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## Comparative techno-economic and life-cycle analysis of precious *versus* non-precious metal electrocatalysts: the case of PEM fuel cell cathodes†

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Sluggish kinetics in the oxygen reduction reaction (ORR) require significant quantities of expensive Pt-based nanoparticles on carbon (Pt/C) at the cathode of proton exchange membrane fuel cells (PEMFCs). This catalyst requirement hinders their large-scale implementation. Single atom Fe in N-doped C (Fe–N–C) electrocatalysts offer the best non-Pt-based ORR activities to date, but their environmental impacts have not been studied and their production costs are rarely quantified. Herein, we report a comparative life-cycle assessment and techno-economic analysis of replacing Pt/C with Fe–N–C at the cathode of an 80 kW PEMFC stack. In the baseline scenario (20 g<sub>Pt/C</sub> vs. 690 g<sub>Fe–N–C</sub>), we estimate that Fe–N–C could reduce damages on ecosystems and human health by 88–90% and 30–44%, respectively, while still increasing global warming potential by 53–92% and causing a comparable impact on resource depletion. The environmental impacts of Pt/C predominantly arise from the Pt precursor while those of Fe–N–C are presently dominated by the electricity consumption. The monetized costs of environmental externalities for both Fe–N–C and Pt/C catalysts exceed their respective direct production costs. Based on catalyst performance with learning curve analysis at 500 000 PEMFC stacks per annum, we estimate replacing Pt/C with Fe–N–C would increase PEMFC stack cost from 13.8 to 41.6 USD per kW. The cost increases despite a reduction in cathode catalyst production cost from 3.41 to 0.79 USD per kW (excluding environmental externalities). To be cost-competitive with a Pt-based PEMFC stack delivering 2020 US Department of Energy target of 1160 mW cm<sup>−2</sup> (at 0.657 V), the same stack with an Fe–N–C cathode would need to reach 874 mW cm<sup>−2</sup>, equivalent to a 200% performance improvement. These findings demonstrate the need for continued Fe–N–C activity development with sustainable synthesis routes in mind to replace Pt-based cathode catalyst in PEMFCs. Based on forecasting scenarios of fuel cell vehicle deployment targets, we find that Pt consumption would be constrained by Pt supply.

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## Introduction

The International Energy Agency predicts that for a global net zero emission scenario by 2050, 60 Mt of hydrogen will be used for power generation.<sup>1</sup> Hydrogen-fed fuel cells, specifically low-temperature polymer electrolyte membrane fuel cells (PEMFCs), are a promising zero CO<sub>2</sub> emission technology with high-technology readiness to fulfill sectors of power generation.<sup>2</sup> This is due to PEMFCs' load flexibility and therefore wide range of applications within stationary and portable power generation, and particularly transportation.<sup>2</sup> PEMFCs' rising prominence in transportation applications is supported by global Government targets of 2.5 million fuel cell electric vehicles (FCEV) on the road by 2030.<sup>3</sup>

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Pt-based nanoparticles on carbon (Pt/C) catalysts are commonly implemented in commercial PEMFCs to facilitate the electrochemical reactions.<sup>4</sup> The reaction kinetics at the cathode, where the oxygen reduction reaction (ORR) occurs, are far more sluggish than at the anode and, as a result, the cathode has previously required 80–90% of the total Pt within the PEMFC.<sup>5</sup> Hence, Pt-based cathode catalysts have been projected as the largest single component cost of an 80 kW PEMFC stack at production rates of 500 000 stacks per year.<sup>6</sup> The price volatility of Pt and subsequent catalyst cost sensitivity would also heavily impact fuel cell costs, making wider-scale commercialization difficult.<sup>7</sup> In terms of Pt supply, South Africa provided 65–74% and Russia 10–14% of the world's Pt between 2017 and 2021.<sup>8</sup> Moreover, industrial recycling of Pt has remained low to date (total <30%, combustion automotive <55%).<sup>8,9</sup> This dependency on fresh Pt supplies is unfavorable since highly reliant on global supply chains and trade agreements. As such, in the UK and EU, Pt is considered a critical raw material due to its high global supply risk and economic vulnerability.<sup>9,10</sup>

Research efforts have been directed at replacing Pt from the PEMFC cathode, also with a view to bringing PEMFC system costs down to the 30 USD per kW target set by the US Department of Energy (DOE).<sup>6,11</sup> The most promising alternative catalysts are those based on metal single atoms coordinated to nitrogen within a conductive carbon scaffold, classified as M–N–C, where M refers to a transition metal (Fe, Mn and/or Co).<sup>12</sup> Of these, Fe–N–C catalysts show the highest ORR activity,<sup>13</sup> and have the potential for low production cost, while relying only on highly abundant and widely accessible elements. The activity and stability of Fe–N–C catalysts in PEMFCs have recently improved substantially.<sup>14</sup> However, state-of-the-art Fe–N–C synthesis require several steps usually with two energy intensive pyrolysis steps. Simple, scalable approaches to single atom Fe–N–C synthesis are also available,<sup>15,16</sup> but this is typically detrimental to the ORR activity. Companies such as Pajarito Powder, LLC (USA)<sup>17,18</sup> Nisshinbo Holdings Inc. (Japan)<sup>19,20</sup> and lately Celcibus AB (Sweden),<sup>21</sup> have invested in commercial development of Fe–N–C catalysts. Pajarito Powder recently demonstrated initial H<sub>2</sub>–air PEMFC performance with their Fe–N–C cathode catalyst comparable to commercial Pt/C, despite degradation still falling short of US DOE targets (60–119 mV vs. <30 mV loss US DOE target at 0.8 A cm<sup>-2</sup> after 30 000 cycles between open circuit potential and 0.6 V).<sup>17</sup>

With the technology readiness level and performance of Fe–N–C catalysts continually improving, it is important to assess their environmental impact and commercial viability compared to Pt-based catalysts through life-cycle assessment (LCA) and techno-economic analysis (TEA). Both LCA and TEA have been conducted on PEMFCs, including some analysis of the critical materials,<sup>22</sup> yet with a limited focus on the Pt-based electrocatalyst.<sup>23</sup> Stropnik *et al.* estimated that a mere 0.75 g of Pt contributes around 60% of the total environmental impact of manufacturing a 1 kW PEMFC system.<sup>24</sup> In terms of global warming potential (GWP), this is not surprising since Pt production emits ~10 000 kg CO<sub>2</sub>-eq per kg in comparison to only

~1 kg CO<sub>2</sub>-eq per kg for Fe.<sup>25</sup> Stropnik and co-workers subsequently found for a 48 kW PEMFC stack and in an optimistic scenario that climate change impact could be reduced by 54% if 95% of the Pt were to be recycled.<sup>26</sup> Recycled Pt has been demonstrated in PEMFC,<sup>27</sup> and EU reports estimate that 76% of Pt in end-of-life PEMFCs is already recoverable using laboratory-scale processes.<sup>28,29</sup>

Another recent EU report comparing a carbon nanotube-based Fe–N–C and Pt/C cathode catalyst predicted that the GWP, human toxicity, abiotic depletion, and eutrophication impacts of Fe–N–C could be greater than those of Pt/C.<sup>30</sup> The same report found that the fluorine-containing Nafion<sup>TM</sup> membrane and ionomer could be the dominating component for environmental impacts for comparable performance systems.<sup>30</sup> Nevertheless, alternative membranes have recently been developed such as Pemion<sup>TM</sup> that eliminate environmental concerns of per- and polyfluoroalkyl substances, which could soon be banned in the EU.

In 2005, Gasteiger *et al.* calculated that for a costless Pt-free cathode catalyst to be economically viable in a PEMFC, it would need to have at least 10% of the activity of Pt under equivalent conditions.<sup>31</sup> While this has been achieved by Fe–N–C catalysts in the meantime, one also needs to account for the material and manufacturing costs of these catalysts. In 2015, a TEA comparison at 500 000 80 kW PEMFC systems per year between a polyaniline-based Fe–N–C catalyst with low material cost of 74–129 USD per kg and a specialized Pt-based alloy catalyst at ~41 000 USD per kg concluded that the Fe–N–C would require a power density improvement from 330 mW cm<sup>-2</sup> to 475 mW cm<sup>-2</sup> at 0.5 V to compete with the Pt-based catalyst, but this was based purely on material costs.<sup>32</sup> Moreover, to the best of our knowledge, the environmental life cycle impacts of Fe–N–C catalyst production have not been quantified yet, let alone comparisons between the environmental impacts of Pt/C and Fe–N–C electrocatalysts in a PEMFC cathode.

Herein, we conduct a comparative techno-economic and environmental analysis of replacing Pt/C electrocatalyst with a Fe–N–C electrocatalyst for the ORR in an 80 kW PEMFC stack, considering only the initial ORR activity and neglecting degradation. We conduct a cradle-to-gate LCA, including the propagation of key process uncertainties on predicted environmental impacts in a sensitivity analysis, and we investigate a range of possible future scenarios. We also perform a detailed cost analysis of both catalysts under a large-scale deployment scenario of 500 000 PEMFC stacks per annum using a learning curve method.

