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DOI:

[10.1016/j.jclepro.2020.124058](https://doi.org/10.1016/j.jclepro.2020.124058)

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Document Version

Peer reviewed version

Citation for published version (Harvard):

Jonsson, C, Awais, M, Pickering, L, Degri, M, Zhou, W, Bradshaw, A, Sheridan, R, Mann, V & Walton, A 2020, 'The extraction of NdFeB magnets from automotive scrap rotors using hydrogen', *Journal of Cleaner Production*, vol. 277, 124058. <https://doi.org/10.1016/j.jclepro.2020.124058>

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The extraction of NdFeB magnets from automotive scrap rotors using hydrogen

Christian Jönsson, Muhammad Awais*, Lydia Pickering, Malik Degri, Wei Zhou, Andy Bradshaw, Richard Sheridan, Vicky Mann, and Allan Walton

School of Metallurgy and Materials, University of Birmingham, B15 2TT Edgbaston, UK

*Corresponding author: M.Awais@bham.ac.uk

Abstract

Scrap containing NdFeB is a valuable resource for the production of NdFeB magnets as the demand for these materials grows. One of the challenges is to recover the rare earths or the NdFeB alloy powder in a clean and cost effective manner so that it can be re-processed into new magnets rather than becoming lost to landfill. Work on using hydrogen to process scrap magnets (HPMS) has been shown to be successful when targeting hard disk drives. Currently, there is a lack of information on reliable methods to separate out NdFeB from other scrap sources such as automotive drives. In the near future, with increasing sales and electrification of cars, the automotive sector could be an important source for Dy containing magnets. In this paper, the hydrogen processing of scrap magnets has been demonstrated as an extraction method for NdFeB from automotive rotors for the first time, with the aim to examine the viability of this recycling process and learn lessons for design for recycling. Thus leading to the sustainable production of these components. Significant challenges were outlined when applying the hydrogen process to rotors with embedded magnets. After the extraction, further process steps may also be needed to separate epoxy coatings, as sieving could only reduce the carbon content to 1420 ppm, compared to 770 ppm in the base alloy. The gravimetric measurements also confirmed that Dy additions increase both the initiation and absorption time for hydrogen decrepitation. Hence, a higher hydrogen pressure will be required to speed up the process.

Highlights:

- Hydrogen can be used to extract NdFeB magnets from the automotive rotors, however, many challenges are presented due to rotors being designed without recycling in mind.
- The importance of design choices that have a negative impact on the recycling of automotive rotors are investigated in this work, so can be avoided in future designs.
- Higher Dy additions slow down the kinetics of the hydrogen decrepitation reaction by 75 % at room temperature and 1.2 bar hydrogen pressure.

- Scrap magnets with 8.5 wt% Dy took longer to initiate the reaction and a significantly higher hydrogen pressure is required for quick processing of the scrap.
- The carbon content could only be reduced to 1420 ppm, compared to 770 ppm in the starting material, by sieving the extracted powder to <45 μm because the epoxy coating is breaking up at a similar rate to the HD powder.

Keywords

Recycling, hydrogen decrepitation, automotive, NdFeB, critical materials

1. Introduction

The rare earth elements (REEs) are defined as critical materials in the EU and the US, due to their high economic importance combined with their supply risk (European Commission, 2017, 2014; U.S. Department of Energy, 2011). The rare earth market is, to a large extent, driven by the increasing use of neodymium-iron-boron (NdFeB) magnets, which are employed in electronics, industrial and automotive motors, and wind turbine generators. Recycling of end-of-life NdFeB magnets could be part of the solution to alleviate the supply risks for REEs outside of China. At present, however, less than 1 % of REEs are recycled, due to a lack of incentives, inefficient collection, and technological difficulties (Reck and Graedel, 2012). In a study, Habib et al. revealed that primary supply would be unable to meet the forecasted demand of Nd and Dy by 2050 and recycling is unlikely to close the wide gap between future demand and supply by 2050. This means that a highly accelerated rate of Nd and Dy mining will be required to meet the demand. However, it has been suggested that secondary supply from recycling can meet almost 50 % of the demand in the long term i.e. by 2100 (Habib and Wenzel, 2014). Another study estimates that the recycling of EOL magnets could supply 68 % of light REE (Nd and Pr) and 11 % of heavy REE (Dy and Tb) demand of NdFeB magnets by 2030 (Schulze and Buchert, 2016).

Automotive scrap has the potential to contain a significant amount of NdFeB typically with increased Dy content. The amount of REEs present in fully electric vehicles varies with the type/model of vehicle. Typically, an EV contains 4-5 kg of REE compared to less than a kilo in a new petrol/diesel car (Alonso et al., 2012). To get a bigger perspective on the potential of recycling of these REE from the electric vehicles, IEA estimated that the EV sales reach would reach 44 million vehicles per year by 2030 (IEA, 2019). If these magnets are recycled efficiently, it can create a parallel supply chain to the primary route. Thus, becoming an important source to meet future demand as highlighted by other researchers (Alonso et al., 2012; Du and Graedel, 2011; Habib and Wenzel, 2014). The key, however, is accessing this material in a cost effective and efficient manner. Work is being carried out to address the issue

of the low rates of recycling of these critical materials. The creation of a systematic labelling system that encompasses the key metrics for recycling can enable end users of magnetic parts to select more environmentally friendly designs and can also ensure that the magnet containing products can end up at the correct recycling facility (MaXycle, 2018; Burkhardt; et al., 2019; Burkhardt; et al., 2020).

Despite a huge surge in the demand for electric vehicles and their recycling potential, there is not much literature available looking at the viability of recycling NdFeB containing scrap from automotive sources. Therefore, this study is aimed to identify the challenges involved to recover NdFeB magnets from end of life automotive scrap. The work also investigates the viability of recycling automotive rotors using hydrogen and provide feedback for design for recycling.

