

On-demand manufacture of circular 3D-printable resins†

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Recent progress in circular 3D-printable photocurable resins that enable closed-loop recycling marks a significant step forward in reducing wasteful manufacturing methods and non-recyclable printed plastics. However, with 3D printing technologies shifting from prototyping to full-scale production, the demand for high-scale processes and easily tuneable resin compositions can benefit from the design of automated systems. Herein, we report on the on-demand preparation of a circular 3D-printable resin, achieved through a continuous flow approach. A supported enzyme (Lipase B from *Candida antarctica*) was used to promote a green esterification of the lipoic acid with biobased alcohols to prepare circular biobased photocurable resins. The supported enzyme was employed for the preparation of a packed bed reactor and was easily recycled and reused to achieve the continuous production of lipoate-based photocurable resins with tuneable composition. Lastly, the environmental impact of the developed on-demand manufacturing process was compared to the previously reported esterification protocols through life cycle assessment, showing the effectiveness of continuous enzymatic flow synthesis in enhancing environmental performance across multiple areas, from human health to ecosystem impact and resources.

Introduction

Vat photopolymerisation is a widely adopted additive manufacturing (3D printing) method that relies on light-activated polymerisation to selectively solidify liquid resins held in a vat.^{1,2} The additive manufacturing market, currently valued at \$0.9 billion, accounts for a significant share of the broader photopolymer market, which exceeds \$2.8 billion in total, and it is projected to expand at a compound annual growth rate of 16.9% as photopolymer-based 3D

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printing technologies become faster and shift from prototyping to full-scale production.³ However, this growth must be accompanied by advancements in photopolymer resin technology to ensure that additive manufacturing contributes to a truly sustainable and circular plastic economy.

Most resin components are derived from non-renewable fossil resources, and many resin-based processes generate waste in the form of partially cured material that cannot be reused.^{4–7} Additionally, the majority of 3D-printed objects made from permanently cross-linked resins are neither degradable nor recyclable, further exacerbating environmental damage and creating significant end-of-life challenges. To tackle this issue, closed-loop recycling of 3D-printed polymers, in which the printed material is fully depolymerised to a resin and re-printed, has been explored. Circular photocurable resins have been focused on the introduction of dynamic bonds that can break down under certain conditions within permanently cross-linked polymer networks, allowing partial depolymerisation of printed parts.^{8–16} Recently, we reported on circular lipoate-based resins entirely obtained from renewable feedstocks that can be 3D-printed into high-resolution parts, efficiently deconstructed, and subsequently reprinted in a circular manner.¹¹ Yang *et al.* have also reported a circular additive manufacturing process to produce high-performance polymer parts containing dithioacetal bonds that can be decomposed into reprintable liquid resins under a specific condition.¹⁶

3D printing has been successfully employed to prepare sophisticated architectures in flexible devices and multimaterial objects that can have varied properties, including conductivity, self-healing capacity, and recyclability.^{17–19} Through vat photopolymerisation, it is possible to achieve the rapid production of customised 3D-printed components. Tuneable resins are conventionally formulated by varying the ratio of monomer manually,^{20,21} whereas the need to move to a large-scale production and the demand for precise control over resin composition can benefit from the design of automated systems. In this regard, the flow approach can be leveraged to develop an automated method to prepare photocurable resins with easily tuneable composition. Applying flow chemistry techniques to synthesis offers several advantages over traditional batch synthesis methods.^{22–25} Not only does it allow for greater repeatability and easier scale-up, but it can also be tuned to deliver bespoke products (*e.g.* resin with specific composition).^{26,27} While flow chemistry has largely benefited from the use of 3D printing (*e.g.* to print continuous flow reactors),²⁸ there have been no reports of using flow systems to directly synthesise photopolymer resins for 3D printing.