## Methodology

### Life-cycle analysis

LCA is a prominent methodology for quantifying the environmental burdens associated with products, processes, or activities across their life-cycle—from extraction and processing of raw materials to recycling and final disposal.<sup>33</sup> Through aggregating the environmental burdens under a set of impact cat-



egories and indicators, LCA results can be easily understood and used to guide sustainable product and process design. LCA has been applied extensively to compare the production routes of chemicals and fuels, including early-stage technologies,<sup>34</sup> but its application to catalyst production has remained scant.<sup>35</sup>

Herein, we apply LCA to quantify and compare the environmental impacts of manufacturing Pt/C and Fe–N–C electrocatalysts for use in a PEMFC cathode. Of all the PEMFC stack components, such focus on the cathode electrocatalyst (Fig. 1a) is motivated by its large repercussion on PEMFC efficiency, cost, lifetime, and environmental impact, with substantial ongoing research on displacing Pt-based cathode catalyst. The two electrocatalysts are compared on a dry basis, so both ionomer and inks for depositing catalysts are ignored in the LCA. The assessment is conducted using OpenLCA 1.11 and the database ecoinvent 3.6.<sup>36</sup> The LCA comprises four phases per the ISO standards (14040:2006 and 14044:2006/AMD 2:2020): (1) goal and scope definition, (2) inventory analysis, (3) impact assessment, (4) interpretation.<sup>37</sup>

### Goal and scope

The goal of the LCA is to compare the environmental impacts for the manufacturing phase of Pt/C and Fe–N–C cathode electrocatalysts for an 80 kW PEMFC stack for light duty vehicle applications. The aim is to help stakeholders and policy-

makers take decisions for future investments in PEMFC cathode catalyst development, as well as informing the research community and public. Accordingly, the geographical location is chosen as the global level in the baseline scenario.

We adopt a cradle-to-gate scope for the LCA (Fig. 1b), which includes all processes from raw material extraction to the electrocatalyst production and where allocation is not required since Pt/C and Fe–N–C are the sole products. The use phase of the electrocatalysts is accounted for through setting the functional unit as the mass of cathode catalyst (grams) needed for a PEMFC stack to provide power of 80 kW at 0.657 V, including the effect of electrocatalyst amount in a *post hoc* scenario analysis. Despite Fe–N–C currently being less stable than Pt/C, Fe–N–C stability is rapidly improving<sup>14</sup> so we make the assumption herein that Fe–N–C will eventually reach equivalent operation lifetimes to Pt/C. Regarding the end-of-life, while our baseline scenario assumes no recycling for the spent electrocatalysts, the effect of Pt recycling is investigated as part of the *post hoc* scenario analysis instead. Incorporating more details on the treatment of the spent electrocatalysts, degradation of the catalyst or PEMFC lifetime into the LCA is beyond the scope of this paper and left for future research.

The main performance characteristics of the analyzed baseline 80 kW PEMFCs are summarized in Table 1. The Pt/C cathode electrocatalyst is based on the 2020 US DOE target and performance of  $0.125 \text{ mg}_{\text{Pt}} \text{ cm}^{-2}$  for light duty fuel cell vehicles using PEMFC technology,<sup>7,38</sup> resulting in around 20 g of Pt/C catalyst per functional unit (80 kW); refer to section A of the ESI for details.<sup>†</sup> For the Fe–N–C electrocatalyst, PEMFC performance is assumed equivalent to a Pajarito Powder Fe–N–C catalyst with  $290 \text{ mW cm}^{-2}$ ,<sup>17</sup> leading to an estimated 690 g of Fe–N–C catalyst per functional unit (see section A of ESI<sup>†</sup>). The baseline scenario, therefore, compares 20 g of Pt/C against 690 g of Fe–N–C.

### Life cycle inventory

**Pt/C catalyst synthesis.** The Pt/C synthesis process (Fig. 2a) is based on reported polyol syntheses.<sup>39,40</sup> It uses chloroplatinic acid since it is the most common Pt salt precursor in the industry.<sup>41</sup> The scale-up procedure involves mixing 40 mM chloroplatinic acid with a solution containing 0.4 M NaOH in ethylene glycol.<sup>39</sup> Next, 1 M HCl is added to the Pt colloidal solution to precipitate the nanoparticle suspension, prior to

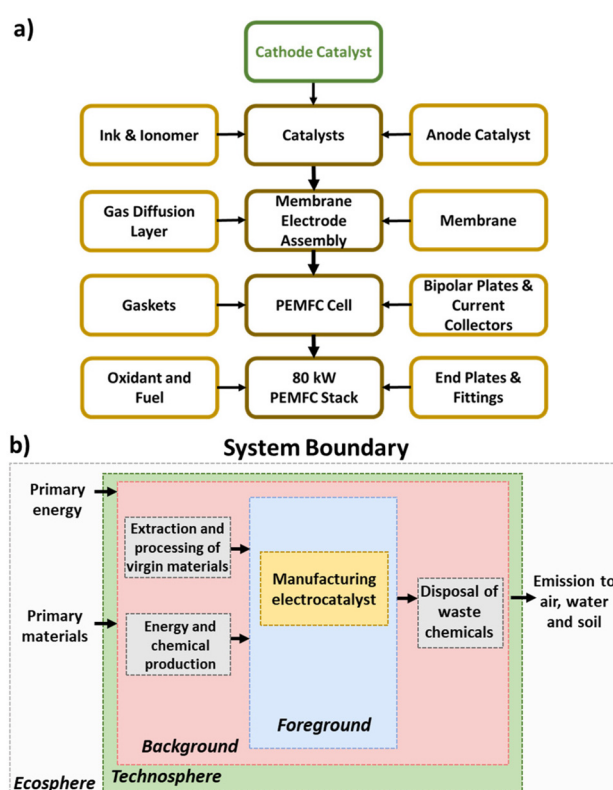


Fig. 1 (a) PEMFC stack component hierarchy, with highlighted focus on the cathode catalyst. (b) System boundary of the PEMFC cathode electrocatalyst LCA.

Table 1 Performance, operating conditions, and components of the PEMFC stack in the baseline scenario for Pt/C and Fe–N–C. Operating temperatures and pressure are assumed to be identical in both systems (80 °C, 1 bar<sub>g</sub>, H<sub>2</sub>–air)

Parameter	Pt/C	Fe–N–C
Power density at 0.657 V (mW cm <sup>-2</sup> )	1160	290
Cathode catalyst loading (mg <sub>Pt</sub> cm <sup>-2</sup> or mg <sub>Fe–N–C</sub> cm <sup>-2</sup> )	0.125	2.5
Active cells (–)	380	11 035
Cell active area (cm <sup>2</sup> )	185	25
Total cathode catalyst required (g)	20	690



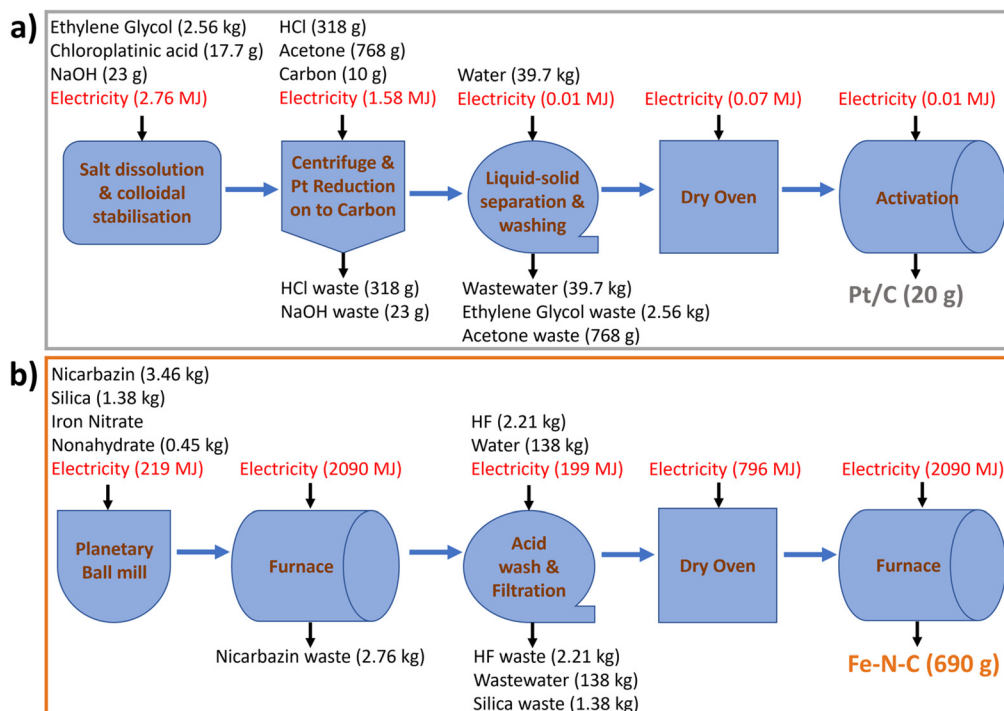


Fig. 2 Process diagrams with material and energy inventories to produce baseline (a) 20 g Pt/C, (b) 690 g Fe–N–C.

centrifugating the mixture three times with supernatant discarded. The recovered Pt nanoparticles are mixed with carbon black in acetone before mixing and drying. The resulting Pt/C powder is then dried in an air oven before undergoing heat activation in a furnace<sup>35,42</sup> to produce the final Pt/C catalyst.

