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H₂-driven biocatalytic O-demethylation of lignin derived aromatics in a closed-loop flow system powered by water electrolysis

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Lignin is an abundant and renewable source of aromatic compounds, yet its utilization remains limited due to its recalcitrance and heterogeneity. Recent developments have enabled the catalytic fractionation of lignin into low molecular weight aromatics, which may be transformed into higher-value compounds. Here, we present a scalable biocatalytic platform for the selective O-demethylation of lignin-derived aromatic compounds, which integrates an O₂-tolerant, NAD⁺-reducing soluble hydrogenase from *Cupriavidus necator* for cofactor regeneration and NADH-dependent cytochromes P450 and Rieske-type monooxygenases. The process was implemented in a closed-loop flow system featuring dialysis membrane-entrapped multi-enzyme modules. H₂ and O₂ were precisely supplied via gas addition modules powered by water electrolysis. This configuration achieved >99% substrate conversion, high atom efficiency, and effective real-time management of the inhibitory byproduct formaldehyde. The hydrogenase-based cofactor regeneration system exhibits robust tolerance to formaldehyde and is adaptable to a broad range of gas-dependent biocatalytic processes, thereby advancing green, resource-efficient chemical production from renewable biomass.

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1. A sustainable O-demethylation system was developed for lignin valorization by leveraging three oxidoreductase systems. By coupling H₂-driven cofactor regeneration in a flow setup, this work demonstrates that lignin valorization can be driven by renewable electricity, with minimal waste and high selectivity.
2. The described process utilizes enzymatic catalysis to facilitate the O-demethylation of LDACs from renewable lignin biomass under mild reaction conditions. It achieves high product yields (>99% substrate conversion), and incorporates water electrolysis to enable an electro-driven process.
3. Future work could focus on reutilizing formaldehyde (a byproduct), broadening substrate scope, and integrating further gas-dependent oxidoreductases.

Introduction

Lignin is a highly abundant biopolymer that constitutes up to 40% of lignocellulosic biomass.^{1,2} As the demand for sustainable alternatives to fossil fuels grows, the carbon-rich aromatic polymer has been increasingly regarded as a resource for platform chemicals.^{1,3–7} Despite its potential, lignin remains

largely underutilized in industrial applications due to its structural complexity, heterogeneity, and recalcitrance.^{8,9} Given the lack of cost-effective methods to transform lignin into high-value products, the majority is burned for heat and power.² The fractionation of lignin biomass yields heterogeneous mixtures of lignin-derived aromatic compounds (LDACs), which are particularly promising substrates for biocatalytic processes.¹⁰ Techniques such as reductive catalytic fractionation (RCF) efficiently break down lignin into LDACs such as alkylguaiacols, *para*-methoxybenzoic acids, and vanillic acid.^{11–14} As lignin is highly methoxylated, the majority of LDACs contain at least one methoxy group. LDACs can be transformed into valuable targets by chemical methods including hydrodeoxygenation, oxidative cleavage, and O-demethylation.^{15,16} However, these methods typically require harsh conditions, often involving high temperatures and pressures and the use of corrosive reagents.^{17–22} For example, O-demethylation via

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acid hydrolysis typically requires high temperatures (120–180 °C), pressurized systems, and strongly acidic conditions (pH < 1). This results in substantial energy input and the formation of corrosive acidic waste, complicating downstream neutralization and purification steps.¹⁷ These challenges result in low selectivity, increased operational costs, and limited scalability. An alternative strategy for *O*-demethylation is biocatalysis, whereby enzymes are used under mild conditions to achieve high selectivity (Fig. 1).^{23–27} Enzymes that catalyze the *O*-demethylation of LDACs include cytochromes P450 (P450s) and Rieske-type oxygenases (ROs).^{25–33} Both classes of oxygenases are widely distributed in bacteria, enabling growth on a variety of LDACs. For example, *Rhodococcus jostii* RHA1 (RHA1) harbours AgcAB and PbdAB, two-component P450 systems that *O*-demethylate alkylguaiaicols (**1a**) and *p*-methoxybenzoates (**2a**),^{24,27} respectively (Fig. 2A). In these systems, AgcB and PbdB are reductases that contain a [2Fe-2S] ferredoxin and a flavin-binding domain, while AgcA and PbdA are heme-containing monooxygenases. Similarly, RHA1 and *Pseudomonas putida* KT2440 (KT2440) utilize VanAB, a two-component RO that catalyzes the *O*-demethylation of vanillate (**3a**).^{34,35} In analogy to the P450 systems, VanA and VanB are oxygenase and reductase components, respectively.

A potential complication of biocatalytic *O*-demethylation is the formaldehyde that is produced. More specifically, *O*-demethylation by P450s and ROs generates one equivalent of formaldehyde (Fig. 2A).²³ Formaldehyde is an electrophilic and protein-reactive molecule that inactivates enzymes through covalent modification of nucleophilic residues such as cysteines, lysines and histidines.³⁶ The inactivation of the enzymes reduces the efficiency and operational stability of the biocatalytic process. Ultimately, this can hinder conversion efficiency and poses scalability challenges.³⁷ Enzymatic *O*-demethylation reactions typically rely on electron transfer from reduced cofactors like NAD(P)H.³⁸ These cofactors are required in stoichiometric amounts to sustain enzymatic activity and can significantly increase operational costs.^{38,39}