Herein, we report on the preparation of an on-demand circular biobased resin by leveraging continuous flow production (Fig. 1). Lipoate-based

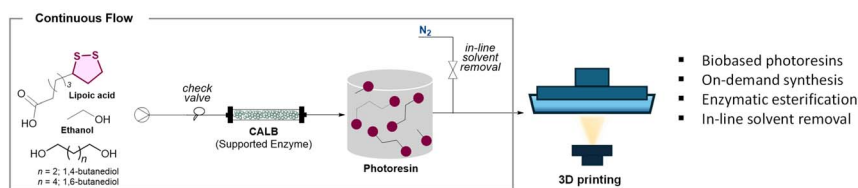


Fig. 1 On-demand photocurable resin synthesis under continuous flow conditions followed by off-line 3D printing developed in this work.



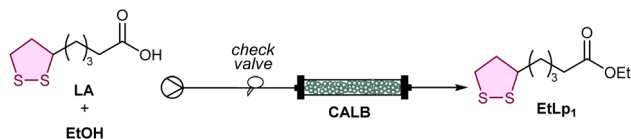
photocurable resins have been synthesised using a sustainable enzymatic esterification. The environmental feasibility of the circular resin production under continuous flow conditions was lastly assessed through life cycle assessment analysis, proving it to be a more sustainable strategy than the previously reported synthetic protocols.

Results and discussion

Enzymatic esterification under continuous flow conditions

Liquid resin formulations employed in 3D printing are usually made of a reactive diluent and multivalent crosslinkers. The reactive diluent, ethyl lipoate (EtLp₁) synthesis was carried out under continuous flow conditions (Table 1), exploring a range of solvents including dichloromethane (CH₂Cl₂), and greener ones, such as 2-methyl tetrahydrofuran (2-MeTHF) and methyl ethyl ketone (MEK). A single syringe system was used by combining ethanol (1.05 equiv.) and a solution of lipoic acid (LA) in CH₂Cl₂ (2.42 M). Initially, a packed-bed reactor was prepared using 0.5 g of supported Lipase B from *Candida antarctica* (CALB), and the reagents were flowed at a flow rate of 0.1 mL min⁻¹, resulting in a residence time of 25 minutes (see ESI† for details). Fractions of the EtLp₁ product were collected at 1 h intervals, and their composition was monitored using ¹H NMR spectroscopy. Under these conditions, a 65% EtLp₁ yield was detected when the steady state was achieved (Table 1, entry 1). To minimise the amount of the solvent employed in the process, the concentration of the solution was doubled; however, that led to a gelation of the solution within the reactor, making that condition impracticable for the production of photocurable resins (Table 1, entry 2).

Table 1 Screening of the reaction conditions for the enzymatic synthesis of EtLp₁ under continuous flow conditions^a



Entry	Catalyst/g	Solvent	Flow rate/mL min ⁻¹	τ/min	EtLp ₁ /%
1	0.5	CH ₂ Cl ₂	0.1	25	65
2 ^b	0.5	CH ₂ Cl ₂	0.1	25	—
3	0.8	CH ₂ Cl ₂	0.1	35	88
4	0.8	CH ₂ Cl ₂	0.07	50	94
5	0.8	2-MeTHF	0.07	40	70
6	0.8	2-MeTHF	0.05	65	85
7	0.8	MEK	0.07	50	73
8	0.8	MEK	0.05	80	72
9 ^c	2.0	CH ₂ Cl ₂	0.11	80	95
10 ^c	2.0 + (0.5 g MgSO ₄)	CH ₂ Cl ₂	0.1	80	96

^a Reaction conditions: LA (1 equiv.), EtOH (1.05 equiv.), solvent (2.42 M), CALB (x g), flow rate, rt. ^b Solvent (4.85 M). ^c EtOH (1.20 equiv.). τ stands for residence time.



To increase the conversion of LA into the corresponding EtLp₁, the catalyst amount and flow rate were then optimised. The catalyst amount was initially increased to 0.8 g, leading to an increased residence time of 35 min. Under these conditions, an 88% EtLp₁ yield was obtained using a 0.1 mL min⁻¹ flow rate (Table 1, entry 3). The results observed were consistent with previously reported experiments on continuous esterification, as conversion enhancement is seen with increasing the catalyst bed height.²⁹ A further increase in EtLp₁ yield (94%) was achieved by lowering the flow rate to 0.07 mL min⁻¹ (Table 1, entry 4).