Initial lab-scale (55 mg Pt/C production) values of sodium hydroxide, ethylene glycol, chloroplatinic acid, hydrochloric acid, deionized water, and carbon black are for 50 wt% Pt/C<sup>39</sup> and adjusted to 46 wt% Pt/C, while acetone is from scaling an 8 g Pt/C lab process.<sup>40</sup> Energy inputs are from a previously

reported simulated 1 kg Pt/C manufacturing process,<sup>35,42</sup> scaled down to 20 g, and assuming 90% efficiency of electric heat ovens and excluding the carbon support precursor processing. The reactants are scaled in stoichiometric ratios with 20% relative reduction in solvents applied during scaling.<sup>43</sup> Material losses in Pt/C production, reported to be ~3% in manufacturing quotes, are ignored. All unreacted inputs are considered waste chemicals in the outlet stream. The foreground inventories for baseline 20 g Pt/C electrocatalyst are summarized in Table 2 (left column) and reported on the

Table 2 Manufacturing inputs and outputs for Pt/C and Fe–N–C catalyst baseline scenarios

Pt/C (20 g)		Fe–N–C (690 g)	
Input	Amount	Input	Amount
Acetone (g)	768	Activated silica (kg)	1.38
Carbon black (g)	10.0	Hydrogen fluoride (kg)	2.21
Chloroplatinic acid (g)	17.7	Iron nitrate nonahydrate (kg)	0.345
Ethylene glycol (kg)	2.56	Nicarbazin (kg)	3.46
Sodium hydroxide (g)	23.3	Water, deionised (kg)	138
Water, deionised (kg)	39.7	Electricity (MJ)	5393
Hydrochloric acid (g)	318		
Electricity (MJ)	4.43		

Output	Amount	Output	Amount
Acetone, waste (g)	768	Silica, waste (kg)	1.38
Hydrochloric acid, waste (g)	318	Fe–N–C Catalyst (g)	690
Pt/C (g)	20	Hydrogen fluoride, waste (kg)	2.21
Sodium Hydroxide, waste (g)	23.3	Nicarbazin, waste (kg)	2.76
Wastewater (kg)	39.7	Wastewater (kg)	138
Ethylene glycol, waste (kg)	2.56		

process diagram in Fig. 2a (further details in Table S2†). The lack of data for chloroplatinic acid in Ecoinvent 3.6 is circumvented through creating a separate process based on literature data (eqn (S1) and Table S1†).<sup>7</sup>

**Fe–N–C catalyst synthesis.** The preparation method and reactants for the Fe–N–C electrocatalyst (Fig. 2b) is based on patents filed by the University of New Mexico.<sup>44–46</sup> The N–C precursor (nicarbazin) is mixed with the Fe precursor (iron nitrate nonahydrate) and the hard template silica (Cab-O-Sil amorphous fumed) in a planetary ball mill for 1 h. Next, the mixture is pyrolysed ( $N_2$ , 100 cc min<sup>−1</sup>) at 10 °C min<sup>−1</sup> up to 900 °C and held for 1.5 h, before being cooled naturally. The resulting material is washed with 25% HF for 12 h to remove the silica template, followed by subsequent washing with copious amounts of deionized water until reaching neutral pH, using a filtration pump for 4 hours. The wet material is then dried in an air oven at 85 °C for 8 h, before repeating the same pyrolysis process to form the final Fe–N–C catalyst.

All energy inputs are scaled linearly from 25 g Fe–N–C production, obtained from laboratory data (Table S5†), and by assuming that the whole production process, including large-scale furnaces, could be electrified at scale. The reactants are scaled in stoichiometric ratios and water as solvent is scaled with a 20% relative reduction.<sup>43</sup> Material losses during production are neglected, aside from during pyrolysis. Based on expert opinion and previous laboratory data from the University of New Mexico, 80% of nicarbazin precursor is assumed to be wasted in the Fe–N–C process due to the initial pyrolysis.  $N_2$  gas flow during pyrolysis is not considered due to the low environmental impact found in OpenLCA. The amount of required HF is calculated based on the amount of silica (eqn (S2)†), with an extra 20% added to ensure complete consumption of silica. Since  $SiF_4$  is unavailable in ecoinvent 3.6, its environmental impact is assimilated to that of HF, under the assumption of equal input and output flows of HF. All output streams other than the catalyst are treated as a waste in the baseline scenario (see Scenario subsection below). The foreground inventories for baseline 690 g Fe–N–C are summarized in Table 2 (right column) and reported on Fig. 2b (further details in Table S6†). Data for iron nitrate nonahydrate and nicarbazin are also unavailable in ecoinvent 3.6. For the former, a separate process is created, with inputs determined using weight distribution data (Table S3†). For the latter, an equimolar mix of 2-nitroaniline and 2-pyridinol is assumed to be equivalent to the nicarbazin precursor (Table S4†). Any energy inputs for producing these two precursors are furthermore omitted.

## Impact assessment

Both the foreground and background inventories are translated into environmental impacts during the life-cycle impact assessment (LCIA) phase using a characterization method.<sup>47,48</sup> Our assessment follows the hierarchist perspective, which is based on the cultural theory of scientific agreement and common policy principles, adopting a medium timeframe of 100 years for the environmental impacts. The LCIA method-

ology of choice is ReCiPe 2016,<sup>49</sup> and of the 17 midpoint indicators in ReCiPe 2016, a particular emphasis is on: *global warming potential*, caused by greenhouse gas emissions, with a reference time of 100 years per the Kyoto Protocol and Paris agreement; *fossil fuel scarcity* and *mineral resource scarcity*, to capture the depletion of both fossil and non-fossil resources; and *marine eutrophication* and *freshwater eutrophication*, to describe the enrichment of the aquatic ecosystems with nitrogenous and phosphorous compounds, respectively. Highly relevant to the present assessment are also the midpoint indicators of *acidification potential* and *human toxicity potential*, which capture the impacts of acidifying and toxic substances both on human health and on the environment. Since ReCiPe 2016 fails to account for HF in these two indicators,<sup>50</sup> we used CML 2001 instead in those cases.

The midpoint indicators are further aggregated into endpoint categories for three areas of protection: *human health*, expressed in terms of disability-adjusted life-years (DALY), measures the number of years that a person is disabled after a disease or accident; *ecosystem quality* measures the local species loss integrated over time (expressed in species × year); and *resource scarcity* monetizes the burdens attributed to future mineral and fossil resource extraction (expressed in US\$ 2013). The last two endpoint indicators are quantified again using the ReCiPe 2016 methodology. To account for the impact of HF, the human health indicator is calculated by converting point units in ReCiPe 2008 to DALY in ReCiPe 2016 (world normalization factor of 73.3 and average weighting factor of 400).<sup>51</sup> The selected characterization methods for both midpoint and endpoint impacts are summarized in Table 3.

Alongside quantifying the environmental impacts, our assessment considers their monetary valuation as externalities, or monetisation in short. This procedure translates the endpoint environmental burdens into monetary units, thereby enabling a direct comparison both between categories and with manufacturing costs. Weidema and coworkers<sup>52,53</sup> first developed monetisation factors for the human health damage categories, based on the annual income generated through extending a person's life by one year, and later extended the approach by extrapolating from human health to ecosystem

**Table 3** Midpoint and endpoint impact categories used

Impact	Category	Characteristic model	Units
Mid-point	Global warming potential	ReCiPe 2016	kg CO <sub>2</sub> -eq
	Acidification potential	CML 2001	kg SO <sub>2</sub> -eq
	Fossil resource scarcity	ReCiPe 2016	kg oil-eq
	Marine eutrophication	ReCiPe 2016	kg N-eq
	Freshwater eutrophication	ReCiPe 2016	kg P-eq
	Human toxicity potential	CML 2001	kg 1,4-DCB-eq
	Mineral resource scarcity	ReCiPe 2016	kg Cu-eq
End-point	Human health	ReCiPe 2008	DALY
	Ecosystem quality	ReCiPe 2016	species × year
	Resource	ReCiPe 2016	USD 2013



damage. The monetisation factors (Table S7†) used herein are those by Dong *et al.*,<sup>54</sup> which are based on the hierarchist perspective.