Enzymatic cofactor regeneration systems such as formate dehydrogenase (FDH) and glucose dehydrogenase (GDH) have been utilized to mitigate the high costs of cofactor supply, but recycling requires auxiliary organic substrates and generate byproducts.⁴⁰ In contrast, an approach which utilizes “soluble” NAD⁺-reducing hydrogenase (SH), is atom-efficient and produces no unwanted byproducts.^{41–45} This enzyme reduces NAD⁺ to NADH using molecular hydrogen (H₂).^{46,47} More specifically, SH oxidizes H₂ at the Ni-Fe center and transfers electrons through a series of Fe-S clusters to an FMN-binding domain, where NAD⁺ is reduced.⁴⁸ Unlike most hydrogenases, which are deactivated by trace amounts of molecular oxygen (O₂), SH from *Cupriavidus necator* is highly O₂-tolerant and has thus been incorporated into many biocatalytic systems for cofactor regeneration.^{48,49} Indeed, SH an ideal partner for O₂-dependent enzymes such as P450s and ROs (Fig. 1). The integration of SH-mediated cofactor regeneration with the electrochemical production of H₂ and O₂ further facilitates electro-driven biocatalysis. Thus, water electrolysis generates H₂ and O₂, which are required for SH-coupled and O₂-dependent enzymatic processes, respectively. Although H₂ and O₂ can be produced *in situ* using a two-electrode system, this method showed low faradaic efficiency, required highly acidic conditions, and exhibited low liquid-to-liquid gas transfer efficiency.⁴¹ *Ex situ* water electrolysis using a proton exchange membrane (PEM) electrolyzer is more energy-efficient, producing H₂ and O₂ with >60% voltage efficiency and >95% faradaic efficiency. A practical and safe strategy for introducing electrolytically generated gases into biocatalytic flow systems involves the integration of gas-permeable tubing into a closed-loop setup. This configuration enables continuous and controlled delivery of H₂ for cofactor regeneration while preventing enzyme deactivation.^{42,50} Another strategy for whole-cell catalysis is to use the culture broth directly as catholyte during water splitting, providing H₂ to the cells without gas-phase transfer. This has been demonstrated in zero-gap electrolysis systems, for example with *Cupriavidus necator* producing isopropanol from CO₂, in hybrid setups coupling a zero-gap cell

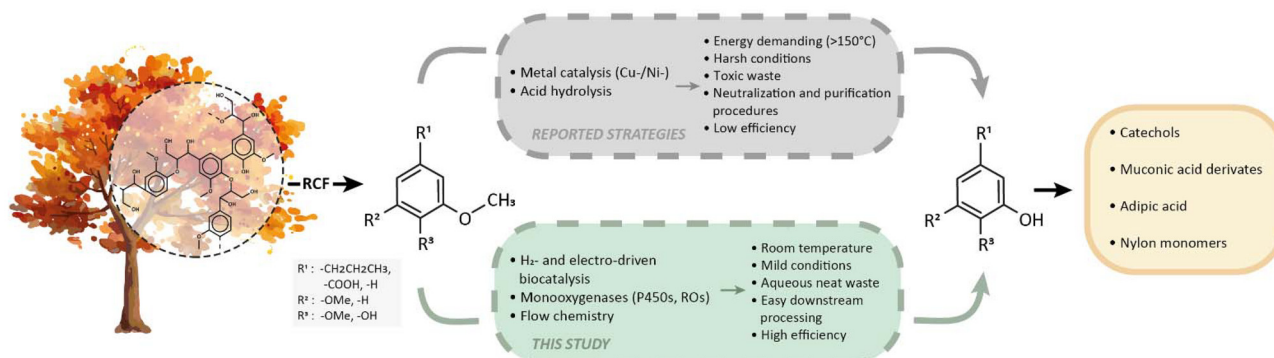


Fig. 1 Schematic representation of the lignin valorization pathway. In grey, the typical LDACs (lignin-derived aromatic compounds) valorization strategies reported in the literature, based on metal catalysis or acid hydrolysis. In green, our proposed biocatalytic approach for LDACs valorization using H₂- and electro-driven systems under mild conditions. R¹, R², and R³ denote typical, yet not exhaustive, substituents found in LDACs. RCF: reductive catalytic fractionation.



