Fusion of Glutamate Dehydrogenase and Formate Dehydrogenase Yields a Bifunctional Efficient Biocatalyst for the Continuous Removal of Ammonia

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A novel fusion protein has been rationally designed, combining the hexameric glutamate dehydrogenase from Clostridium symbiosum with the dimeric formate dehydrogenase from Candida boidinii. The former enzyme consumes ammonia for the reductive amination of α-ketoglutarate using NADH, while the latter biocatalyst regenerates continuously the cofactor. This enzymes fusion opens new perspectives for the detection and the removal of ammonia. The bifunctional biocatalyst has been successfully created, expressed, and then characterized. The two fused protein domains retained identical properties and catalytic activity of the individual enzymes. Additionally, the immobilization on a methacrylate resin optimized the assembly providing a reusable and stable biocatalyst. This is an example of immobilization of a fusion protein, so that efficiency and sustainability of the process are enhanced. The immobilized biocatalyst could be recycled 10 times retaining still half of the initial activity. Such preparation outperforms the co-immobilized wild-type enzymes in the conversion of 300 mM of ammonia, which could be carried out also in continuous mode.

Keywords: fusion protein, protein engineering, ammonia removal, glutamate dehydrogenase, formate dehydrogenase, enzyme immobilization

INTRODUCTION

A fusion protein is a molecule consisting of two or more protein domains incorporated into one single complex. Naturally occurring fusion proteins are widespread, and their modular organization has been identified as an important evolutionary phenomenon (Béguin, 1999; Yu et al., 2015). Artificial fusion proteins may be designed to achieve improved properties or new functionality (Yu et al., 2015). For this purpose, the genetic combination of two proteins to generate a bifunctional enzyme complex has evolved over the years (Lindbladh et al., 1992). The creation of one single protein presenting the biocatalytic activity of two distinct enzymes can dramatically simplify the phases of expression and purification with a significant impact on time, materials and costs spent for the manipulation. This feature translates into sustainable processes that are extremely attractive in line with the growing demand of environmental protection policies.

Furthermore, the very close proximity of the two protein domains yields a higher catalytic efficiency of the two sequential reactions. In fact, the molecule being produced from one enzyme could directly migrate to the active site of the second enzyme without first diffusing to the bulk environment, through substrate channeling (Wheeldon et al., 2016).

Fusion enzyme engineering is one of the promising approaches for the efficient in situ regeneration of cofactors for cost-effective biotransformations (Liu and Wang, 2007). As a
matter of fact, the cofactor is too expensive to be applied in stoichiometric amounts, therefore a recycling system is always incorporated to enable a cofactor-dependent reaction in an economically feasible manner (Aalbers and Fraaije, 2019). Some examples of successful cases reporting the development of fusion systems possessing a protein domain to restore the cofactor have been highlighted in the literature (Prachayasittikul et al., 2006; Hölsch and Weuster-Botz, 2010; Sührer et al., 2014; Fang et al., 2015; Jiang and Fang, 2016; Aalbers and Fraaije, 2017; Zhang et al., 2017; Aalbers and Fraaije, 2019; Hartley et al., 2019). For example, the fusion between a NADPH-accepting mutant of formate dehydrogenase from *Mycobacterium vaccae* (MycFDH) and a ketone reductase (KR) showed beneficial effects in the asymmetric reduction of pentafluoroacetophenone in whole-cell biotransformations (Hölsch and Weuster-Botz, 2010). Later, the fusion of an alcohol dehydrogenase (ADH) with a cyclohexanone monoxygenase (CHMO) has been employed for the synthesis of ε-caprolactone from cyclohexanol, exhibiting a better performance than the combined individual enzymes (Aalbers and Fraaije, 2017). However, only one of the six cloned variants of ADH/CHMO fusion protein maintained the original ADH activity, while the CHMO domain was active in all the cases. This work highlighted how genetic fusion does not always produce fully active enzymes, where structural changes and the orientation of the protein domains play a crucial role in the ability to form the assembled quaternary structure. Indeed, the oligomer interface of one domain may be hindered by the second protein, preventing the correct assembly needed for stability and activity (Aalbers and Fraaije, 2017).

Fraaije and co-workers described the construction of three different ADHs fused with an NADPH oxidase (NOX). Each enzyme retained its catalytic properties and the coupling of the two proteins offered a rapid qualitative screening system for the detection of ADH/NOX oxidase activity to make large mutant libraries (Aalbers and Fraaije, 2019). It is worthy to mention a further example, where enzymes were engineered to both retain and recycle their cofactor (NAD+ and ATP) with the final purpose of producing a drug precursor in a three-step continuous-flow reactor system. In this case, the fusion of three enzymes had little effect on the kinetic performance or thermal stability of each domain. Moreover, the enzyme fusions have been successfully immobilized with 86–98% yield and high recovered activity (Hartley et al., 2019).

Glutamate dehydrogenase (GluDH) from bovine liver is the enzyme of choice for the quantification of ammonia in solution. It catalyzes the reductive amination of α-ketoglutaric acid to glutamic acid using NADPH and ammonia, which is measured by monitoring the decrease in absorbance at 340 nm caused by the oxidation of the cofactor (Kun and Kearney, 1974). This enzyme has been successfully applied for the development of biosensors as well as the generation of the commercially available ammonia assay kits (Azmi et al., 2009). These systems are mainly used for the quantitative determination of ammonia in food, water and biological samples, such as blood and urine (Seshadri et al., 2006; Azmi et al., 2009). Since high concentrations of ammonia are toxic for humans (hyperammonemia) and for aquatic ecosystems, the quantification of ammonia becomes essential to diagnose an abnormality in nitrogen homeostasis in humans and is a key parameter for water quality, exploited for human consumption or use (like swimming pools), and for all the aquatic environments (Barsotti, 2001; Azmi et al., 2009; Mérian et al., 2014; Lin et al., 2019; Gogoi et al., 2021).