## Interpretation

The final phase of LCA entails interpreting the results and checking that the conclusions are well-substantiated, including uncertainty quantification and scenario analysis. Of the multiple sources of uncertainty that can affect the calculations, we focused on uncertainty in foreground inventory data since we derived the Fe–N–C process inventories from scaled-up lab procedures.<sup>55</sup> We also defined a range of catalyst scenarios to account for future technological advances, with a view to reducing catalyst loading and raw material use and improving production processes.

**Uncertainty quantification.** We conducted an uncertainty analysis through jointly propagating the uncertainty in the foreground process inputs to all the midpoint and endpoint impact indicators, using Monte Carlo analysis with 1000 Sobol samples. The uncertainty ranges of  $\pm 20\%$  around the nominal process input values for both catalyst syntheses (Tables S8 and S9†) were informed by expert opinions. Due to the lack of further data, these uncertain parameters are assumed to follow triangular probability distributions.<sup>56</sup> In a second step, we used the same sampled scenarios in a global sensitivity analysis (GSA) to quantify how much each uncertain input contributes to the overall impact variance. We used the software SobolGSA for this, which constructs random sampling-high dimensional model representations (RS-HDMR) surrogates to compute Sobol sensitivity indices.<sup>57,58</sup> Note that each uncertain input contribution adds up separately to the impact variances by construction, so the first- and total-order Sobol indices are identical and sum up to 1. Refer to section B of the ESI† for further details.

**Scenario analysis.** In addition to the baseline assumption of 100% virgin Pt supply, we defined two additional scenarios with 55% and 75% Pt recycling to reflect current industrial automotive Pt recycling,<sup>8,9</sup> reported lab-scale results,<sup>28,59,60</sup> and EU guidelines.<sup>29</sup> For simplicity, these two scenarios omit the infrastructure and energy requirements for the Pt recycling process. Therefore, the recycled Pt could also be interpreted as Pt reductions, which is in agreement with the EU's target of 71% Pt reduction by 2030 (from 2017 values).<sup>61</sup> We considered another scenario with a reduction in Pt/C from 20 g to 15 g as Pt amounts in PEMFCs are continually reduced.<sup>4</sup> On the other hand, we did not analyze any recycling scenario for Fe–N–C as it is made of cheap or abundant materials. To reflect the recent synthesis of a silica templated Fe–N–C catalyst with comparable performance yet without using any HF,<sup>62</sup> we defined a separate scenario with no HF input or output. Since current Fe–N–C catalyst synthesis requires substantial amounts of electricity, we explored electricity mix based on the EU average, Poland, and Sweden, in addition to the global average electricity mix. Since production of Fe–N–C catalyst at scale would rather use a muffle furnace than a tube furnace, we considered yet another scenario that assumes the power

consumption for producing 690 g Fe–N–C with muffle furnace would be the same as producing 25 g Fe–N–C with tube furnace, which is consistent with values communicated by furnace manufacturers. Finally, we defined two scenarios of improved catalyst performance, where the required catalyst amounts were lowered to 380 g Fe–N–C and 15 g Pt/C.

## Techno-economic analysis

We conducted a dynamic cost estimation based on learning curves to determine the influence of the catalyst production cost compared with other PEMFC components on the total PEMFC stack cost at varying production scales, in line with previous PEMFC production studies.<sup>63,64</sup> We adopted a production rate ranging between 1000 and 500 000 stacks per year, where the lower rate is the feasible limit for automobile mass production and the upper rate represents an individual top selling automobile model.<sup>41</sup> Note that we only considered the stack cost, while omitting the cost of the PEMFC balance of plant.

**Dynamic cost estimation using learning curves.** A learning curve models the human activity of accumulating knowledge or experience by cumulative production. In eqn (1), the product cost at production level  $i$ ,  $Y_i$  depends on the cumulative number of products,  $X_i$ , the progress ratio,  $r$ , and a given constant  $A$ . The progress ratio  $r$  is related to the parameter  $F$ , which describes how much the production cost is reduced upon doubling the cumulative production through eqn (2).<sup>63,65</sup> For instance, an  $F$  value of 80% means for the cost to be reduced by 20% each time the cumulative production volume is doubled.

$$Y_i = AX_i^{-r} \quad (1)$$

$$F = 2^{-r} \quad (2)$$

Typical values for  $F$  range between 74–95%.<sup>63,64</sup> We adopt a nominal  $F$  value of 86.4% for the catalysts, which is based on baseline Fe–N–C (690 g) reaching the equivalent production cost of the 380 g Fe–N–C + muffle furnace scenario at 500 000 stacks per year. The values from Mock and Schmid<sup>64</sup> are applied for the rest of PEMFC components. We estimated the production costs for Pt/C and Fe–N–C from quotes from Alibaba and other relevant manufacturers, with processing costs based on calculated utility consumption and factors for plant costs from industry knowledge (Tables S10 and S11†).

For Pt/C cathode catalyst, a simple process achieving 2020 US DOE target performance is modelled with the market price of bulk precursors and electricity applied. Since the state-of-the-art de-alloyed Pt-based cathode PEMFC catalysts that meet the US DOE performance targets possess higher material and processing costs, the calculated Pt/C cathode and anode catalyst process costs are applied as the value at 500 000 stacks per year in the learning curve (Table S12†). We calculated the Pt/C anode catalyst cost assuming an equivalent production Pt/C process, but with  $0.025 \text{ mg}_{\text{Pt}} \text{ cm}^{-2}$  loading (cost divided by five). We based the costs of the other PEMFC components in the Pt/C system at 1000 stacks per year on previous values from



James *et al.* for their 2020 scenario.<sup>7</sup> For comparison, we also conducted a separate learning curve calculation based on the cost of a PtCo/C cathode catalyst from James *et al.*<sup>7</sup> (Table S13†).

For Fe–N–C cathode catalyst, market prices for bulk precursors and electricity were also used. We based the cost at 1000 stacks per year on material and process costs from the modelled Fe–N–C catalyst synthesis. We selected the production rate as 1000 stacks per year as we assumed that the performance of Fe–N–C can be improved, and thus the production costs reduced, at higher production rates. The total required coating area is 3.9-times greater for Fe–N–C than Pt/C due to the lower power output of the catalyst. Therefore, we scaled the cost of Fe–N–C PEMFC components at 1000 stacks per year in proportion with the increased surface area in the Fe–N–C system relative to the Pt/C system (Table S14†).

**Sensitivity analysis.** Historical records indicate that the price of Pt has doubled and halved over the past 20 years compared to the average price,<sup>66</sup> therefore we conducted a sensitivity analysis by varying the price of both Pt/C and Fe–N–C catalysts from –50% to +100%. Due to volatile market electricity prices, we carry out sensitivity on the electricity price with variation between –75% to +200%. We also carried out a sensitivity analysis in terms of the *F* value adopted in the learning curves (eqn (2)) between 74–95%.

**Pt demand forecasting.** Based on forecasts of PEMFC-based fuel cell electric vehicle production<sup>3,67</sup> we estimate future Pt consumption from PEMFC. We consider 10% of total annual Pt production<sup>8</sup> as the consumption limit for deployment of a new technology,<sup>68</sup> such as PEMFC for transportation. We based the analysis on 80 kW PEMFC light-duty vehicles and their Pt usages and recycling targets.<sup>29,38,61</sup>

## Results

### Life-cycle assessment

The LCA results for all seven mid-point indicators of baseline Pt/C (20 g) and Fe–N–C (690 g) PEMFC cathode catalysts are summarized on the bar charts and spider chart in Fig. 3 (see also Table S17†). Substituting Fe–N–C for Pt/C reduces impacts in the categories of mineral resource scarcity, acidification potential, and freshwater eutrophication; yet impacts are predicted to increase in the other four categories of human toxicity potential, global warming potential, fossil resource scarcity, and marine eutrophication at the same time (Fig. 3h). In particular, the global warming potential (Fig. 3a) and fossil resource scarcity (Fig. 3b) of Fe–N–C (1130 kg CO<sub>2</sub>-eq, 286 kg oil-eq) are, respectively, 70% and 35% higher than those of Pt/C (659 kg CO<sub>2</sub>-eq, 213 kg oil-eq). This is attributed to the high electricity demand from the 3-hour long two-step high temperature pyrolysis process during Fe–N–C synthesis (Tables S5 and S6†), which is typical of Fe–N–C materials and greatly exceeds the energy requirement of Pt/C production. In terms of human toxicity potential too (Fig. 3e), the impact of Fe–N–C (7140 kg 1,4-DCB-eq) is 3-times greater than for Pt/C (2300 kg