With the aim of designing a sustainable and scalable esterification protocol, alternative green solvents with low boiling points were also explored. The use of 2-MeTHF as the solvent resulted in a lower EtLp₁ yield (70%, Table 1, entry 5) at a 0.07 mL min⁻¹ flow rate, while a decrease to 0.05 mL min⁻¹ led to an EtLp₁ yield of 85% (Table 1, entry 6). The use of 2-MeTHF led to reduced process efficiency, most likely a consequence of its inferior resin swelling properties compared to CH₂Cl₂, which in turn resulted in less effective mass transfer (Fig. S12 and S13†). MEK was also tested, achieving 73% EtLp₁ formation at room temperature with a 0.07 mL min⁻¹ flow rate (entry 7, Fig. S14†). Decreasing the flow rate to 0.05 mL min⁻¹ showed no improvement (Table 1, entry 8).

To explore the process scalability, a large-scale reactor was prepared with 2 g of supported enzyme. The scale-up was tested using CH₂Cl₂, and the results achieved were consistent with the small-scale experiments (Table 1, entry 9). Moreover, the use of drying agents (*e.g.*, MgSO₄ or molecular sieves) to suppress ester hydrolysis during esterification yielded negligible improvement (Table 1, entry 10). Finally, to understand the stability of the developed system, a long-run experiment was performed. Fractions were collected every hour, and EtLp₁ formation was stable for 13 h, with more than 36 g of EtLp₁ produced in that time frame (Fig. 2a).

To explore the preparation of photocurable resins under flow conditions, multivalent crosslinkers were then synthesised following the same protocol used for the reactive diluent, EtLp₁. Previously reported lipoic acid-based crosslinkers¹¹ were initially targeted. Glycerol and sugar-based isosorbide and isomanide were tested; however, their enzymatic esterification at room temperature was inefficient. Hence, alternative biobased alcohols, 1,4-butanediol and 1,6-hexanediol, were considered.^{30,31} The synthesis of the hexyl lipoate derivative (HexLp₂) was carried out by combining 1,6-hexanediol (1 equiv.) and a solution of LA in CH₂Cl₂ (2.42 M). Reagents were initially supplied at a flow rate of 0.1 mL min⁻¹ and fractions were collected at 1 h intervals and analysed using ¹H NMR spectroscopy. Under these conditions, HexLp₂ was obtained with an 86% yield (Fig. 2b), while using a flow rate of 0.08 mL min⁻¹ was shown to lead only to a slightly higher yield (89%). Lastly, decreasing the flow rate to 0.05 mL min⁻¹ showed consistent data with higher flow rates, with only a marginal increase in yield (90% yield of HexLp₂).

Subsequently, the synthesis of butyl lipoate ester (ButLp₂) was carried out under flow conditions using the same conditions optimised for HexLp₂ by combining 1,4-butanediol (1 equiv.) and a solution of LA in CH₂Cl₂ (2.42 M). Using a large-scale reactor (with 2 g of supported enzyme) with a flow rate of 0.1 mL min⁻¹ led to ButLp₂ being obtained with an 84% yield when the steady state was reached (Fig. 2c).