2. Literature Review

Extraction of NdFeB magnets from end of life vehicles is very labour intensive work. It has been shown to be profitable to dismantle the rotor manually due to the amount of Cu in the windings (Bast et al., 2014). Previously, Walton et al. (2015) presented a recycling process, HPMS (Hydrogen Processing of Magnet Scrap), which uses hydrogen to extract NdFeB magnets from the hard disk drives (HDDs) in the form of demagnetised, hydrogenated alloy powder. The VCM-containing corner was cropped off to concentrate the magnet-containing feed and provide a route out for the NdFeB powder. The cropped corners are distorted to fracture the magnets prior to hydrogen exposure for activation. The corners are then loaded into a porous drum inside the hydrogen reactor, which is subsequently tumbled to liberate the powder from the assemblies. Milling and sieving is performed to remove impurities, especially the Ni coating, before the alloy powder can be reprocessed, for example by sintering into new fully dense magnets. Life Cycle Analysis (LCA) has shown that by producing re-sintered magnets using HPMS, energy usage can be reduced by 88 % compared with primary production (Sprecher et al., 2014)

Extracting NdFeB magnets from automotive rotors comes with a different set of challenges compared to that of HDDs. The removal of the motor from a car and then the magnet-bearing rotor is labour intensive work. Moreover, automotive rotors come in a wide range of designs, using different materials, and with the magnets placed in various positions, making it difficult to find a universal method for NdFeB extraction. However, the MORE (MOtor REcycling) project stated that stripping down to rotor/stator level is profitable, even with German labour costs, due to the recovery of Cu from the stator windings (Bast et al., 2014). This did not include the potential value of the recovered NdFeB. One method was developed to punch out

buried magnets for reuse and another was to shear off surface mounted magnets. The undamaged magnets could then be reused after removal of the adhesive residues.

Automotive scrap has a potentially higher value due to the increase in Dy content. The increase in Dy is required to improve performance at typical operating temperatures (Gauder et al., 1988). The two main phases in the NdFeB type magnet microstructure are rare earth (RE) rich, an alloy predominantly rich in Nd but will contain other rare earths such as Dy. The matrix phase consists of $\text{Nd}_2\text{Fe}_{14}\text{B}$, this is the magnetic phase. The Fe can be substituted for Co and the Nd for other rare earths (Sagawa et al., 1985).

During the hydrogen decrepitation reaction, the formation of rare earth hydrides at the grain boundary generates exothermic heat that initiates hydrogen diffusion into interstitial sites of the $\text{Nd}_2\text{Fe}_{14}\text{B}$ matrix phase (Harris et al., 1987). Regarding Dy, it has been shown that, at 20bar and room temperature, it takes longer to form a hydride from Dy than from Nd (Hirano et al., 2006) and that the $\text{Dy}_2\text{Fe}_{14}\text{B}$ phase can absorb less hydrogen than its Nd counterpart (Bartolomé et al., 1991). However, Saje et al., (1992) did not see a difference in reaction rate when performing hydrogen decrepitation (1 bar, 20 °C) of the $\text{Nd}_{16-x}\text{Dy}_x\text{Fe}_{76}\text{B}_8$ (at.%, $x = 0-3$) alloy with varying Dy content. There is some disagreement as to whether the Dy content of the magnets may affect the hydrogen decrepitation of sintered NdFeB (which the HPMS process is based around), where the grain boundary phase plays a key role for the initiation and propagation of the reaction.

After the hydrogen decrepitation, the hydrogenated NdFeB powder needs to be purified to remove all the impurities. This purification process has to be done in an inert atmosphere to avoid oxidation. Various authors have noted that both oxygen and carbon have been shown to have detrimental effects on the sintering process for NdFeB (Lopes et al., 2012; Sasaki et al., 2015; Zakotnik et al., 2009). After this, the powder can be inertly milled to single crystal using the jet mill and sintered to new magnets by following the commercial production route (Malik Degri, 2014).

3. Experimental Design

To look at the feasibility of extraction of hydrogenated NdFeB from magnetic rotors from automotive scrap, the process was broken down into two separate stages. The first stage, to look at the feasibility of using hydrogen processing of magnet scrap (HPMS) applied to automotive rotors and whether pre-processing of the rotors is required to get the magnets to

successfully react with hydrogen at room temperature and 1.5 bar. The second stage was to study the kinetics of the reaction hydrogen with the magnets of different Dy content to assess the influence of Dy on hydrogen decrepitation.

3.1. Extraction from Rotors

Pre-separated discarded rotors, from an unnamed motor company, were used in this study, see Figure 1. The grade was determined to be N42SH and the composition of these magnets was measured to be $\text{Nd}_{9.08}\text{Pr}_{2.62}\text{Dy}_{1.38}\text{Fe}_{78.99}\text{Al}_{1.06}\text{Nb}_{0.17}\text{B}_{5.81}$ (atomic %) and traces of Gd 0.04 at%, Co 0.71 at%, Cu 0.13 at%, Mn 320 ppm, Si 410 ppm and N 170 ppm. The oxygen and carbon contents were 760 ppm and 690 ppm respectively. These rotors contain ten surface-mounted epoxy-coated magnets on a ferritic steel disk with a diameter of 25 cm. The magnets were either non-, semi- or fully embedded in the thermosetting plastic.