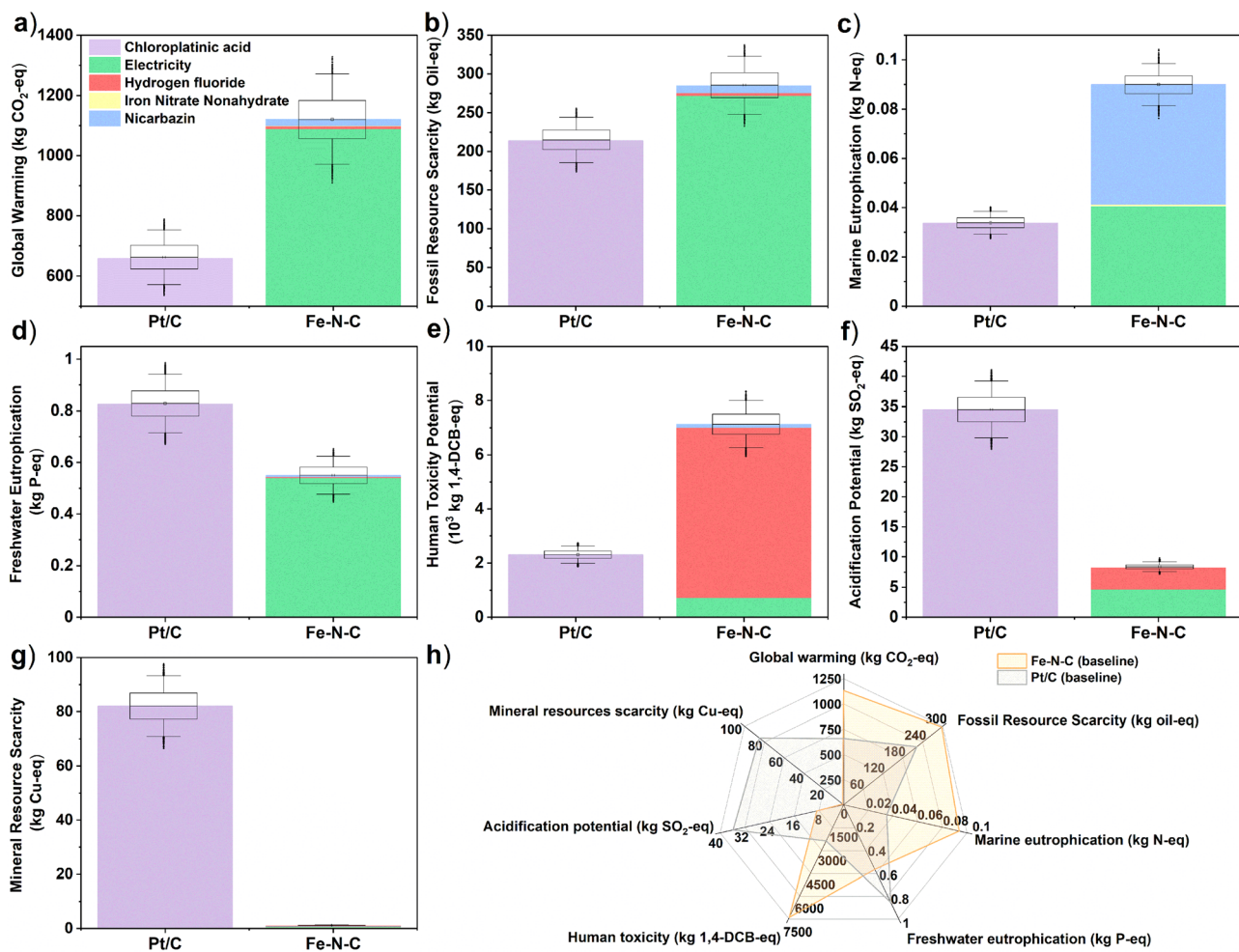
1,4-DCB-eq), mostly due to the production and disposal of large volumes of HF (Table S6†). The adoption of Fe–N–C also trebles marine eutrophication (Fig. 3c) impacts (0.09 kg N-eq) compared to Pt/C (0.03 kg N-eq) due to N release from the nicarbazin precursor (60% contribution) used in Fe–N–C synthesis. The situation is reversed for freshwater eutrophication (Fig. 3d), acidification potential (Fig. 3f), and mineral resource scarcity (Fig. 3g), where Fe–N–C impacts (0.55 kg P-eq, 8.4 kg SO<sub>2</sub>-eq, 1.0 kg Cu-eq) are reduced by, respectively, 34%, 75% and 99% compared to those of Pt/C (0.83 kg P-eq, 8.4 kg SO<sub>2</sub>-eq, 1.0 kg Cu-eq). Most Pt/C impacts in all these categories are caused by Pt mining and refining operations, which carry a significantly higher burden than obtaining the raw materials of Fe–N–C. Note that the inter-quartile ranges in predicted midpoint indicators, which are obtained by propagating the foreground inventory uncertainty, do not overlap for any of the comparisons in Fig. 3a–g, thereby asserting the conclusions. The global sensitivity analysis results indicate that the dominant factors contributing to this uncertainty are the amounts of electricity, HF and nicarbazin used for Fe–N–C production (Table S16†), and the amounts of chloroplatinic acid and electricity used for Pt/C production (Table S15†).

The LCA results for all three endpoint damage categories of human health, ecosystems quality, and resource scarcity are summarized on the bar charts in Fig. 4a–c for the baseline Pt/C (20 g) and Fe–N–C (690 g) scenario (see also Table S22†). They depict a different reality than the midpoint indicators, whereby the Fe–N–C catalyst clearly outperforms Pt/C in terms of ecosystems quality (–89%) and human health (–36%) and does marginally better in terms of resource scarcity (–9%). The midpoint contributions to endpoints (Table S19†) can help understand this shift.

Regarding ecosystems quality, the impacts of both land use ( $4.6 \times 10^{-5}$  species × year) and acidification potential ( $0.7 \times 10^{-5}$  species × year) associated with Pt/C production far exceed the dominant impact of Fe–N–C production due to global warming ( $0.3 \times 10^{-5}$  species × year). In terms of human health, the larger impact of particulate matter formation for Pt/C ( $1.4 \times 10^{-3}$  DALY) overpowers the larger climate change impact for Fe–N–C ( $1.0 \times 10^{-3}$  DALY), while human toxicity is also lower for Fe–N–C after accounting for cancerous human toxicity (conversion factor  $3.3 \times 10^{-6}$  DALY per kg 1,4-DCB eq) alongside non-cancerous human toxicity (conversion factor  $2.3 \times 10^{-7}$  DALY kg<sup>−1</sup> 1,4-DCB eq) caused by HF alone. Finally, regarding resource scarcity, the environmental burden caused by Pt mining on mineral resources is balanced by the larger burden of Fe–N–C production on fossil resources due to its much larger energy requirement.

These conclusions are confirmed by the inter-quartile ranges in predicted endpoint indicators, which show no overlap between Fe–N–C and Pt/C in the human health (Fig. 4a) and ecosystems quality (Fig. 4b) categories, and a limited overlap in the resource scarcity (Fig. 4c) category. The global sensitivity analysis finds that the dominant uncertainty factors are the amounts of chloroplatinic acid and electricity for Pt/C production (Table S20†), as well as the amounts of