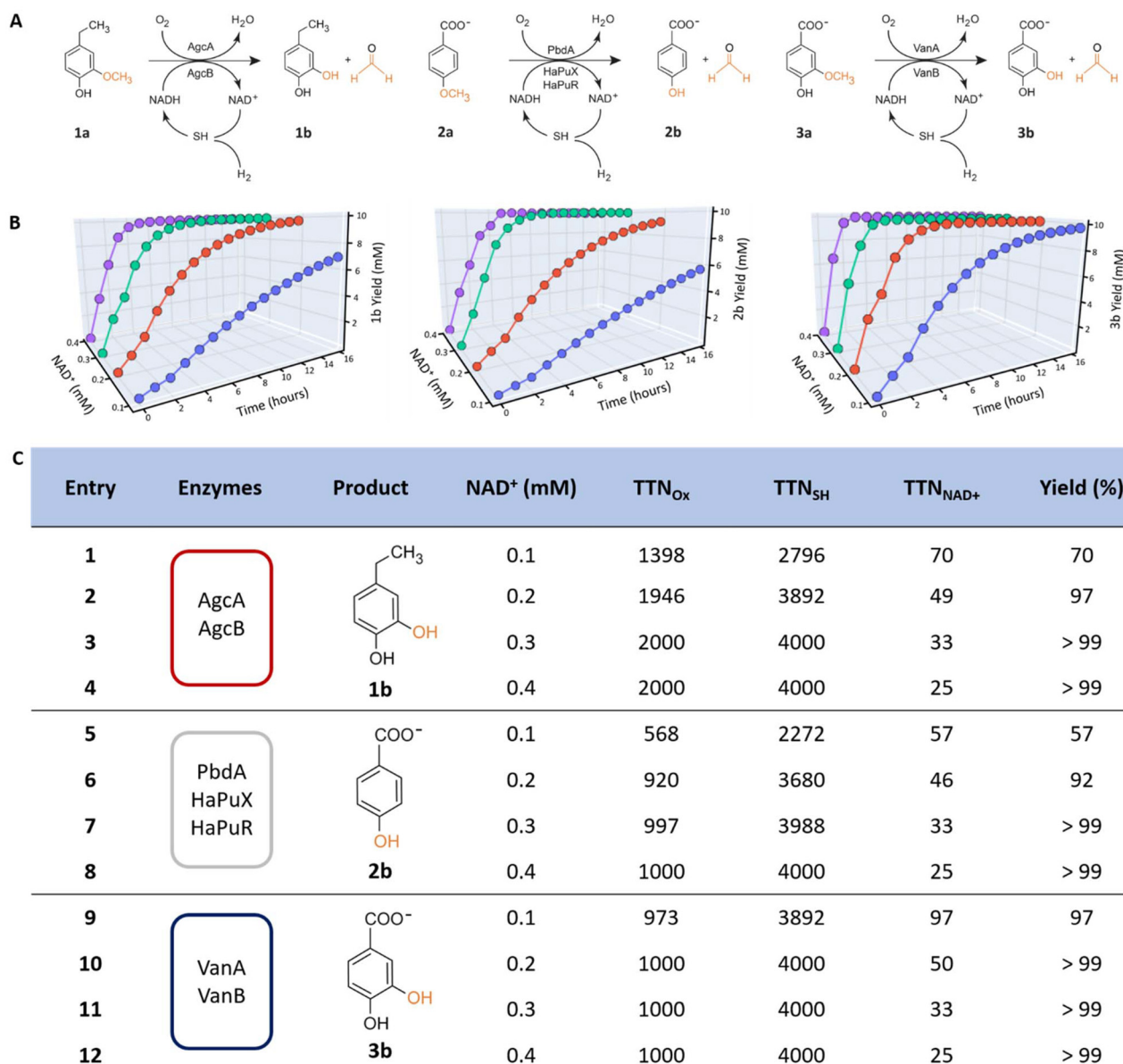


Fig. 2 Comparative analysis of H₂-driven O-demethylation reactions. (A) Reaction schemes of the three studied monooxygenase systems. Batch reactions were carried out in 8 mL gas-tight vials, each containing 1 mL of reaction mixture prepared in 20 mM phosphate buffer (pH 8.0) with 0.1–0.4 mM NAD⁺, 10 mM substrate, and enzyme concentrations optimized for each system (5 μM for AgcA and AgcB, 10 μM for VanA and VanB, 10 μM for PbdA, 5 μM for HaPuX, 1 μM for HaPuR, and 2.5 μM for SH). **1a** was used as the substrate for AgcA/AgcB, **2a** for PbdA/HaPuX/HaPuR, and **3a** for VanA/VanB. Gas mixtures: 4% H₂, 20% O₂, and 76% N₂. All reactions were performed using H₂ as the sole electron source. (B) Time-course profiles of product formation over 16 hours at different NAD⁺ concentrations (0.1–0.4 mM). From left to right, the 3D plots represent AgcA/AgcB, PbdA/HaPuX/HaPuR, and VanA/VanB, respectively. (C) Summary table of reaction performance across the tested NAD⁺ concentrations. The table includes the final product yield (mM), total turnover numbers for the oxygenases (TTN_{Ox}: $n_{\text{product}}/n_{\text{oxidoreductase}}$), soluble hydrogenase (TTN_{SH}: $n_{\text{product}}/n_{\text{SH}}$), and NAD⁺ (TTN_{NAD⁺}: $n_{\text{product}}/n_{\text{NAD}^+}$).

to methanogens using wastewater-derived catholyte, and in water-splitting biosynthetic platforms where electrochemically generated H₂ drive CO₂ conversion with high efficiency.^{51–53}

Here, we report a scalable biocatalytic system for the electro-driven valorization of LDACs by integrating O-demethylation with H₂-driven NADH regeneration using SH. The valorization potential of three LDAC O-demethylase systems, AgcAB, PbdA/HaPuXR, and VanAB, was assessed. To

address formaldehyde accumulation and process scalability, the reactions were adapted to a closed-loop flow setup with continuous formaldehyde removal *via in situ* evaporation. The setup incorporates immobilized and entrapped enzymes, enhancing enzyme stability, reusability, simplifying downstream processing, and improving control over formaldehyde production.^{53–56} The system was optimized to operate safely and efficiently under non-flammable H₂ concentrations.



Tubing with different gas permeability was integrated into the flow system to precisely regulate the release of H₂ and O₂ to the reactor. This study showcases the potential for lignin valorization as well as the future broader application of these systems for NAD(P)H- and FMN₂-dependent oxidoreductase reactions,⁵⁷ offering a scalable and sustainable pathway for biocatalytic chemical production.

Experimental section

Preparation of P450s and ROs

O-Demethylase components were produced as His-tagged proteins in *Escherichia coli* BL21 (DE3). AgcA and AgcB from *Rhodococcus rhodochrous* EP4 (EP4) were produced using pET15a.²⁷ VanA and VanB from KT2400, PbdA from RHA1, and HaPuX and HaPuR from *Rhodospseudomonas palustris* were produced using pET28a.^{24,34} Expression was induced in cultures at an OD₆₀₀ of 0.6–0.7 using IPTG. Cell pellets were harvested by centrifugation and stored at –80 °C until use. Proteins were purified using Ni-NTA affinity chromatography (Thermo Scientific), and protein purity was verified by SDS-PAGE. Protein concentrations were determined using BCA Protein Quantification Assay (SERVA). Detailed purification protocols, including buffer compositions and reconstitution procedures, are provided in the SI.

Preparation of SH

SH from *C. necator* was homologously produced using cells harboring the pGE771 plasmid.⁴⁸ Cultures were grown heterotrophically in a mineral salt medium supplemented with fructose, glycerol, trace elements, and metal ions. Cells were harvested by centrifugation and lysed anaerobically using a French press. SH was purified using a Strep-Tactin XT 4Flow column (IBA), and protein purity and concentration were verified by SDS-PAGE and the BCA Protein Quantification Assay (SERVA). Detailed protocols, including media composition, buffer formulations, and purification procedures, are provided in the SI.

Gas composition reactions setup and analysis

The batch reactions were conducted in 8 mL gas-tight vials, each containing 1 mL of a reaction mixture composed of 20 mM HEPES, pH 8.0, 1 mM NAD⁺, and 10 mM substrate. Enzyme concentrations were 5 μM AgcA, 5 μM AgcB, 10 μM VanA, 10 μM VanB, 10 μM PbdA, 5 μM HaPuX, 1 μM HaPuR, and 2.5 μM SH. The gas headspace composition was controlled using a gas mixing station (SI section S3.2). The O₂ concentration was kept constant at 20%, while H₂ concentrations were varied from 2% to 20% in increments of 2%, with nitrogen comprising the remaining balance. The vials were bubbled with the gas mixture before sealing with gas-tight septa. Reactions were initiated by adding enzymes *via* a gas-tight syringe, followed by incubation at 30 °C with shaking at 200 rpm. After overnight incubation, 50 μL samples were quenched 1:1 with acetonitrile and analyzed by HPLC. Product for-

mation was quantified using UV absorbance by comparing peak areas to standard compounds. Detailed HPLC protocols, including column specifications, gradient settings, and wavelength monitoring, are provided in the SI.