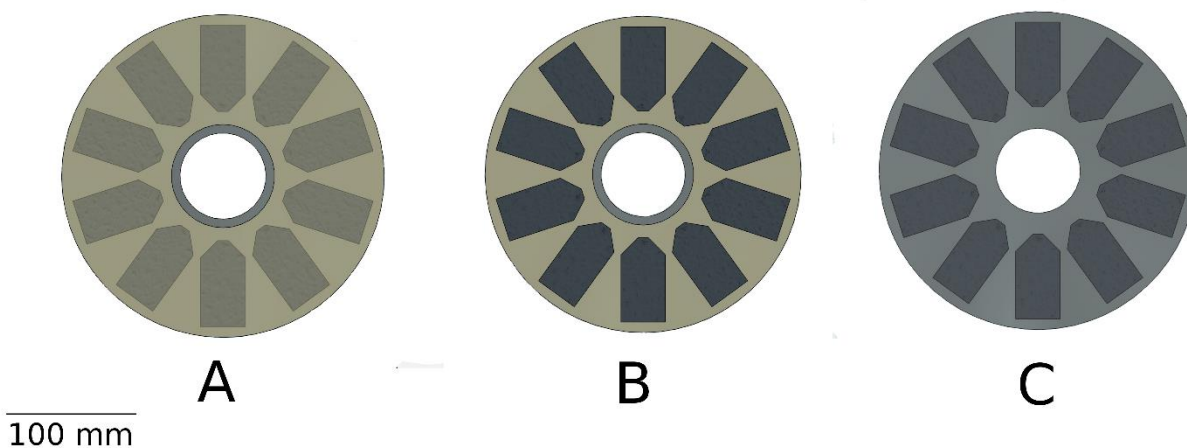


Figure 1: A model of the automotive rotors studied in this work, all containing epoxy-coated magnet slots on a ferritic steel plate. The magnet slots are either a) fully embedded, b) semi-embedded or c) non-embedded in a thermosetting plastic.

The HPMS processing was carried out using a 300 L reaction chamber, shown in Figure 2. The experiments started by loading rotors from the top of the vessel, followed by argon flushing and evacuation to 10^{-2} mbar. Hydrogen was then introduced to a pressure of 1.2-1.5 bar in order to initiate the hydrogen decrepitation reaction and the pressure within the vessel was maintained manually throughout the experiment until the pressure had been stable for several hours. All processing was performed at room temperature and the hydrogen was removed at the end of the reaction. The recycling rig provided the option of connecting the argon inlet to a pneumatic spinner that was connected to the assembly holder. As argon is fed, the spinner agitated the assemblies, liberating the NdFeB powder, which would fall into

the collection pot. The collection pot was then be sealed by the closing of two ball valves and transferred inertly to an argon-filled glovebox with oxygen level <10 ppm.

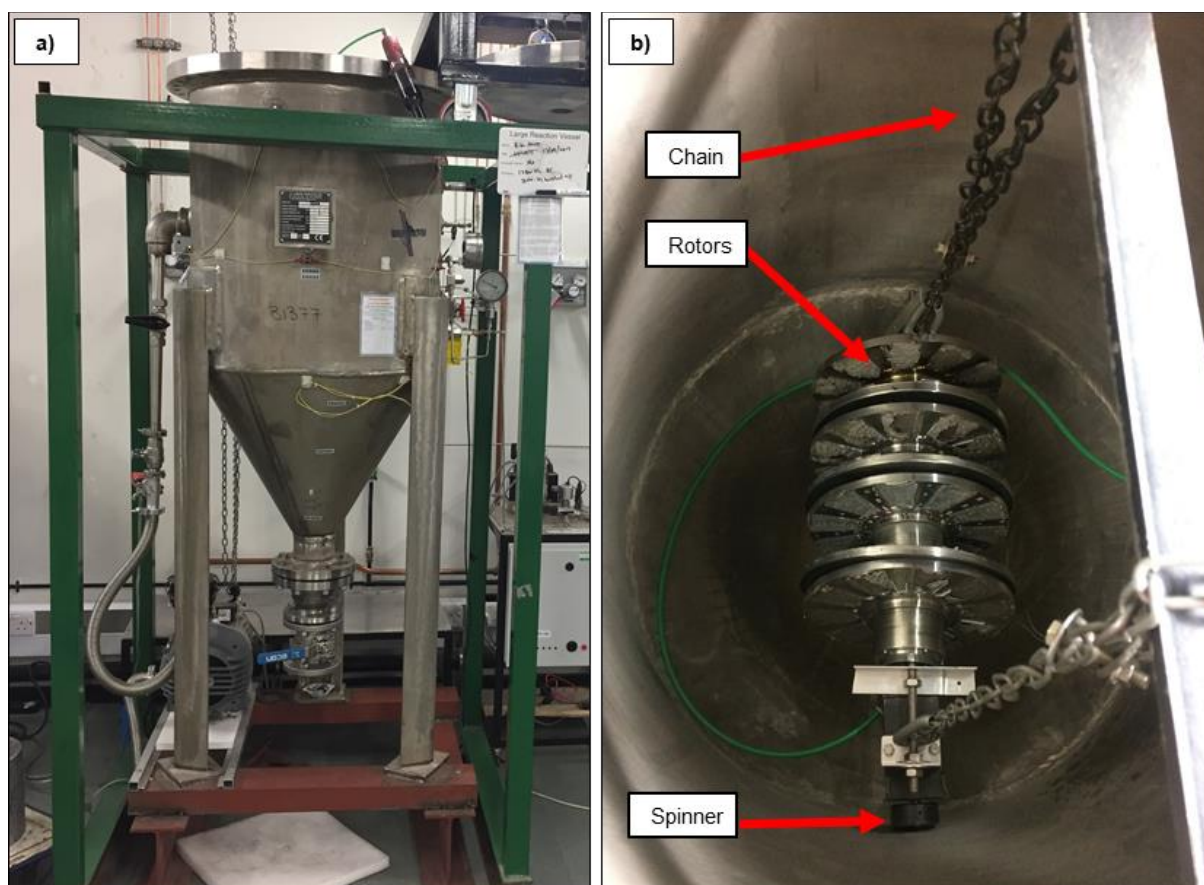


Figure 2: The 300 L reaction vessel used for HPMS processing, shown from the outside (a), and from the inside (b) with the rotors loaded.