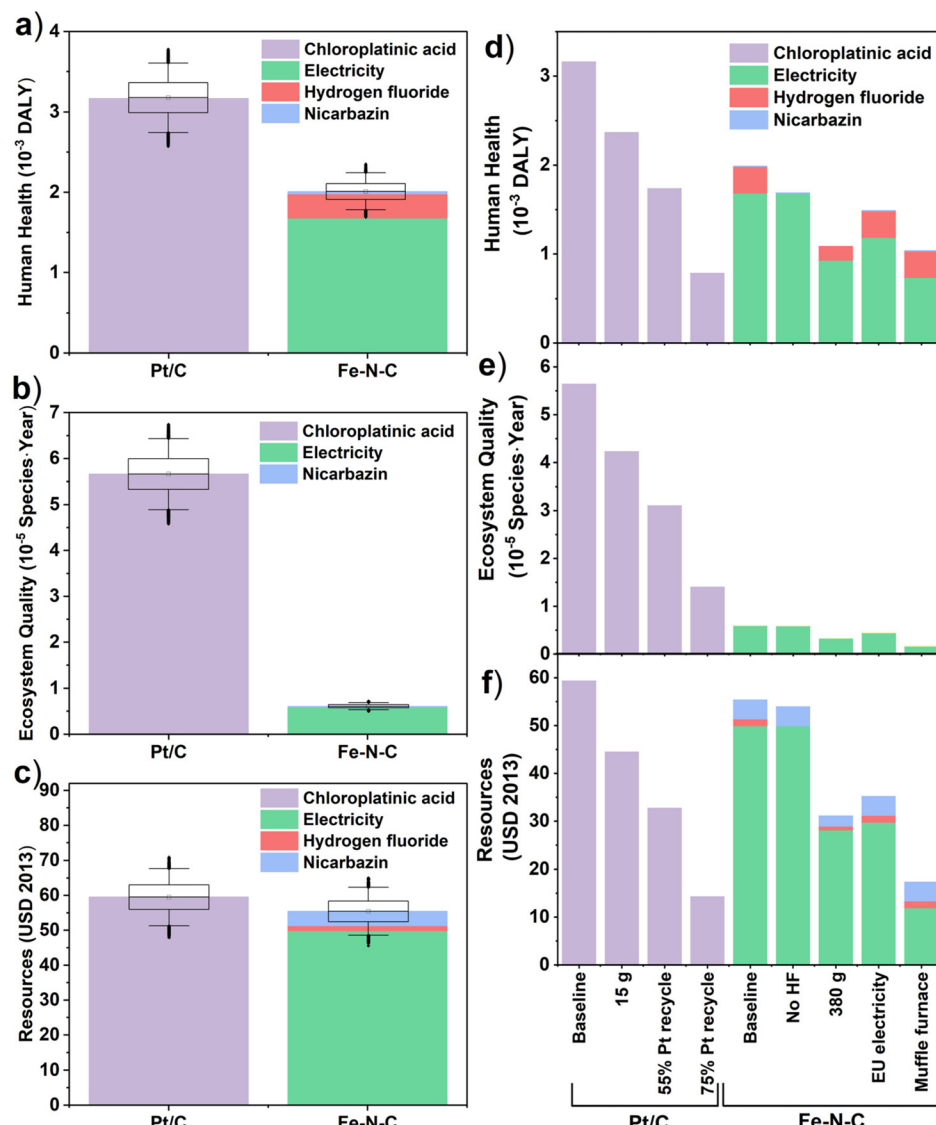


**Fig. 3** Comparison of midpoint impact indicators of baseline Pt/C (20 g) and Fe–N–C (690 g). (a) Global warming potential (kg CO<sub>2</sub>-eq). (b) Fossil resource scarcity. (c) Marine eutrophication (kg N-eq). (d) Freshwater eutrophication (kg P-eq). (e) Human toxicity potential (kg 1,4-DCB-eq). (f) Acidification potential (kg SO<sub>2</sub>-eq). (g) Mineral resource scarcity (kg Cu-eq). The main activities contributing to each mid-point indicator are reported on each bar. The boxes on each bar represent the inter-quartile range of uncertainty scenarios, with central line showing the median and central dot the mean, and the lower and upper whiskers extending the box to the 5<sup>th</sup> and 95<sup>th</sup> percentiles, outliers represented by dots. (h) Spider chart summary of baseline Fe–N–C and Pt/C midpoint indicators.

electricity and, to a lesser extent, HF for Fe–N–C production (Table S21†). Though it should be noted that end-point indicators are subject to larger uncertainty than mid-indicators,<sup>69</sup> especially regarding uncertainties in the life-cycle (background) emissions and the LCIA characterization factors, which were not quantified here. It would thus be important, as part of future work, to study in greater depth the burden-shifting occurring between the global warming impact (greater for Fe–N–C) and both end-point impacts of human health and ecosystems quality (greater for Pt/C).

The alternative scenarios of Fe–N–C and Pt/C are compared to baseline scenarios for endpoint categories in Fig. 4d-f (see Table S22† for breakdown). This is complemented with a similar comparison for midpoint categories in Fig. S1† (with breakdown in Table S17†). For Fe–N–C production, the scenario with muffle furnace presents the largest improvement potential in all three damage areas of human health (−48%),

ecosystems quality (−70%) and resources (−68%). This was expected given the dominant burden of electricity consumption on Fe–N–C impacts, but nonetheless interesting that using a muffle furnace could present even greater benefits than reducing the amount of Fe–N–C catalyst to 380 g from the baseline scenario. Swapping global to EU electricity supply could also present significant environmental benefits in terms of human health (−26%), ecosystems quality (−25%) and resources (−35%) due to the reduced carbon intensity. Although this could increase both freshwater (+16%) and marine (+11%) eutrophication potentials at the mid-point level due to the higher share of brown coal (lignite) in current EU supplies compared to the global average.<sup>49</sup> The large impact of electricity mix on the environmental performance of Fe–N–C production is further exemplified on Fig. S2 and S3† (with breakdown in Tables S18 and S23†), where a mix with high share of renewables and low carbon intensity such as Sweden



**Fig. 4** Left: comparison of endpoint impact indicators of baseline Pt/C (20 g) and Fe–N–C (690 g). (a) Human health ( $10^{-3}$  DALY). (b) Ecosystem quality ( $10^{-5}$  species  $\times$  year). (c) Resource scarcity (USD 2013). The boxes on each bar represent the inter-quartile range of uncertainty scenarios, with central line showing the median and central dot the mean, and the lower and upper whiskers extending the box to the 5th and 95th percentiles. Right: scenarios of Fe–N–C (baseline, no HF, 380 g, EU electricity) and Pt/C (baseline, 15 g, 55% and 75% Pt recycling for endpoint categories: d) Human health ( $10^{-3}$  DALY). (e) Ecosystem quality ( $10^{-5}$  species  $\times$  year). (f) Resources (USD 2013).

could dramatically reduce all three damage areas of human health (−76%), ecosystems quality (−72%) and resources (−81%), while a mix heavily reliant on coal such as Poland would significantly worsen the impacts on human health (+35%) and ecosystems quality (+69%). By contrast, the scenario without HF would bring more modest benefits, with a 16% reduction in human health impacts compared to the baseline scenario and marginal impact on other endpoints. At the midpoint level, the largest benefit of eliminating HF is on human toxicity (−88%) and acidification potential (−55%), but these two categories do not carry much weight when aggregated into the human health endpoint indicator (compare Table S19†). Regarding Pt/C production, reducing the amount

of catalyst or increasing Pt recycling has a very significant benefit in all environmental impact categories, which agrees with the dominant burden of chloroplatinic acid on Pt/C impacts. Within the best-case, 75% Pt recycling in particular, reductions from the baseline Pt/C scenario are close to 75% in all endpoint and midpoint impact categories. The 75% Pt recycling scenario is also more favorable than any of the Fe–N–C improvement scenarios in terms of impacts on human health and resources, although Fe–N–C still carries significantly less burden on endpoint ecosystems quality and midpoint acidification potential and mineral resource scarcity compared to any of the Pt/C scenarios. It is also noteworthy that the predicted impacts on human health and resources in the muffle

furnace scenario of Fe–N–C production remain within 20–30% of those of the optimistic 75% Pt recycling scenario, confirming the large improvement potential of Fe–N–C catalyst.

### Techno-economic analysis

Material, utility, and plant costs of cathode catalysts are summarized in Tables S10 and S11.<sup>†</sup> The material costs for Fe–N–C are only 50.6 USD kg<sub>Fe–N–C</sub>, with over 70% coming from the N–C precursor (nicarbazin). Meanwhile Pt/C material costs are several hundred times higher at 13 600 USD kg<sub>Pt/C</sub>, with 98% of the cost from the Pt precursor (chloroplatinic acid). Normalizing material costs per kW PEMFC cathode, Fe–N–C is calculated as 0.44 USD per kW, while Pt/C is 3.39 USD per kW (Fig. S4a<sup>†</sup>). If Pt recycling was implemented the Pt/C cost could be comparable to Fe–N–C, and the material feedstock cost could also become comparable to baseline Fe–N–C if 75% Pt recycling was achieved (0.85 USD per kW). As expected from these results, Pt/C production costs show an approximate 1 : 1

ratio with feedstock price variance, while Fe–N–C is relatively insensitive (Fig. S4b<sup>†</sup>). If the price of Pt were to double, as occurred in 2008, while Fe–N–C feedstock prices remained the same, the material cost of the cathode PEMFC Pt/C would become 15-time higher than Fe–N–C. On the other hand, Fe–N–C production costs are highly sensitive to electricity price variance while Pt/C is not (Fig. S4c<sup>†</sup>).

Learning curve analysis of Pt/C and Fe–N–C is shown in Fig. 5a and b (see Tables S12 and S14<sup>†</sup> for details), with Fe–N–C and Pt/C PEMFC component breakdown shown in Fig. S5.<sup>†</sup> At 500 000 stacks per year, the Pt/C cathode makes up 25% of the PEMFC stack cost and 3.4 USD per kW, while Fe–N–C makes up only 2% of stack cost and is 0.79 USD per kW. This result relates to the total PEMFC stack cost being far lower for Pt/C than Fe–N–C cathode stacks, at 13.8 USD per kW compared to 41.6 USD per kW, respectively. This difference is caused by the 3.9-times greater required surface area of the Fe–N–C cathode system, which was assumed to increase the