Batch reactions setup and analysis

Time-course experiments were performed in 8 mL gas-tight vials containing 20 mM HEPES, pH 8.0, 0.1–0.4 mM NAD⁺, and 10 mM substrate (ethylguaiacol, **1a**, for AgcAB, vanillate, **2a**, for VanAB, and methoxybenzoate, **3a**, for PbdA/HaPuXR) with a final volume of 1 mL. Enzyme concentrations were 5 μM AgcA, 5 μM AgcB, 10 μM VanA, 10 μM VanB, 10 μM PbdA, 5 μM HaPuX, 1 μM HaPuR, and 2.5 μM SH. The vials were bubbled with a gas mixture containing 4% H₂, 20% O₂, and 76% N₂, sealed with gas-tight septa, and reactions were initiated by adding the enzymes *via* a gas-tight Hamilton syringe. Reactions were incubated at 30 °C with shaking at 200 rpm, and samples were collected hourly for 16 h. Each sample was quenched 1:1 with acetonitrile and stored on ice before HPLC analysis as previously described. Formaldehyde formation was monitored over time *via* derivatization with 2,4-dinitrophenylhydrazine (DNPH), followed by GC-FID analysis. Detailed methods are provided in the SI.

Formaldehyde resistance analysis

The formaldehyde tolerance of the enzyme systems was assessed by conducting NAD⁺ reduction (for SH) and NADH oxidation (for AgcA/AgcB, PbdA/HaPuX/HaPuR and VanA/VanB) assays in 2 mL quartz cuvettes, using a Cary 60 UV-Vis spectrophotometer. For comparison, a commercially available FDH from *Candida boidinii* (Merck) was used as an alternative NAD⁺ reducing system. Reactions were maintained at 30 °C, with the following enzyme concentrations: 2.5 μM for SH, 5 μM for AgcA/AgcB, 10 μM for VanA/VanB, 10 μM for PbdA, 5 μM for HaPuX, and 1 μM for HaPuR. Formaldehyde concentrations were varied between 2 and 20 mM, with 0.4 mM NAD⁺ used in SH reactions and 0.4 mM NADH used in oxidoreductase assays. Reactions were initiated by adding the enzymes directly into the cuvette. Progress was monitored by measuring absorbance at 365 nm, using an extinction coefficient of 3.3 mM⁻¹ cm⁻¹ for NADH/NAD⁺ quantification. Reaction slopes were recorded, and enzyme activities were normalized relative to activity in the absence of formaldehyde.

Electro-driven biotransformation with immobilized enzymes

The reaction was carried out in a closed-loop flow system scaled to 200 mL volume using a peristaltic pump. H₂ and O₂ were produced using a lab-scale PEM electrolyzer (H-TEC education) operated at 3.9 V and 2.21 A (H₂ 33 mL min⁻¹, O₂ 16 mL min⁻¹) for 5 min before addition of aromatic substrate. Approximately 3.8 h (228 min) after substrate addition, the PEM electrolyzer was operated at 3.5 V and 1.03 A (H₂ 21 mL min⁻¹, O₂ 10 mL min⁻¹) to ensure full conversion of substrate. H₂ and O₂ were directly transferred into the flow volume *via* gas addition modules using polyvinylmethyl siloxane (PVMS) and polytetrafluoroethylene (PTFE) tubing, respectively. The



different permeabilities and lengths of these tubings, described in the SI, enable precise and controlled gas delivery. The reaction medium, containing 50 mM vanillate and 0.3 mM NAD⁺ in 20 mM Tris-HCl, pH 8.0 at 30 °C, was pumped through the system at 3 mL min⁻¹. A flow Clark-type H₂ sensor (Unisense) and an optical O₂ sensor (PreSens) were added to the flow setup to allow online monitoring of dissolved gases to maintain safe H₂/O₂ concentrations. SH and catalase (TCI chemicals) were immobilized on Amberlite FPA54 resin through non-covalent adsorption, at 1 mg SH and 5000 U catalase per gram of resin. This resin was packed into a C10/10 column (Cytiva) as a cofactor regeneration unit with a bed volume of 4 mL. VanAB were contained in a semi-permeable 5 kDa dialysis membrane (Carl Roth) and immersed in a 250 mL Schott bottle with magnet inside for increased diffusion as a biotransformation unit. The temperature of the biotransformation unit was maintained at 30 °C with a hot plate. The cap of the biotransformation unit was opened during the first 2 hours of the reaction for evaporation of formaldehyde. Thin-layer chromatography (TLC) and high-performance liquid chromatography (HPLC) were used to monitor reaction progress, and formaldehyde levels were determined by gas chromatography. At the end of the reaction, the product was extracted with ethyl ether, dried, and crystallized for further analysis by NMR. Details are provided in the SI.

Results and discussion

Benchmarking safe and efficient H₂-driven *O*-demethylation

The enzymatic transformation of LDACs offers a promising strategy for converting renewable biomass into value-added chemicals. We selected three representative *O*-demethylases to further explore the biocatalytic potential of these enzymes: AgcAB from EP4, VanAB from KT2440, and PbdA from RHA1 paired with HaPuX and HaPuR from *R. palustris*. HaPuX and HaPuR, the ferredoxin and ferredoxin reductase, respectively, from a homologous P450 system, were used instead of PbdB due to the O₂-lability of the cognate reductase (SI Table S3).²⁴ Similarly, AgcAB from EP4 was selected due to the greater stability of the reductase *versus* AgcB from RHA1. AgcAB, VanAB, and PbdA/HaPuXR catalyze the *O*-demethylation of ethylguaiacol (**1a**), *p*-methoxybenzoate (**2a**), and vanillate (**3a**), to ethylcatechol (**1b**), hydroxybenzoate (**2b**), and protocatechuate (**3b**), respectively (Fig. 2A). To overcome the constraints of supplying stoichiometric NADH, we explored the use of SH from *C. necator* to efficiently regenerate NAD⁺ using H₂ as the reductant.