The sieving of the extracted powder was performed in 250 g batch sizes in an argon-filled glovebox using a sieve stage with 2000 μm , 500 μm , 250 μm , 150 μm , 90 μm , and 45 μm mesh size sieves (25 cm diameter) for 30 minutes. ICP-OES and carbon analysis was carried out on the fraction of material that was remaining on every sieve and the fraction that had passed through all of the sieves to identify any separation and removal of the epoxy coating.

3.2. Gravimetric Measurements of Hydrogen Absorption

Gravimetric measurements of the hydrogen absorption were carried out on uncoated sintered magnets with compositions $\text{Nd}_{31.1-x}\text{Dy}_x\text{Pr}_{0.60}\text{Fe}_{64.78}\text{Co}_{3.00}\text{Ga}_{0.22}\text{Cu}_{0.15}\text{Al}_{0.15}$ ($x=1.00$ (composition A), $x=6.80$ (B), $x=8.45$ wt.% (C), supplied by Magneti Ljubljana). These magnets were chosen as they only differ in the Nd/Dy ratio so the effect of Dy could be investigated. The samples were cut into cylinders of 3.65 mm diameter using an electrical discharge machine and ground

to 152 mg under an argon atmosphere where they were also stored. The samples were quickly ground in air on each flat side before the measurements, to get the weight down to 150 g and to keep the surface conditions constant. The gravimetric measurements of the hydrogen decrepitation process were performed using an Intelligent Gravimetric Analyser (IGA-001) from Hiden Isochema Ltd. The microbalance of the IGA has long-term stability of $\pm 1 \mu\text{g}$, a weighing resolution of 0.2 g, and pressure resolution of $\pm 0.02 \%$. After loading the samples, the system was left open to air for a set time for air exposure (5, 30, and 60 minutes) to determine the effect of oxidation after preparation of the rotors for the HPMS process and then evacuated to 10^{-4} mbar pressure. Hydrogen was introduced at a rate of 200 mbar/minute to the set pressure (1.2, 2.5, 5.0 or 10.0 bar), where it was held constant.

3.3. Analytical Methods

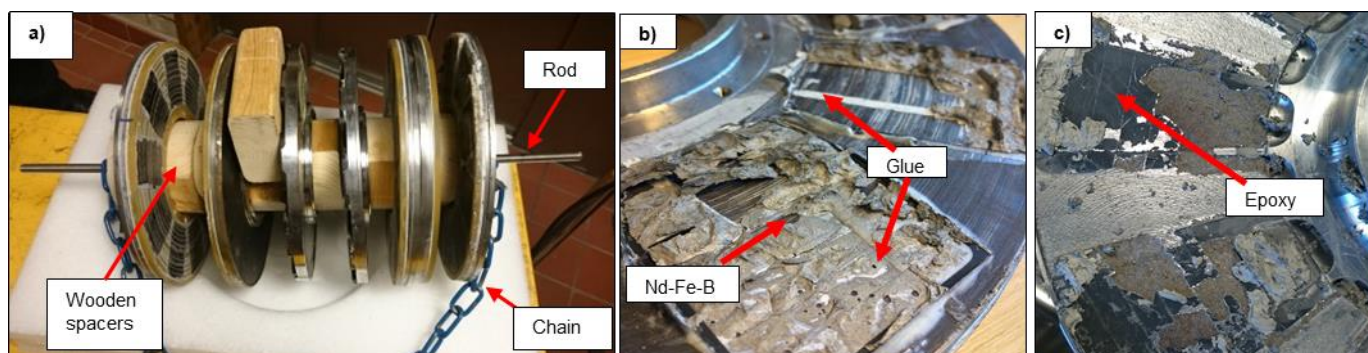
A JEOL 7000 scanning electron microscope (SEM) was used for microstructural imaging of the magnets, using a 20 kV accelerating voltage in backscattered mode, as well as compositional analysis using energy-dispersive x-ray spectroscopy (EDX). Optical microscopy, using a Laborlux 12HL microscope with Leitz lenses from 10x-100x, was used to visually study the sieving fractions and the epoxy coating before and after thermal demagnetisation treatments. Chemical analysis of the sieving fractions and starting magnets was performed by Less Common Metals Ltd., using a Perkin Elmer Optima 5300 DV inductively coupled plasma optical emission spectrometer (ICP-OES). Carbon analysis was performed by Magneti Ljubljana using an ELTRA CS-800, unless otherwise stated.

4. Results and Discussion

4.1. Extraction of NdFeB from Rotors Using Hydrogen

Seven semi-embedded rotors and non-embedded rotors were processed separately in two experiments. All the magnets in individual slots were scored in concentric circles using an angle grinder to provide a fresh surface for the hydrogen, as shown in Figure 3, 15-30 minutes before being loaded into the hydrogen vessel. The rotors were placed on a non-magnetic

201 metal rod, with wooden spacers between them so that the rotors would not stick together. The
202 rod was fastened to the top of the inside of the reactor with a chain, as shown in Figure 2b.



203

204 **Figure 3: A non-embedded rotor scored, a) placed on a rod with wooden spacers. b) After hydrogen decrepitation**
205 **at 1.2-1.5 bar for 135 hours, some material was still left in the corners of some slots.**

206 The hydrogen decrepitation was carried out at a pressure range of 1.2–1.5 bar, for a total
207 duration of 114 and 135 hours for the non-embedded and semi-embedded rotors, respectively.
208 The experiment was allowed to continue for such long time because the hydrogen pressure
209 was monitored throughout and the experiment was not ended until the pressure had been
210 stable for several hours. This was taken as indication that the reaction has completed. The
211 extraction was different between the semi- and non-embedded rotors. The semi-embedded
212 rotors were agitated using argon run pneumatic spinner after 18 hours of hydrogen exposure,
213 and it was observed from a small viewing window in the lid of the reactor that powder was
214 liberated from the assemblies. In the case of the non-embedded rotors, it was observed that
215 the powder self-liberated from the assemblies, so argon spinning was only carried towards the
216 end of the experiemnt to make sure all the powder was dropped into the collection pot.