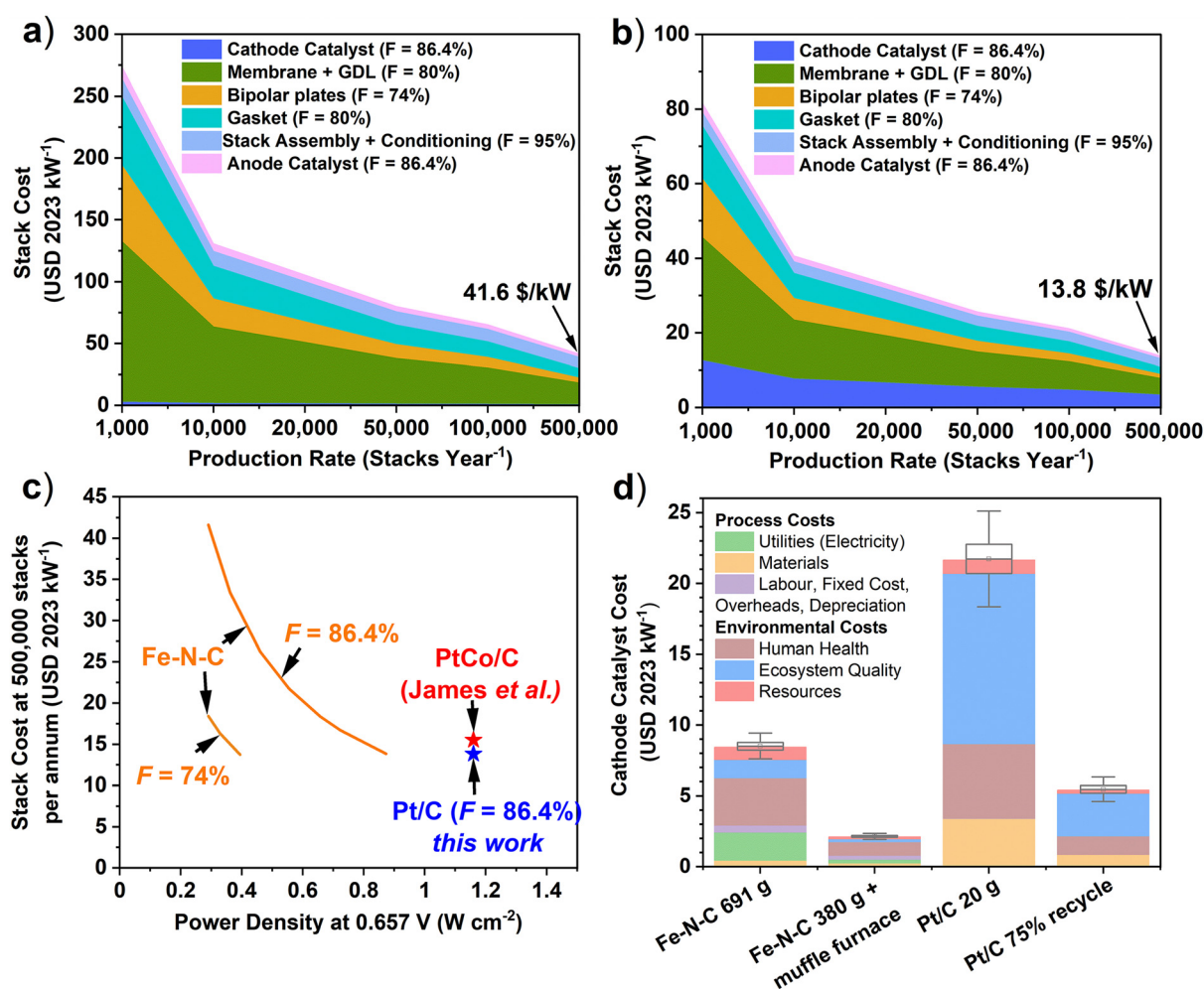


Fig. 5 Learning curve analysis of 80 kW PEMFC stack cost with production rate for baseline cathode catalysts with  $F = 86.4\%$  (a) Fe–N–C (690 g<sub>FeNC</sub>). (b) Pt/C (20 g<sub>Pt/C</sub>). (c) Variation of PEMFC stack cost at 500 000 stacks with power density for different  $F$  values for Fe–N–C compared to target Pt-based 80 kW PEMFC systems, including PtCo/C of James *et al.* (2018)<sup>7</sup> (d) Breakdown of the total process and environmental production costs for baseline Fe–N–C (690 g<sub>Fe–N–C</sub>), Pt/C (20 g<sub>Pt/C</sub>), Fe–N–C 380 g<sub>Fe–N–C</sub> + muffle furnace and 75% Pt recycling scenarios. Box and whisker plots are drawn using the endpoint uncertainty scenarios, with monetization applied.

costs of other components by the same factor. This increased area completely negates the benefit of the reduced cathode catalyst cost in the Fe–N–C system. For instance, in a Fe–N–C cathode based PEMFC, the Pt-based anode could end up using more Pt than in an all Pt-based PEMFC, if the Pt consumption at the anode scaled with area.

The PEMFC stack cost is therefore sensitive to the Fe–N–C power density performance, as well as the  $F$  value applied, which describes how much the production cost is reduced upon doubling the cumulative production (Fig. S6†). For the modelled system at  $F = 86.4\%$  for the catalysts, the baseline Fe–N–C would need to reach a power density of  $874 \text{ mW cm}^{-2}$  at  $0.657 \text{ V}$ , equivalent to a 200% improvement (reduction to  $230 \text{ g}_{\text{Fe}-\text{N}-\text{C}}$ ) to become cost competitive with the Pt/C based cathode at 500 000 stack per annum (Fig. 4c). At  $F = 74\%$  for all Fe–N–C based PEMFC components, only a 34% improvement to  $0.392 \text{ mW cm}^{-2}$  is required to reach equal costs to Pt/C based system. However, the Fe–N–C stack with  $F = 95\%$  cannot achieve the same Pt/C stack cost (with  $F = 86.4\%$ ) for the power density range considered here. A simple and cheap Pt/C cathode process was modelled here; however, to achieve US DOE performance targets, de-alloyed Pt-based catalysts are typically required. Therefore, the cost of an equivalent performance but more expensive Pt-based de-alloyed cathode catalyst (PtCo/C) was also evaluated (Table S13 and Fig. S7†). De-alloyed PtCo/C is more expensive due to additional processing steps.<sup>7</sup> At 500 000 stacks per year, the higher cost of the PtCo/C precursor ( $3.93 \text{ USD per kW}$ ), which makes up 27.4% of component costs, results in PEMFC stack cost of  $14.3 \text{ USD per kW}$ . This is lower than the  $15.5 \text{ USD per kW}$  calculated by James *et al.* for a 2020 PEMFC auto system with PtCo/C,<sup>7</sup> owing to the learning curve rate applied here. For baseline Fe–N–C to reach equivalent cost to the PtCo/C cathode PEMFC considered here at 500 000 stacks per year would require reaching a power density of  $0.842 \text{ W cm}^{-2}$  at  $0.657 \text{ V}$ , or  $0.781 \text{ W cm}^{-2}$  for the PtCo/C system of James *et al.* (Fig. 5c).<sup>7</sup>

Incorporating monetisation of environmental impacts into the catalyst material and processing costs has a significant impact on the cathode catalyst cost (Fig. 5d). Though it is worth reiterating that the monetisation factors used (Table S7†) carry large uncertainty, so the monetised impacts may only provide an order of magnitude estimate.<sup>52</sup> For all catalyst scenarios, the environmental externalities are higher than the material and processing costs. For baseline Fe–N–C, these environmental externalities add up to  $5.50 \text{ USD per kW}$ , while material and processing costs are  $2.92 \text{ USD per kW}$ . The case is even more severe for baseline Pt/C, with environmental externality costs of  $18.23 \text{ USD per kW}$ , against material and process costs of  $3.41 \text{ USD per kW}$ . Pt/C has a much larger impact on ecosystem quality than Fe–N–C, which leads to the largest monetised externality of  $12.03 \text{ USD per kW}$  (Fig. 3). Taking improved Fe–N–C ( $380 \text{ g}_{\text{Fe}-\text{N}-\text{C}} + \text{muffle furnace}$ ) scenario leads to a reduced total process and environmental externality cost of  $2.10 \text{ USD per kW}$ . Additionally, for 75% Pt recycling scenario, the total cost falls to  $5.41 \text{ USD per kW}$ . Costs associated with labour, fixed costs, overheads, and deprecia-

tions are negligible for Pt/C due to only  $20 \text{ g}_{\text{Pt/C}}$  production required for an  $80 \text{ kW}$  PEMFC, whereas  $690 \text{ g}$  of Fe–N–C results in  $0.49 \text{ USD per kW}$  for these costs. Moreover, electricity costs for baseline Fe–N–C are significant at  $1.98 \text{ USD per kW}$ , and insignificant for Pt/C, although Fe–N–C electricity requirements can be reduced to  $0.28 \text{ USD per kW}$  for the  $380 \text{ g}_{\text{Fe}-\text{N}-\text{C}}$  with muffle furnace.

Finally, we consider if Pt supply can meet future scenarios of Pt demand from PEMFC deployment for FCEV production. This topic has previously recently been considered in detail,<sup>70,71</sup> and we here provide a new perspective on this topic (Fig. S8 and S9†). We based our criterion on Pt demand not exceeding 10% of supply for a new technology.<sup>8</sup> If the sum of global Government targets for FCEV deployment by 2030 (2.5 million FCEV) were to be produced in a single year, Pt consumption would still not exceed 10% Pt supply based on US DOE 2020 Pt specific power density targets values, even with zero Pt recycling. If EU 2030 Pt specific power density and Pt recycling targets were met, Pt supply constraints would not be a limiting factor in PEMFC production, even if the highest 2040 target of FCEV on the road (15 million FCEV<sup>67</sup>) were to be produced in one year. Nevertheless, this analysis does not factor in that the majority of Pt is produced by a single country, South Africa, and the supply of Pt could therefore be subject to disruptions.

