryder and A. P. Abbott, *Green Chem.*, 2021, 4710–4715.
- 23 A. E. Danks, S. R. Hall and Z. Schnepp, *Mater. Horiz.*, 2016, 3, 91–112.
- 24 H. Poerwono, K. Higashiyama, H. Kubo, A. T. Poernomo, S. Suharjono, I. K. Sudiana, G. Indrayanto and H. G. Brittain, *Citric Acid*, Elsevier Masson SAS, 2001, vol. 28.
- 25 D. D. Perrin, B. Dempsey and E. P. Serjeant, pK_a Prediction for Organic Acids and Bases, 1981.
- 26 X. Chen, C. Luo, J. Zhang, J. Kong and T. Zhou, ACS Sustainable Chem. Eng., 2015, 3, 3104–3113.

- 27 W. J. Wolfgong, Handb. Mater. Fail. Anal. with Case Stud. from Aerosp. Automot. Ind, 2016, pp. 279–307.
- 28 X. Xiao, F. Hayashi, H. Shiiba, S. Selcuk, K. Ishihara, K. Namiki, L. Shao, H. Nishikiori, A. Selloni and K. Teshima, J. Phys. Chem. C, 2016, 120, 11984–11992.
- 29 Q. H. Zhang, S. P. Li, S. Y. Sun, X. S. Yin and J. G. Yu, Chem. Eng. Sci., 2010, 65, 169–173.
- 30 K. Momma and F. Izumi, J. Appl. Crystallogr., 2011, 44, 1272–1276.
- 31 H. Nord, Acta Chem. Scand., 1955, 9, 430-437.
- 32 D. G. Eskin, *Ultrasonic degassing of liquids*, Elsevier Ltd, Uxbridge, UK, 2015.
- 33 A. R. Kaiser, C. A. Cain, E. Y. Hwang, J. B. Fowlkes and R. J. Jeffers, J. Acoust. Soc. Am., 1996, 99, 3857–3859.
- 34 D. Tromans, Hydrometallurgy, 1998, 48, 327-342.
- 35 P. R. Taylor, M. de Matos and G. P. Martins, *Metall. Trans. B*, 1983, **14**, 49–53.
- 36 R. G. Bates and G. D. Pinching, J. Am. Chem. Soc., 1949, 71, 1274–1283.
- 37 K. J. Laidler, J. Chem. Educ., 1984, 61, 494-498.
- 38 T. C. Hu, S. Y. Chiu, B. T. Dai, M. S. Tsai, I. C. Tung and M. S. Feng, *Mater. Chem. Phys.*, 1999, 61, 169–171.
- 39 K. Makino, M. M. Mossoba and P. Riesz, *J. Phys. Chem.*, 1983, **87**, 1369–1377.
- 40 L. P. He, S. Y. Sun, X. F. Song and J. G. Yu, *Waste Manage.*, 2017, **64**, 171–181.
- 41 J. C. Chen and W. T. Tsai, *Mater. Chem. Phys.*, 2004, **87**, 387–393.
- 42 R. Smith and A. Martell, *Critical Stability Constants Volume*4: Inorganic Complexes, 1976, vol. 4.
- 43 D. M. Dabbs, U. Ramachandran, S. Lu, J. Liu, L. Q. Wang and I. A. Aksay, *Langmuir*, 2005, **21**, 11690–11695.