We first investigated the effect of gas composition on reaction performance. Specifically, we assessed the minimum H₂ concentration required to sustain full turnover under constant O₂ (20%) and whether efficient operation could be achieved under non-explosive conditions. In 1 mL reactions performed in 8 mL gas tight vials, all three enzyme systems, AgcAB for the conversion of **1a** to **1b**, PbdA/HaPuXR for **2a** to **2b**, and VanAB for **3a** to **3b**, reached >99% substrate conversion when oper-

ated at H₂ concentrations of 4% or higher (SI Fig. S4). This corresponds to 13.6 μmol of H₂, slightly exceeding the 10 μmol of aromatic substrate in solution. At 2% H₂ (~6.8 μmol), conversions dropped sharply to ~60% across the three systems, consistent with a stoichiometric relationship between H₂ and product formation. This highlights the role of H₂ as a stoichiometric reagent under batch conditions. These results also establish that high enzymatic efficiency can be retained even at low H₂ partial pressures, consistent with SH's low *K_M* value for H₂ (<40 μM) and highlighting the system's intrinsic safety and scalability.⁵⁸ Compared to previously reported H₂-driven biocatalytic processes, such as processes involving imine reductases (IREDs) and putrescine oxidases under 50:50 H₂:O₂ ratios, IREDs at 100% H₂, or flavin-dependent oxidoreductases operating with 10% O₂ and 60% H₂, our system achieves complete substrate conversion using just 4% H₂ under atmospheric pressure conditions.^{43,44,59} This positions our platform as a significant advancement in the safe operation of H₂-driven biocatalytic systems.

Moreover, the ~90% match between expected and observed conversion efficiencies under optimized conditions indicates that electron transfer from H₂ to the final product is relatively well-coupled. Each mole of H₂ donates two electrons, which are transferred *via* SH to NAD⁺ and ultimately to the oxygenase through the reductase. Electrons may be lost to O₂ during this process, from either the reductase or the oxygenase, generating ROS. The two-component systems VanAB and AgcAB are well-coupled under certain conditions, as demonstrated previously for VanAB³⁴ and for AgcAB in unpublished data by Wolf *et al.* As the coupling of PbdA to HaPuXR was not known, we tested coupling with varying proportions of the three components. In these studies, a 10:5:1 ratio of PbdA:HaPuX:HaPuR maximized activity and coupling, and was used in subsequent experiments (SI Table S3). Overall, the relatively tight coupling from H₂ to final product reflects minimal loss of electrons throughout the electron transport chain.

We next examined the influence of NAD⁺ concentration on each system's performance. Reactions were conducted with NAD⁺ ranging from 0.1 to 0.4 mM, while maintaining 10 mM substrate and 4% H₂. As summarized in Fig. 2C, VanAB (**3a** to **3b**) demonstrated the most robust profile, with full conversion being achieved using 0.2 mM NAD⁺. Reaction Mass Efficiency (RME) values remained consistently around 21%, indicating that the system performs well even under cofactor-limited conditions, with no gain in atom efficiency above 0.2 mM NAD⁺ (SI Table S4). VanAB's high RME at lower NAD⁺ concentration is comparable to those of conventional chemical processes for lignin valorization but is achieved under significantly milder conditions, enabling efficient substrate conversion with minimal waste.⁶⁰ AgcAB (**1a** to **1b**) required higher cofactor concentrations for stoichiometric conversion ([NAD⁺] = 0.3 mM). Finally, the three-component system PbdA/HaPuXR (**2a** to **2b**) showed lower relative performance. At 0.1 mM NAD⁺, TTN_{ox} reached only 568, yielding 5.7 mM product with an RME of 15.9%. The reductase HaPuR was supplied at a 10-fold lower concentration than PbdA to maintain high coupling (SI



Table S3) and likely restricted the comparative efficiency of this system.

Interestingly, AgcB and VanB have low K_M values (12 and 10 μM , respectively) and high k_{cat}/K_M values (1.3 and 3.0 $\mu\text{M}^{-1}\text{s}^{-1}$, respectively) for NADH.^{27,61} These high efficiencies support fast turnover at relatively low cofactor concentrations, consistent with the high TTN and RME values. Conversely, the K_M of SH for NAD⁺ is approximately 0.5 mM, which is slightly above the maximum tested concentration (0.4 mM). Under these conditions, NAD⁺ concentration is not saturated for SH, resulting in sub-maximum reaction rates. Finally, RME varies not only with cofactor input, but also with the interplay between enzyme efficiency, time, and turnover balance. Identifying the lowest NAD⁺ concentration at which complete conversion occurs without compromising time or yield is critical for scaling. Higher concentrations of NAD⁺ may increase the reaction rate, but lower RME and cost efficiency.

In this study, we tested each of the enzyme systems for activity on their preferred monoaromatic substrate, all of which are LDACs. However, lignin processing frequently produces higher order lignin-derived oligomers.⁶² Although activity of AgcA, PbdA and VanA on lignin-derived oligomers has not been explored experimentally, it is unlikely to be significant given the enzymes' respective substrate specificities. We further note that emergent lignin valorization strategies seek to increase monomeric yields, a priority that aligns with a set-up involving monomer-active enzymes.^{14,63}

By combining TTN, RME, and product yield, our data highlight how careful optimization of cofactor input across different catalytic architectures enables the development of H₂-driven systems that are not only effective, but also sustainable.

Formaldehyde tolerance of the biocatalytic components

The enzymatic *O*-demethylation of LDACs results in the production of formaldehyde (Fig. 2A).^{64,65} In light of formaldehyde's ability to inactivate enzymes,^{37,66} we assessed the formaldehyde tolerance of the different components of our H₂-driven systems. More specifically, we evaluated the relative activities of AgcAB, PbdA/HaPuXR, and VanAB across 0–40 mM formaldehyde (Fig. 3). At the lowest formaldehyde concentration (2.5 mM), all three systems retained over 90% of their initial activity, indicating a tolerance to low concentrations of small aldehydes. However, with increasing formaldehyde concentrations, significant differences in activity attenuation were observed among the systems. VanAB exhibited the highest tolerance, retaining approximately 90% of its initial activity at 5 mM formaldehyde, and maintained around 40% activity at 20 mM. In contrast, AgcAB and PbdA/HaPuXR were more sensitive, with both systems exhibiting ~70% and ~40% activity at 5 and 10 mM formaldehyde, respectively.