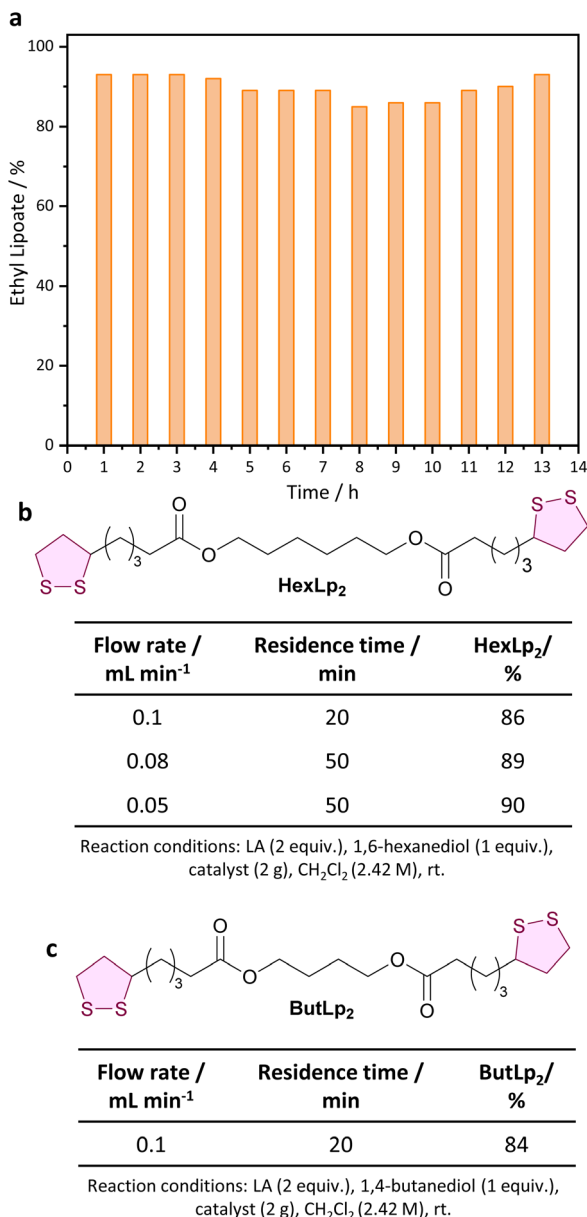


Fig. 2 (a) EtLp₁ long-run continuous flow preparation (reaction conditions: Table 1, entry 4). (b) Screening of the reaction conditions for the enzymatic synthesis of multivalent crosslinkers HexLp₂. (c) Reaction conditions for the enzymatic synthesis of multivalent crosslinkers ButLp₂.

Photocurable resin preparation under continuous flow conditions

Photopolymer resins were prepared by combining the reactive diluent (EtLp₁) and multivalent crosslinker (HexLp₂ or ButLp₂) made by continuous flow enzymatic esterification (Fig. 3a). EtLp₁ and HexLp₂ were both prepared under continuous



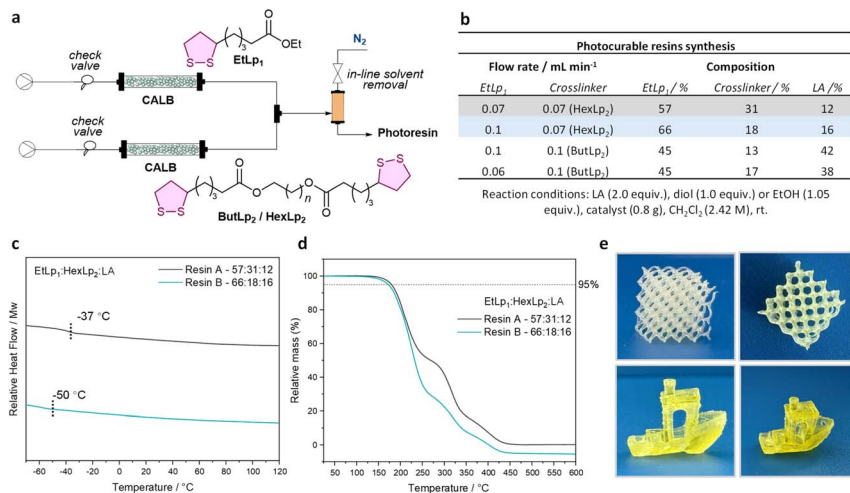


Fig. 3 (a) Continuous flow set-up for photocurable resin preparation. (b) Composition of the combined photocurable resin obtained under continuous flow conditions. (c) DSC thermograms of the second heating cycle of the EtLp₁:HexLp₂ resins (−80 to 130 °C, 10 °C min^{−1} under N₂). *T_g* is indicated by the dashed line. (d) TGA thermograms of the EtLp₁:HexLp₂ resin 2D-photoset post-cured samples (25–600 °C, N₂ atmosphere). *T_{d5%}* is indicated by a horizontal line. (e) 3D-printed complex parts, top and front views.

flow conditions in separate flow reactors, each packed with the supported enzyme (0.8 g) and initially set to a flow rate of 0.07 mL min^{−1}. The outlet of each reactor was connected at a T-junction, and the mixed photocurable resin was collected in a separating funnel. Nitrogen was bubbled through the collected solution while the resin was being prepared to ensure that the final product had minimal solvent content for the following printing process. Aliquots were taken at 1 h intervals and the ratios of EtLp₁, HexLp₂ and lipoic acid of the combined resin were determined using ¹H NMR spectroscopy. As lipoic acid is not fully converted into the ester derivative, we anticipated that the unreacted LA could be incorporated into the material when photocured.