217 **The yield was higher for the non-embedded rotors than for the semi-embedded rotors. The comparison**
218 **between the two rotors is given in**

219

220 Table 1. It was observed on the semi-embedded rotors that a small amount of the magnets
221 was still mechanically stuck in the bottom corners of the slots, as shown in Figure 3. The extent
222 of this would vary from rotor to rotor. However, for the non-embedded rotors, there was some
223 variation in the appearance of the rotors after exposure to hydrogen. Residues of glue and
224 epoxy, and sometimes magnet powder, was observed on some rotors, whereas there were
225 no residues in other cases. This is likely to be due to differences in how the magnets were

affixed to the rotors, and differences in surface materials on the rotor's ferritic stainless steel base plate.

Table 1: Comparison of the extraction of NdFeB from non-embedded and semi-embedded rotors. (* The quoted yield includes a high degree of error because of the presence of epoxy and glue in the liberated powder)

Rotor type	Time (hours)	Yield (%)	Observation
Non-embedded	114	98*	A portion of powder self-liberated. Only a minor amount of powder remained on rotors.
Semi-embedded	135	92*	Chunks of un-liberated NdFeB mechanically stuck on rotors.

It was also observed that the non-embedded rotors reacted quicker than the semi-embedded rotors, both with regards to the shorter reaction time, and a quicker drop in pressure in the early phase of the reaction. It is possible that the embedding slows down the reaction, as it hinders the volume expansion. The volume expansion is key in the decrepitation process, as it leads to cracking (Moosa and Nutting, 1988), and as fracturing occurs this create fresh surfaces for the reaction to propagate.

4.2. Decontamination of Extracted Powders

In order to recycle the extracted powder into new magnets, any auxiliary materials (glue, plastic) and impurities must be removed. Work by Burkhardt et al., has shown that some polymer coatings break up during processing and contaminate the final material, making it unsuitable for reprocessing (Burkhardt et al., 2019). Work by Lopez et al., showed that increases in carbon have an adverse impact on the final magnetic properties (Lopes et al., 2012). This purification process will take place in an inert atmosphere to avoid oxidation of the RE-rich phase. As already discussed in the literature review section 2, both oxygen and carbon

248 will have detrimental effects on the sintering process and magnetic properties of the recycled
249 magnets.

250 Chemical analysis of the extracted powder from the rotors showed an increased carbon
251 content in the extracted powder; 1750 ppm compared to 770 ppm in the base alloy, which is
252 explained by the observed adhesive residues. It was investigated whether sieving could
253 remove these impurities using the same process as for removing nickel from HDD scrap after
254 HPMS . A sieve stage with mesh sizes of 2000 μm , 500 μm , 250 μm , 150 μm , 90 μm and 45
255 μm was used. For the sieving trials, 250 g of powder was loaded the top mesh, and the
256 automatic shaker would agitate the sieves for 30 minutes until the experiment was ended. The
257 powder was collected from each mesh to be further analysed. As can be seen in Figure 4(a),
258 large chunks of glue and other binding materials were separated by the 2000 μm sieve.
259 However, the magnets' epoxy coating remained in all fractions, which was identified using
260 optical microscopy, see Figure 4 (b and c). The epoxy was present either on the surface of
261 the NdFeB powder particles or on its own in large flakes. The carbon content of each fraction
262 is plotted in Figure 4 (d), together with the weight fractions of hydrogenated powder remaining
263 in each sieve upon completion of the 30 minutes.

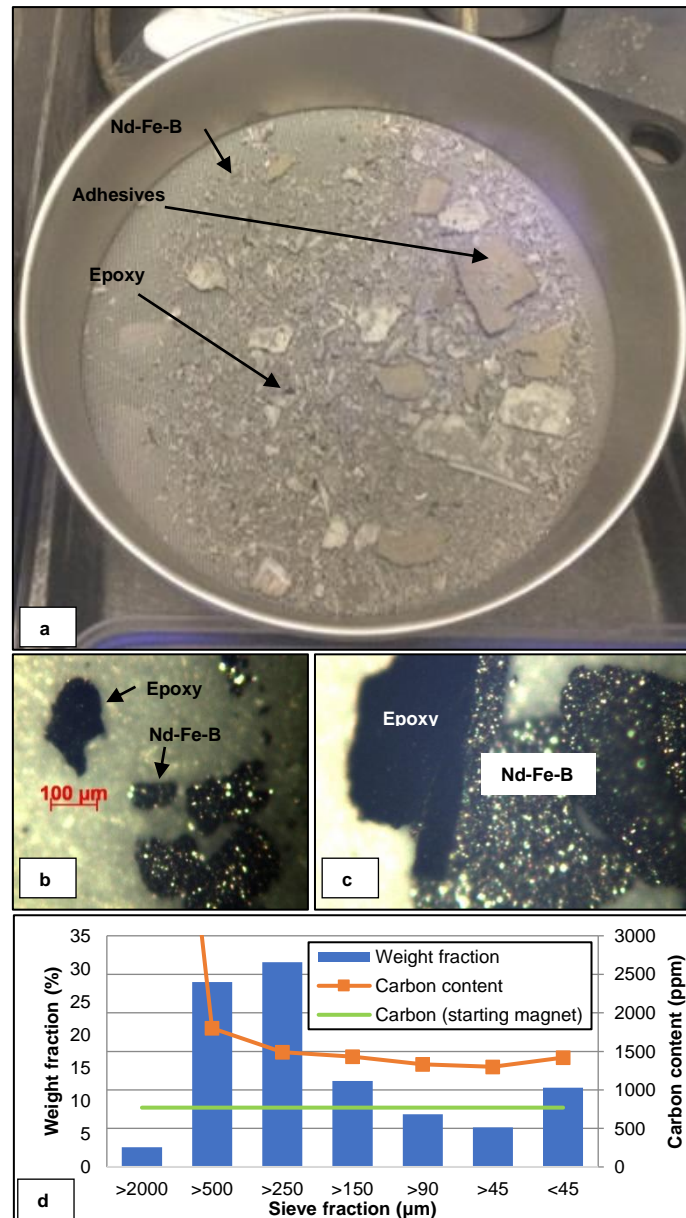


Figure 4: a) The residual impurities left on the 2000 μm sieve stage, however, all fractions contained epoxy from the coatings, either free (b) or on the NdFeB particles (c). This was reflected in the carbon analysis of each sieving fraction (d).