## Discussion

Following results analysis, in this section we provide a discussion of the results in context with previous works and their methods. For global warming normalized to power, Pt/C was calculated to emit  $8.4 \text{ kg CO}_2\text{-eq per kW}$  based on  $0.125 \text{ mg}_{\text{Pt}} \text{ cm}^{-2}$ . Previous reports (summarized in Fig. S8†) calculated between  $6.5\text{--}14.6 \text{ CO}_2\text{-eq per kW}$ , with catalyst loading ranging from  $0.15\text{--}0.60 \text{ mg}_{\text{Pt}} \text{ cm}^{-2}$ .<sup>22,23,42,72</sup> Here, the impact of the preparation of the catalyst inks for deposition was not considered. Differences in environmental impacts also result from changes in impact factors across background database processes, different assumed Pt-based catalyst performance and different Pt compositions. The Pt catalyst has previously been found to make up 6–24% of the total global warming contribution for a fuel cell stack.<sup>22,42,72</sup> As found by Usai *et al.*,<sup>72</sup> the Pt-based catalyst is significant for all other impact categories (33–76%), although other components contribute to the environmental impact of manufacturing PEMFC vehicles as well. For instance, Evangelisti *et al.*<sup>42</sup> previously found that, even with zero Pt loading, fuel cell vehicles in the manufacturing phase would have a higher environmental impact than internal combustion engine vehicles.

The heating temperature, heating rate, time, and number of pyrolysis steps for Fe–N–C catalysts needs to be carefully considered, since this energy intensive processes makes up a significant share of all environmental impact categories through the electricity consumption. James *et al.*<sup>32</sup> estimated that the manufacturing costs in a poly-aniline derived Fe–N–C, produced at 500 000 systems per year, are largely derived from the



acid washing process (49%), while the carbon support makes up the most significant material cost (34%). At this production rate, they estimated the Fe–N–C cost at 74–129 USD per kg,<sup>32</sup> which is comparable to our study (90.8 USD per kg with  $F = 86.4\%$ ). In their study the 1-hour pyrolysis in a rotary kiln only contributed 3% of manufacturing costs, while we scaled the electricity input for Fe–N–C manufacturing process with pyrolysis steps based on a tube furnace process here. In particular, we found that vast reductions in electricity requirement, and therefore cost and environmental impacts of Fe–N–C, could be achieved by using a muffle furnace (Fig. 3d–f). Further energy reductions could also be made when producing Fe–N–C at scale using heat integration methods such as pinch analysis. These reductions would lead the low technology readiness of Fe–N–C catalyst to reach more comparable electricity consumption to the high technology readiness optimized Pt/C process.

With regards to the carbon intensity of electricity, it is worth noting that the EU aims for its electricity production to reach carbon-neutrality by 2040. In this scenario, the impact of Fe–N–C on climate change, human health, ecosystems quality and resources all together would all become negligible compared to those of Pt/C. Already in countries with a high share of renewables in the electricity mix, such as Sweden (where commercial Fe–N–C manufacturer Celcibus AB is based), the environmental impacts of Fe–N–C production would be greatly mitigated (Fig. S2 and S3†). The impact on human toxicity could furthermore be avoided through alternative production methods (without HF, Fig. S1e†).<sup>62</sup>

Considering degradation, Stropnik *et al.* recently developed LCA models that incorporate degradation effects,<sup>73</sup> and James *et al.* recently incorporated durability adjusted operating conditions for TEA models.<sup>74</sup> We did not consider degradation of the PEMFC here. The reason for this is practical light duty automotive application would need to reach the US DOE target of 8000 h,<sup>38</sup> or 7000 h according to EU 2030 targets,<sup>61</sup> while to date high activity Fe–N–C have only been demonstrated for 100s of hours in PEMFCs.<sup>14</sup> Nevertheless, recent work suggests Fe–N–C catalysts can be partially regenerated *in situ* for short periods with simple electrochemical protocols,<sup>75</sup> rather than requiring an *ex situ* recovery process as required for Pt. Fe–N–C based catalysts could also find applications in small, short term back-up power applications (at high production volumes), such as the emergency power/Wi-Fi backpack recently demonstrated by Ballard System Ltd and Nisshinbo.<sup>76</sup>

Considering Pt-based PEMFCs, over an order of magnitude reduction has been achieved in the past three decades (Fig. S9†). For instance, Toyota has achieved 72% reduction from  $0.93 \pm 0.25$  to  $0.26 \pm 0.07 \text{ g}_{\text{Pt}} \text{ kW}^{-1}$  in their 2008 PEMFC model to 2014 1<sup>st</sup> generation Mirai, and a further 58% reduction to  $0.11 \pm 0.03 \text{ g}_{\text{Pt}} \text{ kW}^{-1}$  in their 2020 2<sup>nd</sup> generation Mirai.<sup>4</sup> The EU targets a reduction in PEMFC Pt content to  $0.05 \text{ g}_{\text{Pt}} \text{ kW}^{-1}$  by 2030,<sup>61</sup> although increased Pt instability at these low future target loadings may prevent their realization.<sup>77</sup> Nevertheless, combining ideal scenarios of Pt reduction and recycling targets being met would result in minimal environmental impacts.

In terms of learning curve analysis, typical  $F$  values range between 74–95%<sup>63,65</sup> and 86.4% was selected as the nominal value here. We highlighted in the Results section that the  $F$  value has a significant impact on the stack costs. It is difficult to predict what  $F$  value would be realized; however, based on the previous minimum  $F = 74\%$ , an 80 kW Fe–N–C based PEMFC would need to improve at least 34% from 290 mW  $\text{cm}^{-2}$  to become cost comparative with Pt/C based delivering 1160 mW  $\text{cm}^{-2}$ . Current non-precious metal hydrogen evolution catalysts for proton exchange membrane electrolyzers also show a similar requirement of improved performance under most operating regimes to replace precious metal catalysts.<sup>78</sup>

Future work could consider the effect on Fe–N–C LCA for other common synthetic strategies, such as recent state-of-the-art Fe–N–C derived from zeolitic imidazolate framework 8.<sup>14</sup> Broader environmental contextualization in terms of planetary boundaries or comparison to other technologies, such as batteries, would also provide further critical insights. Future comparison of the use and disposal phase of the catalysts could be developed based on degradation<sup>73</sup> and recycling.<sup>27</sup> Design for manufacture and assembly methodology could be used to track annual cost impact of catalyst developments.<sup>74</sup>

## Conclusions

In this paper, we compared scenarios in the LCA and TEA for manufacturing a silica templated Fe–N–C and a Pt/C cathode catalyst to form 80 kW PEMFCs stack. Electricity consumption and Pt mining are by far the dominant factors for environmental impacts and costs in Fe–N–C and Pt/C catalysts, respectively. Applied monetization factors show the environmental impact cost to be greater than material and processing costs for both catalysts. At 500 000 stacks per year, moving from Pt/C to Fe–N–C could reduce the cathode catalyst material and processing cost from 2.92 to 0.79 USD per kW. However, the increased cost of all other components from the ~4-times greater surface area of the Fe–N–C cathode system results in a PEMFC stack cost of 41.6 USD per kW, compared to 13.8 USD per kW for Pt/C. For the nominal  $F$  value of 86.4%, Fe–N–C based PEMFC would need to reach 874 mW  $\text{cm}^{-2}$  at 0.657 V, equivalent to a 200% performance improvement, to reach the same PEMFC stack cost with the  $0.125 \text{ mg}_{\text{Pt}} \text{ cm}^{-2}$  Pt/C cathode catalyst delivering 1160 mW  $\text{cm}^{-2}$ . This finding highlights further improvements in activity for Fe–N–C cathode catalysts are required, while the large discrepancy in durability between Fe–N–Cs and Pt/C has not been considered. Finally, we consider if Pt supply can meet future scenarios of Pt demand from PEMFC deployment for FCEV production.

## Author contributions

Conceptualization: J. P., A. P., C. P., M.-M. T. Data curation: J. P., A. P., G. L., A. S., A. B. Formal analysis: A. P.,



J. P. Investigation: J. P., A. P., G. L. Methodology: B. C., C. P. Resource: A. S. Software: B. C., G. L., A. B. Supervision: B. C., C. P., I. E. L. S., M.-M. T. Project administration: A. P., J. P. Validation: A. P., B. C., J. P. Visualization: A. P., B. C., C. P., J. P. Writing – original draft: A. P., J. P. Writing – review & editing: all authors. A. P. and J. P. contributed equally to the article.

## Conflicts of interest

There are no conflicts to declare.

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