To determine whether the composition of the resin could be tuned *via* the continuous flow parameters, the flow rate of the EtLp₁ process was increased from 0.07 mL min^{−1} to 0.1 mL min^{−1}. It was hypothesised that increasing the flow rate would result in a higher amount of EtLp₁ as there would be a higher volume of EtLp₁ in the combined resin compared to HexLp₂. On the other hand, increasing the flow rate would also increase the amount of lipoic acid in the final photocurable resin as a lower conversion of EtLp₁ would be achieved with the lower residence time. The resulting resin composition showed that increasing the flow rate of EtLp₁ does lead to an increase in the amount of EtLp₁ and lipoic acid in the final resin and a decrease in the amount of the crosslinker HexLp₂ (Fig. 3b). In doing so, we demonstrated that changing the continuous flow parameters enables the final resin composition to be tuned. This represents a novel approach to tune photocurable resin compositions and therefore prepare on-demand tailored photocurable resins in an automated way. Similarly, a EtLp₁:ButLp₂:LA photocurable resin was prepared with an analogous set-up. Due to the higher



flow rates employed relative to the EtLp₁ : HexLp₂ resin preparation, the EtLp₁ : ButLp₂ resin exhibited an increased lipoic acid content (Fig. 3b). This outcome was anticipated as higher flow rates result in lower conversions.

To explore the material properties of the EtLp₁ : HexLp₂ : LA photopolymer resins, 2D photosets of the resins were prepared by curing 0.1 mL of resin on glass slides (10 min UV irradiation), followed by post-curing for 24 h at 60 °C (Fig. S15†). Differential scanning calorimetry (DSC) and thermogravimetric analysis (TGA) of the obtained photocured polymers were conducted. The glass transition temperature (T_g) for the resin containing a 57 : 31 : 12 ratio of EtLp₁ : HexLp₂ : LA (resin A) was −37 °C, whereas the T_g for the resin containing a 66 : 18 : 16 ratio of EtLp₁ : HexLp₂ : LA (resin B) was −50 °C (Fig. 3c). The difference in T_g can be ascribed to the higher content of HexLp₂ crosslinker, which leads to a higher degree of crosslinking. This proof-of-concept experiment shows that the material properties of the resins can be tuned by preparing the photopolymer resins under flow conditions. Lastly, both resins showed thermal stability up to 170 °C (Fig. 3d), in line with lipoic acid-based photopolymers previously reported.¹¹

To explore the structure–property space of the platform, additional samples of the resin were prepared using ButLp₂. However, we observed a partial gelation of the formulated photocurable resin as soon as the solvent was removed, which proved to be unsuitable for the following application in additive manufacturing.

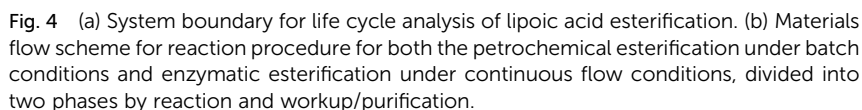
Printing the resin EtLp₁ : HexLp₂ : LA (57 : 31 : 12) on a commercial digital light processing (DLP) printer was achieved using a $\lambda = 405$ nm light source (Fig. 3e). The high-fidelity, complex prints were obtained using a 35 s cure per layer, which corresponds to a build rate of 5.1 mm h^{−1} (excluding the peeling time process), demonstrating that the resins prepared through the continuous flow method remained suitable for DLP printing.