From the carbon analysis, it can be seen that even after sieving down to less than 45 μm the powder contains ~ 1500 ppm of carbon. From the work by Lopez et al., it can be seen that these levels of carbon would have an adverse impact on the magnetic properties of the magnets produced. The magnetic properties of the metal injection moulding (MIM) parts with carbon levels of 1500 ppm had inferior magnetic properties compared to those produced by conventional means with a carbon content of 600 ppm (Lopes et al., 2012).

4.3. Effect of Dy on the Kinetics of Hydrogen Decrepitation

Uncoated, sintered NdFeB magnets with three different Dy contents were exposed to hydrogen. The Dy content of the magnet (A), (B) and (C) was 1.00wt.%, 6.8wt.% and 8.45wt.%, respectively. In the first set of experiments, each magnet composition was exposed to three different lengths of air exposure times: 5, 30 and 60 minutes, before the system was closed and evacuated. These time scales were selected to investigate how long damaged magnets can be exposed to air for before hydrogen extraction takes place, or if the exposure to air would inhibit the hydrogen absorption, as it has been shown that the RE-rich phase starts to oxidise when exposed to air (Meakin et al., 2016).

All nine samples were cut into 3.65 mm diameter cylinders using an electrical discharge machine and ground to 152 mg under an argon atmosphere where they were also stored. Just before measurements were performed, the samples were quickly ground in the air on each flat side, to get the weight down to 150 mg and to keep the surface conditions constant.

The absorption of hydrogen was then carried out at 1.2 bar hydrogen pressure and at room temperature, which are typical processing conditions for the HPMS process, using an IGA. The absorption traces for the 9 experiments are shown in Figure 5 and summarised in Table 2.

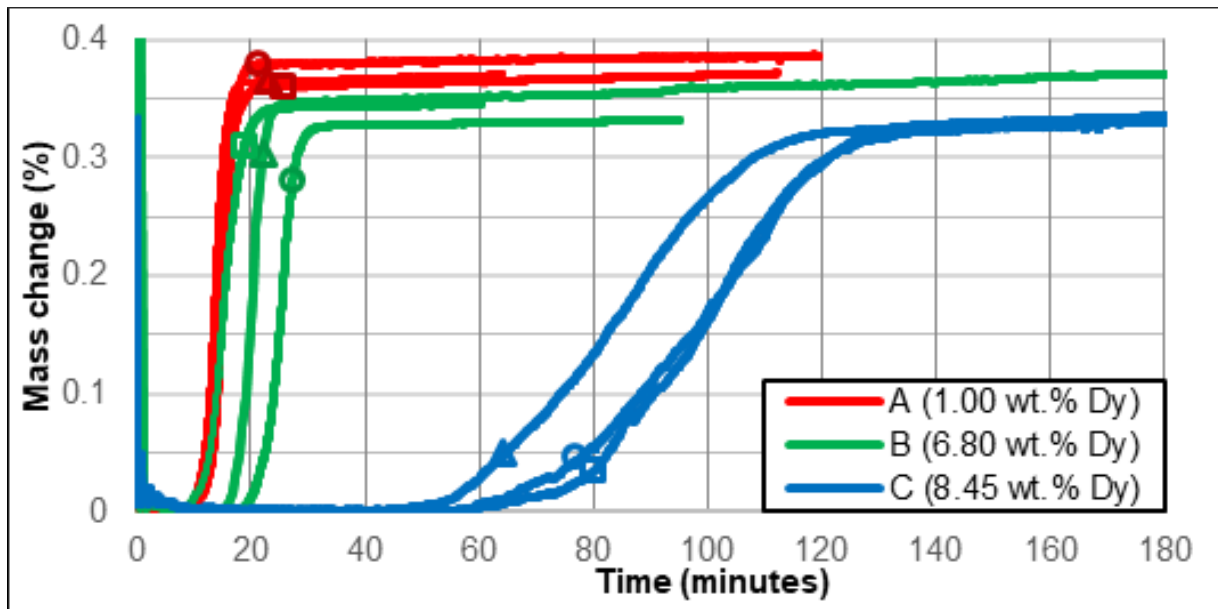


Figure 5: Gravimetric measurements showing the mass change vs time during hydrogen decrepitation at 1.2 bar for magnets (A), (B) and (C), which only differ in Dy content.

The initiation time refers to the time it takes from the start of the hydrogen pressure ramp (the pressure ramp started at $t=0$ minute and reached 1.2 bar at approximately $t=6$ minutes) to the start of the S-shaped curve. Saturation time refers to the time taken until the end of the S-

shaped curve, i.e. when the magnet is fully saturated with hydrogen and absorption is complete. The absorption time is the time from initiation to saturation.

Table 2: Summary of the gravimetric measurements at 1.2 bar hydrogen pressure.