We also assessed the activities of two NADH regeneration enzymes, SH and the commercial *C. boidinii* metal-independent FDH (Fig. 3B). SH exhibited remarkable tolerance, maintaining over 85% activity at 5 mM formaldehyde and retaining over 50% activity at 20 mM. In comparison, the activity of FDH

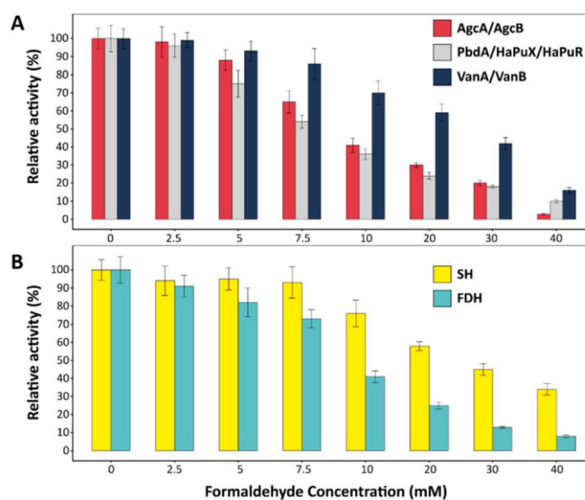


Fig. 3 Differential formaldehyde sensitivity of oxygenases and NAD⁺ reducing modules. (A) The figure illustrates the formaldehyde tolerance of the three oxidoreductase systems AgcA/AgcB, PbdA/HaPuX/HaPuR, and VanA/VanB, expressed as the relative NADH oxidation activity (%) at increasing formaldehyde concentrations (0–40 mM). Reactions were performed in 2 mL quartz cuvettes, enzyme concentrations were 5 μM for AgcA/AgcB, 10 μM for VanA/VanB, 10 μM for PbdA, 5 μM for HaPuX, and 1 μM for HaPuR, with 0.4 mM NADH as the electron donor. (B) Relative activity of the NADH regeneration systems SH (soluble [NiFe]-hydrogenase) and FDH (formate dehydrogenase) under identical conditions. Both systems catalyze the reduction of NAD⁺ to NADH, with SH using H₂ as electron donor and FDH using formate. Enzyme concentrations were 2.5 μM for SH and 3 U mL⁻¹ for FDH, with 0.4 mM NAD⁺. T-Tests revealed that SH maintained significantly higher activity compared to FDH at formaldehyde concentrations ≥ 5 mM ($p < 0.05$), with pronounced significance at ≥ 20 mM ($p < 0.005$).

sharply declined, with more than 50% inhibition observed at 10 mM formaldehyde, and less than 20% activity at concentrations above 20 mM.

Without further study, it is unclear why VanAB is more formaldehyde-tolerant than AgcAB and PbdA/HaPuXR, or whether this property reflects ROs and P450s more broadly. Regarding the cofactor regeneration systems, the remarkable tolerance of SH may be due to its FeS cluster-based mechanism for electron transfer from the NiFe site, which minimizes susceptibility to electrophilic modifications. In contrast, FDH catalyzes direct hydride transfer from formate to NAD⁺ and relies on cysteine residues for activity. These thiol groups are highly reactive toward electrophiles such as formaldehyde, making FDH more susceptible to formaldehyde-induced inactivation.⁶⁷ This observed difference in formaldehyde tolerance is consistent with previous comparative studies on [FeFe]- and [NiFe]-hydrogenases, which provide valuable insights into how redox-active metal centers interact with formaldehydes.⁶⁸ Furthermore, the [NiFe]-active site, in contrast to metal-independent FDHs, is deeply buried within the enzyme, with access controlled by a gas tunnel that limits formaldehyde entry.⁶⁹ This structural feature provides an additional protective mechanism, preventing small reactive species like formaldehyde from directly modifying the metal center.⁶⁸ This struc-



tural advantage makes the SH more suitable for applications involving aldehyde-generating biotransformations. When compared to other NADH regeneration systems such as FDH, SH offers a unique advantage in formaldehyde-rich environments while also minimizing byproduct formation. Despite the retained activity of both the oxidoreductase systems and SH, formaldehyde accumulation during lignin valorization remains a critical challenge in setups that involve a higher substrate load.

Electro-driven *O*-demethylation with closed-loop flow biocatalysis

To enable continuous formaldehyde removal, scale-up, and *in situ* H₂ supply, we designed a closed-loop flow biocatalysis system. Due to its higher formaldehyde tolerance compared to the other two *O*-demethylase systems, VanAB was assessed in the flow system. Our initial attempts to immobilize VanAB using metal ion affinity beads (EziG) resulted in complete loss of activity, likely due to disruption of the direct interaction between the two components that is required for electron transfer. To address this, a closed-loop flow system was developed with a dialysis membrane (MWCO 3.5 kDa) that entrapped VanAB in a 5 mL volume, referred to here as the biotransformation unit.^{70,71} This maintained the two components at relatively high concentrations, maximizing their interaction and therefore activity. SH was immobilized on Amberlite FPA.⁵⁴

an anionic resin, enabling H₂-driven NADH regeneration. To introduce H₂ and O₂ continuously to the bioreactor in the correct proportions, separate gas addition modules were implemented to reduce gas competition due to differential solubility and diffusion rates (Fig. 4A). The H₂ module was positioned in the flow sequence directly upstream of the O₂ module. This arrangement ensured accurate tuning of the gas composition without direct aeration into the reaction volume avoiding inactivation by shear force. Using H₂ and O₂ from water electrolysis, SH-Amberlite facilitated reduction of NAD⁺ for VanAB even at 15% O₂ (Fig. 4C).

The entrapped VanAB in the biotransformation unit exhibited high activity, converting >99% of 50 mM vanillate (**3a**) to protocatechuate (**3b**) within 5 h (Fig. 4B). The biotransformation unit was maintained at 30 °C using a heating panel to enhance VanAB activity. The formation of reactive oxygen species was prevented by adding catalase to the biotransformation unit. The biotransformation unit was open to the atmosphere allowing equilibration with air. This presumably helped keep formaldehyde concentration below 0.5 mM (Fig. 4B). Finally, since the enzymes were immobilized on columns/within the biotransformation unit, the flow volume only contained **3a/3b** and NAD⁺/NADH. This design greatly facilitated a simple product extraction using diethyl ether, where the 200 mL reaction volume yielded 1.542 g (95% isolated yield) of protocatechuate (**3b**).