life cycle assessment and green chemistry metrics

A first estimation of the sustainability of the preparation of circular lipoic acid-based resins was explored using green chemistry metrics (Table S1†). The enzymatic reaction is considered to be a more sustainable approach, with chemicals such as *N*-(3-dimethylaminopropyl)-*N'*-ethylcarbodiimide (EDC), commonly used in esterification reactions, replaced by the supported enzyme. Additionally, the continuous flow approach minimises the purification process, limiting it to the sole solvent removal. For the proposed enzymatic process, the atom economy (AE)³² was equal to 93, meaning that most of the reagents are converted into the products with little waste production. In contrast, the previously used EDC coupling¹¹ pathway (here named as petrochemical process) resulted in a significantly lower value as a consequence of the high amount of waste produced and the lower efficiency of the process (78% yield for the petrochemical process *vs.* 94% for the enzymatic process). The *E* factor³³ for the enzymatic process was calculated to be <0.01, falling into the range of oil refinery processes, while it is equal to 2.57 when the solvent is considered, showing a comparable value with the recently reported Fisher esterification process.³⁴ The use of EDC led to a higher *E* factor (17.04 when the solvent is considered, Table S1†), comparable with those of bulk processes. While these parameters showed that the enzymatic process is more sustainable than the petrochemical one, an absolute assessment of the process sustainability must take into account the energy required for the



In this study, midpoint indicator assessment was conducted using ReCiPe 2016 Endpoint (E) V1.08 to better understand and compare the impact



categories.³⁶ The LCA of petrochemical esterification was conducted based on two stages in the inventory analysis: the synthetic process and the purification (removal of unreacted reagents and by-products) process. The waste treatment processes and emissions to air, water and land were not considered as part of the LCA.

Relative results were generated from the simulation for which the indicator results maximum is set to 100% (Fig. 5 and Table S3†). The petrochemical synthetic process is above the purification process in all the categories. When compared to the enzymatic process, the petrochemical synthesis is still above the purification and the enzymatic synthesis in all categories.

The petrochemical esterification process had the highest environmental impact across all criteria, mainly due to its long reaction time, high solvent use, and toxic reagents like EDC. The workup stage also contributed significantly, primarily through waste generation. In contrast, the enzymatic process showed the lowest impact, benefiting from a biobased catalyst, lower energy use, and more concentrated solutions that reduced solvent removal energy demands. Water consumption in the petrochemical process was 0.099 m³ per unit of product, while the continuous flow method used only 0.002 m³, reducing water usage by over 90% thanks to more efficient reagent use and simpler purification. Land system changes were also significantly lower in the continuous flow process (0.002 m² vs. 0.061 m² per unit), mainly due to reduced solvent and reagent requirements. Carbon emissions were over 95% lower in the enzymatic process compared to the petrochemical route, highlighting the environmental benefits of using a syringe pump instead of a stirring plate. Across all nine planetary boundaries, including ocean acidification, biosphere integrity, and freshwater use, the enzymatic approach showed the least impact. The petrochemical method had the highest toxicity, largely due to chlorinated reagents and solvents.

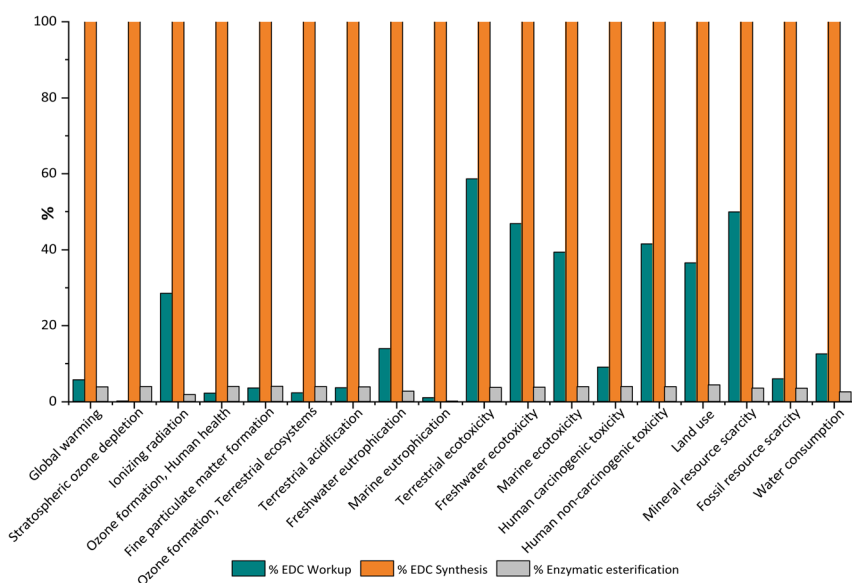


Fig. 5 Environmental impacts due to the synthesis of 100 g of ethyl ester computed using ReCiPe 2016 V1.08.