Sample name (wt.% Dy)	Air exposure (minutes)	Initiation time (minutes)	Saturation time (minutes)	Absorption time (minutes)	Hydrogen absorbed (wt.%)
A (1.00)	5	6.8	20.4	13.7	0.378
A (1.00)	30	8.1	18.6	10.5	0.363
A (1.00)	60	7.8	21.4	13.6	0.359
B (6.80)	5	13.7	27.0	13.3	0.346
B (6.80)	30	16.9	34.2	17.3	0.327
B (6.80)	60	7.8	24.3	16.5	0.342
C (8.45)	5	57.0	135.1	78.1	0.321
C (8.45)	30	42.0	121.9	79.9	0.322
C (8.45)	60	53.6	133.5	80.0	0.326
1.00: Avg. (SD)		7.6 (0.7)	20.1 (1.4)	12.6 (1.8)	0.367 (0.010)
6.80: Avg. (SD)		12.8 (4.6)	28.5 (5.1)	15.7 (2.1)	0.338 (0.010)
8.45: Avg. (SD)		50.9 (7.9)	130.2 (7.2)	79.3 (1.1)	0.323 (0.002)

From Figure 5 and Table 2, the following can be observed:

1. Varying the air exposure time between 5 and 60 minutes prior to hydrogen exposure shows inconclusive results, this could be due to several factors including humidity. Humidity has a big influence on the corrosion behaviour of sintered NdFeB (Yan et al., 2009).

Saje et al., (1992) observed for the $\text{Nd}_{16-x}\text{Dy}_x\text{Fe}_{76}\text{B}_8$ ($x = 0-3$) cast alloy; that air exposure times between 5 and 1200 minutes led to a parabolic increase in initiation time for hydrogen decrepitation at 1 bar pressure and 20 °C.

2. Increasing Dy content leads to delayed initiation of the hydrogen decrepitation reaction; it took on average 7.6 minutes for the magnet (A) to start reacting, 12.8 minutes for the magnet (B), and 50.9 minutes for the magnet (C).

3. Increasing Dy content leads to increased absorption time. On average, it took 12.6 minutes for the magnet (A), 15.7 minutes for the magnet (B) and 79.3 minutes for the magnet (C).

Hirano et al., (2006) have previously shown that Dy metal is more stable against hydrogenation than Nd. In their work, 1 g of pure Nd and Dy metal was individually exposed to 20 bar hydrogen at 273 K, and it was found that Dy had both delayed initiation and absorption times. The saturation time was 7 minutes and 830 minutes for Nd and Dy, respectively. This could explain point 2 and 3 above, as the hydrogen decrepitation reaction both initiates and propagates through the RE-rich grain boundaries. If these both initiate and react more slowly, this would lead to longer initiation and reaction times for the hydrogen decrepitation process as a whole. Furthermore, as the hydrogenation of the matrix phase is accommodated by the exothermic heat from the RE-hydride formation at the grain boundary (Harris et al., 1987), a slower reaction at the grain boundary could mean that the local temperature increase is lower, slowing down the diffusion of hydrogen into the matrix phase. However, Saje et al., (1992) did not see any effect of the Dy content on the hydrogen decrepitation kinetics of the $\text{Nd}_{16-x}\text{Dy}_x\text{Fe}_{76}\text{B}_8$ ($x = 0-3$) cast alloy at 1 bar pressure at 20 °C. This could be explained by the higher RE content of approximately 35.1 wt.% in their work, compared to 31.1 wt.% in this work. With a higher amount of RE-rich phase, the effect of the Dy additions would be smaller as there would be a high amount of Nd present in this phase.

4. Increasing Dy content leads to less hydrogen being absorbed by the material: on average, magnet (A) absorbed 0.367 wt.% hydrogen, magnet (B) 0.338 wt.%, whereas magnet (C) absorbed 0.323 wt.% hydrogen.

This observation is supported by the work by Bartolomé et al., (1991), who showed that the $\text{Dy}_2\text{Fe}_{14}\text{B}$ phase absorbed less hydrogen than its Nd counterpart (Bartolomé et al., 1991). Furthermore, the volume expansion is lower for the Dy compound (3.5 vol.%) than the Nd compound (3.9 vol.%). The decrepitation reaction is based on the differential volume expansion of the matrix and grain boundary phases leading to inter- and trans-granular cracking, leading to pulverisation of the magnet (Moosa and Nutting, 1988). A lower volume expansion may affect the HPMS process with higher Dy contents will not break down to powder as efficiently as the low Dy content magnets, and hence why they

are more difficult to liberate from the assemblies after completion of the hydrogenation reaction.

5. Magnet (C) absorbed hydrogen significantly slower than magnets (A) and magnet (B), although the compositional differences between magnets (B) and magnet (C) are not very large. The Dy content is larger in the RE-rich phase for the magnet (C) which would slow down the reaction. However, the differences in the composition are much larger between magnets (A) and magnet (B), and hence this result is confusing.

Harris et al., (1987) have shown that increased hydrogen pressure led to a decrease in both initiation and saturation time for the $\text{Nd}_{15}\text{Fe}_{77}\text{B}_8$ cast alloy, following a logarithmic relation in both cases. Higher hydrogen pressure could thus potentially be used to speed up the reaction of the magnet (C), and the extraction of NdFeB from the rotors. In a subsequent set of experiments, higher hydrogen pressures were applied to the magnet (C): 2.5 bar, 5 bar and 10 bar. Three runs were performed for every pressure, with 5 minutes of air exposure before each run. The results are shown in Figure 6, where the absorption time is plotted versus hydrogen pressure, and summarised in Table 3.

Table 3: Summary of the gravimetric measurements of the magnet (C) at varying hydrogen pressure.