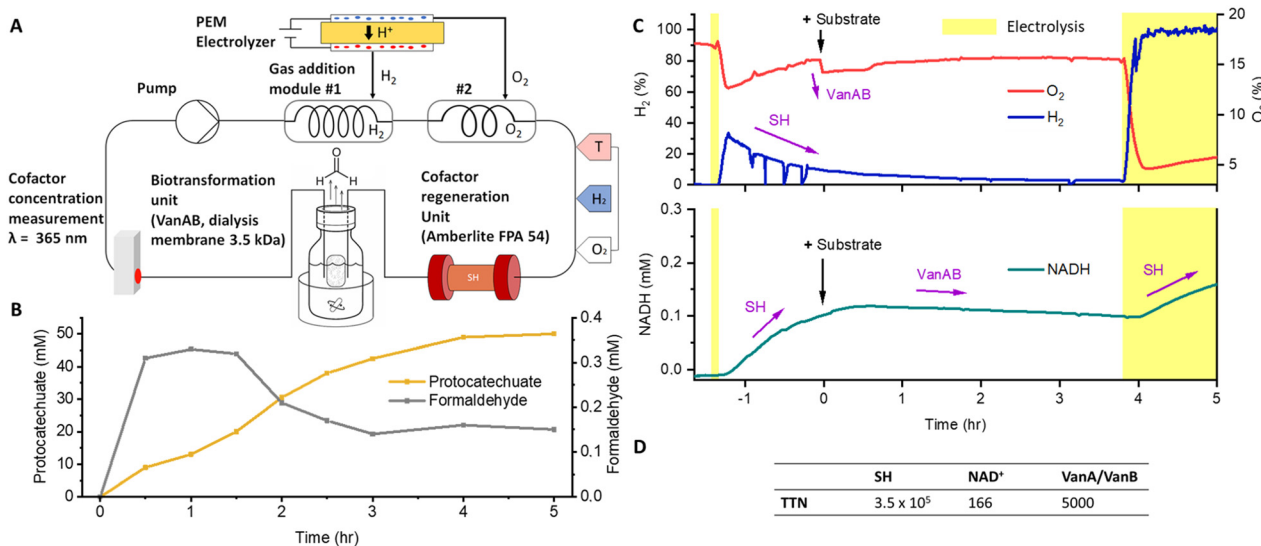


Fig. 4 Integrated flow system for electrodriven biocatalytic *O*-demethylation of vanillate. (A) Flow setup for electro-driven *O*-demethylation of vanillate. H₂ is produced from a PEM electrolyzer using Pt_B/Pt_C as a H₂ evolution catalyst. Two gas addition modules are used to transfer H₂ and O₂ to the 200 mL closed-loop system containing 20 mM Tris-HCl buffer (pH 8.0), 50 mM vanillate (**3a**) and 0.3 mM NAD⁺. H₂, O₂, and temperature sensors together with a spectrophotometer were integrated. 1 mg of SH was immobilized by Amberlite FPA. VanA and VanB were entrapped in a dialysis membrane and integrated as a biotransformation unit. (B) Product protocatechuate yield (dark yellow) and formaldehyde concentration (dark gray) across time points during the upscaled reaction of electro/H₂ driven *O*-demethylation of vanillate. (C) On-line monitoring of H₂, O₂ (up), and NADH (down) during the electro-driven biotransformation. The concentration of H₂ (blue), O₂ (red), and NADH (green) were followed during the time of reaction. The pulsed electrolysis is also depicted (yellow box). NAD⁺ was added before the electrolysis, then substrate was added after NADH reached a plateau. After 3.8 hours substrate addition, H₂ and O₂ were supplied again with a lower rate. Purple arrows indicate the activity of each enzyme. (D) Total turnover numbers of all biocatalysts ($n_{\text{product}}/n_{\text{enzyme}}$) and cofactors ($n_{\text{product}}/n_{\text{NAD}^+}$) involved in the closed-loop flow setup process.



The described flow system can be compared to fed-batch fermentation using VanAB-expressing *P. putida*, where equimolar formaldehyde during the *O*-demethylation reaction limits protocatechuate production to 17.5 g L⁻¹ in 50 mL media.⁷² In contrast, the flow setup enables continuous removal of formaldehyde and offers modular scalability by connecting or expanding reactor units, providing a directly controllable operational format. For gas-dependent transformations, scale-up limitations in gas-to-liquid transfer can be alleviated by extending the gas-permeable tubing within the gas addition module, thereby improving mass transfer. In the flow reaction, the TTN of SH and VanAB reached up to 3.5 × 10⁵ and 5000, respectively, showing high stability of the enzymes (Fig. 4D). The high TTN_{SH} compared to the batch reactions is due to a lower enzyme load and an increased product yield. The TTN_{SH} in our upscaled flow reactions is more than three-fold higher than that of a comparable system for flavin-dependent biocatalysis, where the SH was immobilized on Strep-Tactin XT resin.⁴² This high TTN for SH-Amberlite corroborates previous reports, where Amberlite FPA⁵⁴ served as a stable enzyme carrier for SH reaching TTN values of up to 1.1 × 10⁶ after six cycles.⁴¹ The TTN of entrapped VanAB was three-fold higher than that reported for other RO systems, including a sophisticated hybrid system comprising cumene dioxygenase and the reductase from phthalate dioxygenase (TTN of 1515).⁷³ This illustrates the challenges of the redox interactions in ROs and also demonstrates that they can be effectively stabilized through entrapment for flow applications.

Prior studies applied PVDF (polyvinylidene fluoride) membranes as teabags, entrapping lyophilized whole-cell biocatalysts for stereoselective synthesis.⁷⁰ Similar catalyst containment using dialysis membranes such as PVDF has been used since the 1980s as drug-delivery systems and, more recently, for oxazoline Cu(II) Lewis acid catalysts.^{71,74} Dialysis membranes offer molecular weight cut-off control, ease of handling, and robust structural integrity under various pH/temperature conditions compared to other entrapment methods such as hydrogels.⁷⁵ Also, enzyme entrapment applications serve as an excellent alternative to enzymatic membrane reactors (EMRs), ensuring efficient diffusion, reusability of the enzymes. Finally, entrapment does not require high-pressure systems and maintains enzyme stability. Overall, the high TTN_{VanAB} achieved here establishes that entrapment with dialysis membrane is a viable method for stabilizing multi-component oxidoreductases in flow systems.