Lastly, ReCiPe 2016 Endpoint (E) V1.08 was used to conduct endpoint analysis. Human health, ecosystem quality and resources endpoint indicator assessments have been compared for the petrochemical and enzymatic processes (Fig. 6). Ecosystem quality comprises acidification, ecotoxicity, eutrophication and land use (indicated as species $\times$ year ["Species*yr"]). Human health is related to the impacts of environmental degradation that increases the number and duration of loss-of-life-related diseases (recorded as disability-adjusted life year (DALY)). For resources, it is closely related to the depletion rate of raw materials and energy sources (recorded as extra costs in US dollars for future mineral and fossil resource extraction (USD)). Results are evaluated based on the future energy surplus requirements needed to produce lower-quality energy and minerals. For all the categories, the enzymatic esterification showed a lower impact than the petrochemical process (Fig. 6b). By combining the precise midpoint results with the aggregated endpoint outcomes, we provided a robust and comprehensive assessment of the environmental performance of the seven processes, capturing both detailed category-specific impacts and their broader environmental effects. In summary, the implementation of the enzymatic esterification under continuous flow conditions resulted in the reduction of more than 90% based on the average outcomes observed in the respective categories of the endpoint method.

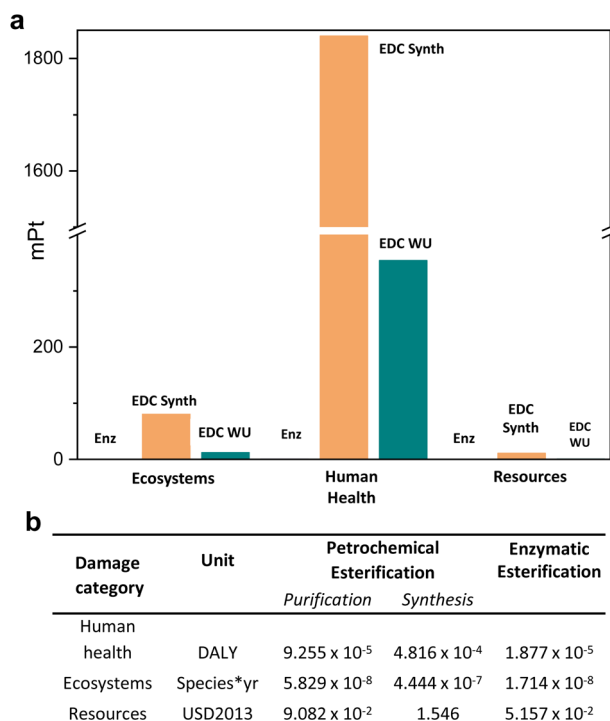


Fig. 6 (a) ReCiPe 2016 Endpoint (E) V1.08 endpoint indicators for producing 100 g of ethyl ester. 'Enz' stands for enzymatic synthesis, 'EDC synth' stands for petrochemical synthesis, and 'EDC WU' represents the workup step for the petrochemical esterification. (b) Environmental impacts due to the stearic acid esterification process (100 g) computed using ReCiPe 2016 Endpoint (E) V1.08.



Data availability

The data supporting this article have been included as part of the ESI.†

Author contributions

A. B. and A. P. D. conceived the work. All authors designed the experiments. N. N. M., D. G., M. R. E. and A. B. performed and analysed the experiments. N. M. aided in the LCA. A. B. directed the research and prepared the manuscript. All authors contributed to manuscript revisions and have given approval to the final version of the manuscript.

Conflicts of interest

There are no conflicts to declare.

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