Pressure (bar)	Air exposure (minutes)	Initiation time (minutes)	Saturation time (minutes)	Absorption time (minutes)	Amount absorbed (wt.%)
1.2	5	57.0	135.1	78.1	0.321
1.2	30	42.0	121.9	79.9	0.322
1.2	60	53.6	133.5	80.0	0.326
2.5	5	81.0	124.1	43.1	0.320
2.5	5	80.2	143.4	63.2	0.326
2.5	5	64.5	121.6	57.1	0.335
5.0	5	26.7	69.0	42.3	0.328
5.0	5	40.2	85.6	45.4	0.294
5.0	5	70.0	108.8	38.8	0.335
10.0	5	50.9	69.3	18.4	0.340
10.0	5	20.0	57.9	37.9	0.326
10.0	5	46.8	75.2	28.4	0.337
1.2: Avg. (SD)		50.9 (7.9)	130.2 (7.2)	79.2 (1.1)	0.323 (0.003)

2.5: Avg. (SD)		75.2 (9.3)	129.7 (11.9)	50.5 (10.3)	0.327 (0.008)
5.0: Avg. (SD)		45.6 (22.2)	87.8 (20.0)	42.2 (14.1)	0.319 (0.003)
10.0: Avg. (SD)		39.3 (16.8)	67.5 (8.8)	28.2 (9.8)	0.334 (0.008)

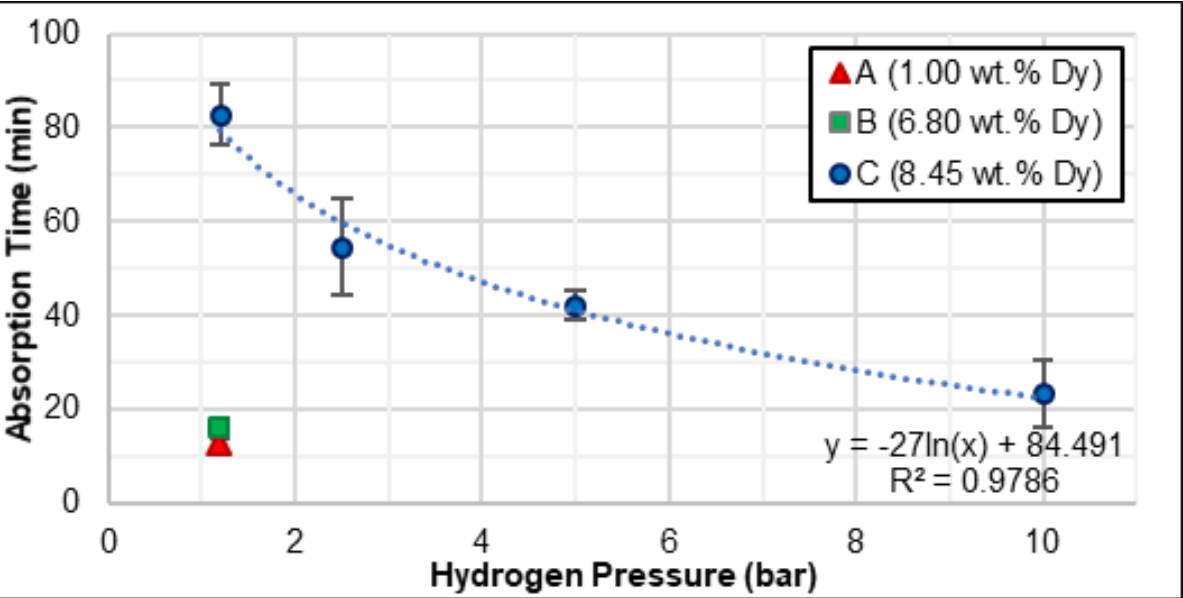


Figure 6: Absorption time vs hydrogen pressure, generated from gravimetric measurements.

There is a clear decrease in absorption time with increasing pressure, which appears to follow a logarithmic relation very similar to what Harris et al., have shown for a $\text{Nd}_{16}\text{Fe}_{76}\text{B}_8$ cast alloy (Harris et al., 1987). However, there was no clear reduction in initiation time and the initiation data is inconclusive. Increasing the pressure from 1.2 bar to 5 bar and 10 bar, the absorption time decreased from 79 minutes to 42 and 28 minutes, respectively. This result means that it is likely that the extraction from the rotors could be significantly quicker by increasing the hydrogen pressure.

5. Conclusions

This paper is an early attempt at processing one particular design of automotive rotor and it is important to note that there are many different designs where the magnets are often more heavily embedded. However the data gathered and discussed in this paper can provide some initial information to help with design for recycling. For example if the systematic eco labelling outlined in the MaXycle project could be applied, then the composition of magnet would be known which would guide the process conditions for HPMS. Alternatively by choosing an appropriate binding material which does not break apart to the same extent on hydrogen

processing it would make the purification stage much easier. This is a focus of the on-going SUSMAGPRO project in the EU.

Furthermore, it is clear that a pre-treatment stage is required for the HPMS process, as is the case with hard disk scrap, to initiate the reaction. Once the HPMS powder is generated then in this case the epoxy-coating could not be entirely separated from the hydrogenated powder with the technique used in this work. The carbon content was reduced from 1 % (10,000 ppm) to 1490 ppm with a 250 µm sieve but on further fine sieving the carbon content was still at 1420 ppm, which is significantly higher than 770 ppm in the base alloy. It is not clear whether different polymer coatings would interact in the same way and a much larger sample size of drive motors would be required to investigate this. There may be a detrimental effect on the magnetic properties of the recycled magnets if this powder was used to sinter magnets without further purification, although it would require further work to confirm this. Further purification methods need to be examined to reduce the carbon content down to acceptable levels as well as choosing appropriate binding materials to avoid this.

The hydrogen decrepitation reaction was shown to be significantly slower for the rotor magnets compared to lower Dy content magnets, which could be attributed both to the lower rare earth content and higher Dy content of these magnets. Gravimetric measurements confirmed that increasing Dy contents slows down absorption times, which has previously been unclear. Therefore a higher pressure would be required to process automotive magnets. It is also important to note that through time automotive magnets have been gradually decreasing in Dy content.

Acknowledgements

The research leading to these results has received funding from the European Community's Seventh Framework Programme ([FP7/2007–2013]) under grant agreement no. 607411 (MC-ITN EREAN: European Rare Earth Magnet Recycling Network). Project website: <http://www.erean.eu>.

Muhammad Awais received funding from the European Community's Horizon 2020 Programme (H2020/2014–2019) under Grant Agreement no. 674973 (MSCA-ETN DEMETER). This publication reflects only the authors' view, exempting the Community from any liability. Project website: <http://etn-demeter.eu/>.

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