To facilitate optimization of the bioreactor, we monitored various parameters online during the reaction, including the electrocatalytic evolution of the gases and their enzymatic consumption (Fig. 4C). Faradaic efficiency was improved by pulsing the PEM electrolysis. After 5 min of electrolysis (3.9 V, 2.21 A), 25% dissolved H₂ and 12% dissolved O₂ were reached in the flow system (Fig. 4C). At 3.8 h after the addition of substrate, the electrolysis was activated again (3.5 V, 1.03 A) for 1.2 h to further increase the reaction rate. A sharp decrease in dissolved O₂ concentration upon substrate addition was

observed, indicating O₂ consumption by VanAB during *O*-demethylation. In the flow setup, the spectrophotometer for monitoring NADH was positioned downstream of the biotransformation unit. As a result, the detected NADH concentration reflected its level post-enzymatic consumption. Four hours after substrate addition, an increase of NADH concentration was observed, indicating the completion of VanAB reaction. This was confirmed by GC (Fig. 4B) and online TLC monitoring. The higher NADH concentration may also reflect increased SH activity due to higher H₂ concentrations from water electrolysis (Fig. 4C).

We also determined the faradaic efficiency for the electrodriven *O*-demethylation of **3a** to **3b** in the flow system. Five hours after substrate addition, with 99% conversion, the faradaic efficiency of the flow system was 18% (SI section S4). However, the electron contribution to product formation was three-fold higher (62%) when measured to 4 h after substrate addition. At that timepoint, only 12 min of electrolysis was performed, despite near-complete vanillate turnover. The lower faradaic efficiency observed at 5 h suggests that during prolonged operation, a significant portion of electrons supplied during electrolysis no longer contributes to product formation. This may reflect mostly complete substrate conversion, although the open biotransformation unit must result in H₂ loss and thus lowered efficiency. The high faradaic efficiency relative to substrate conversion in this system suggests that only 5–10 min of electrolysis is sufficient to drive the reaction in the flow system. The faradaic efficiency of bioelectrocatalytic systems can reach up to 100% when electron transfer from the electrode to the enzymes is directly and optimally controlled.⁷⁶ Nevertheless, the 62% faradaic efficiency achieved with the current design is over 400-fold higher than that reported for another closed-loop biocatalytic flow setup.⁴² While high atom efficiency and faradaic efficiency were demonstrated in this system, optimizing the balance between reaction time with electrolysis duration is critical to further optimizing the process's energy efficiency. We evaluated waste production using the environmental factor (*E*-factor).^{79,80} For the 200 mL reaction performed here, the *E*-factor was 3.6 and the *E*⁺-factor was 2850 (Table 1 and SI Table S6). This indicates that less waste was formed than in conventional NADH regeneration systems used other oxidoreductase reactions.^{77–80} For instance, glucose/GDH-based regeneration produces stoichiometric quantities of gluconolactone or gluconic acid as waste products. In contrast, H₂-driven regeneration using SH generates

Table 1 Environmental and biocatalytic parameters of flow system for vanillate conversion

Parameters	Values
<i>E</i> -Factor	3.6
<i>E</i> ⁺ -Factor	2850
EcoScale	88
Carbon efficiency	87.5%
Atom efficiency	76.2%
RME	90.3%



no organic byproducts. This is reflected in high EcoScale atom and carbon efficiencies (Table 1). Overall, the H₂/electro-driven biocatalytic *O*-demethylation process demonstrates significantly lower waste generation than previously reported electro-biocatalytic approaches. Reuse of the nicotinamide cofactor offers an effective route to increase the operational TTN. NAD (H) can be retained or recaptured during product isolation using established separation strategies such as membrane processes that exploit size and Donnan exclusion.⁸¹ Cofactor immobilization on functionalized polymers or silica supports is another option, though typically accompanied by diffusional limitations, reduced turnover, and higher material costs.⁸² These constraints should be carefully considered when evaluating immobilization for continuous operation. As an outlook, the demethylated carboxylates could be recovered using established aqueous-phase separations, including pH-shift liquid-liquid extraction, capture and release on anion-exchange or polymeric adsorption resins (with optional in-flow product removal).⁸³

Conclusions

This study presents a scalable, energy-efficient biocatalytic system for valorizing LDACs. More specifically, our bioreactor for electro-driven *O*-demethylation integrates water electrolysis with H₂-driven NADH regeneration and a simple entrapment system for multicomponent oxygenases. We optimized the activity of three such oxygenases, AgcAB, PbdA/HaPuXR, and VanAB, with the H₂/SH cofactor regeneration system, demonstrating the feasibility of using these enzymes at low concentrations of H₂ and NAD⁺ while maintaining tolerance to formaldehyde. The closed-loop flow system achieved scalable and optimized electro-driven biocatalytic *O*-demethylation. Using VanAB for proof-of-concept, we established a flow system that can valorize LDACs in high atom efficiency. In future, the system could be further refined to support continuous product release by integration of an *in situ* product extraction module and optimized by scalable gas transfer. The inclusion of gas sensors within the setup also allowed real-time monitoring of the interplay between water electrolysis and biocatalysis, providing valuable insight for future process development. This platform allowed for the implementation of pulsed electrolysis and improved faradaic efficiency, as well as ensuring safe and scalable biotransformation. Importantly, the described platform provides a foundation for broader applications involving NAD(P)H-utilizing multicomponent oxidoreductases systems such as soluble methane monooxygenase as well as other P450s and ROs. Looking ahead, future scale-up should focus on integrating a low-temperature in-line capture unit to condense and retain evaporated formaldehyde, thereby enabling its selective removal and continuous recovery. Such a capture stream creates opportunities for sustainable valorization: recovered formaldehyde can be converted into chiral amines *via* IRED-catalyzed reductive amination,⁸⁴ or transformed into higher carbon products through formolase-mediated C–C

bond formation.⁸⁵ In parallel, further optimization of electrolysis parameters, combined with cofactor immobilization or entrapment, is expected to enhance process robustness and long-term operation. These improvements may also expand compatibility with alternative reaction media, including organic solvents and deep eutectic systems,⁸⁶ thereby offering expanded possibilities for future green biotransformations.

Author contributions

L. L., D. C., and G. L. conceptualized the study; D. C., G. L., P. N., and M. E. W. designed and conducted the experiments; D. C. and G. L. performed the formal analysis; L. L. secured funding for the project; L. L., P. R. F. C., and L. E. supervised the research and provided guidance throughout the study; D. C. and G. L. prepared the original manuscript draft; L. L., P. R. F. C., L. E., D. C., and G. L. reviewed and edited the manuscript; all authors contributed to data analysis and discussions.

Conflicts of interest

There are no conflicts to declare.

Data availability

The data supporting this article have been included as part of the supplementary information (SI). Supplementary information is available. See DOI: <https://doi.org/10.1039/d5gc05054e>.

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