Ammonia commonly enters the environment through municipal, industrial, agricultural, and natural processes (Geadah, 1985). Ammonia concentrations in water vary seasonally and regionally. In natural waters, concentrations of total ammonia are generally less than 0.1 mg/L. Higher levels of ammonia normally indicate organic pollution (McNeely et al., 1979). In aquaculture systems, it is at most 5 mg/L (0.28 mM), while domestic sewage typically contains 20–40 mg/L of ammonia (1.1–2.2 mM). Dairy effluent, refinery effluent and mineral industry effluent may reach the 100–140 mg/L (5.5–7.8 mM) (Gogoi et al., 2021). However, many circumstances lead to a higher release of ammonia, like accidental spills, the use and disposal of cleansing agents that contain ammonia, urban runoff, accidental releases of ammonia-rich fertilizer, intensive farming, and the decomposition of livestock wasters (CEQG, 2011). For example, during the period from 1992 to 1995, groundwater samples at an old municipal landfill site in Grinsted (Denmark) contained ammonia levels higher than 1 g/L (>55 mM) (Kjeldsen et al., 1998). In 1991–1993, ammonia was found in the surface water collected at the Porto Alegre landfill (Brazil) at a maximum concentration of 1.63 g/L (90 mM), while the maximum amount of ammonia was 1.83 g/L (101 mM) in the leachate (Kuajara et al., 1997).

The methods to quantify ammonia are based normally on biosensors for concentrations up to 500 μmol/L, or assay kits (up to 800 μmol/L) (Azmi et al., 2009; Li et al., 2020), therefore the direct quantification of higher concentrations of ammonia is not possible.

On the other hand, there are biotransformations, for instance, where ammonia can be steadily produced on a millimolar scale as a by-product, so theoretically its continuous removal by a different enzyme can shift the reaction equilibrium towards the formation of the product of interest, increasing the overall biocatalytic efficiency of the desired reaction. Similarly, if the reagents and products of those biotransformations cannot be
Chimera to mimic the hexameric form, known to be the predominant with size exclusion chromatography (Pettersen et al., 2004). After mimicking the assembly shown in Supplementary Figure S6, the loop to connect both proteins was modelled using the DaReus server (Karami et al., 2019). The final PDB file has been obtained after local minimization with openMM software, which is a toolkit for molecular simulation using high performance GPU code (Eastman et al., 2017).

**Genetic Construction**

The pET28b(+) vector harboring the formate dehydrogenase gene was used as template for PCR amplification (Fwd: 5′-CTCAAGGGCATTTGCTTAGAGCTAAACTGAGCACCCACAC-3′; Rev: 5′-GCTAGCCG TGCCACACCGGC CGGATCTTTTTATCGTTTTAC-3′). The glutamate dehydrogenase gene was amplified from pRSETb-GluDH to obtain the insert fragments (Fwd: 5′-GGCGGTGTTGTTGCC AGCGCTAGCATGAGCAAGATGTGTGACAG- 3′; Rev: 5′- TGTTGTTGTGCTGCTAGGTGACCTTTACCAAGCAAT GCCGTTGAG-3′). The primers were designed to incorporate a short peptide linker in the fusion protein (italics) with flanking regions containing restriction sites (underlined). *BamHI* and *NheI* have been chosen for this purpose, since they are not contained in any other site of the entire construction and the translated amino acids follow the requirement described in the previous statement. Indeed, *BamHI* restriction site encodes glycine (GGA) and serine (TCC), while *NheI* translates to alanine (GCT) and serine (AGC). The length of the linker resulted to be a compromise where the distancing between the two protein domains was effectively supplied, and the primers involved in the gene cloning were not too long for being inefficient. The linker consisted of the following protein sequence: Gly-Ser-Gly-Gly-Gly-Ala-Ser (9 amino acid residues). The genetic construction was designed to include two additional restriction sites at the very beginning of FDH gene (*EcoRI*) and at the very end of GluDH DNA sequence (*SacI*). This creates a modular and versatile system which allows the exchange of the genes to develop other fusion proteins using FDH as a cofactor recycling system, as well as the incorporation of a third fusion domain which would be involved in the catalytic reaction system.

The genes were amplified by PCR using Q5 High-Fidelity DNA polymerase and the obtained fragments (insert and vector) were used for cloning the fusion construct with the Gibson Assembly Cloning kit. To transform host cells with the assembled product, 2 µl of the Gibson Assembly reaction mixture were added to the chemically competent *E. coli* DH5α cells (included in the kit), following the protocol given by the supplier. After overnight growth on a LB agar plate supplemented with 50 µg/ml of kanamycin, colonies were picked and grown in liquid LB with kanamycin (50 µg/ml). Then the plasmid was isolated (Thermo Scientific GeneJET Plasmid Miniprep kit) and sent for sequencing to confirm the correct genetic construction.

Through the Gibson assembly method, the linker and the glutamate dehydrogenase sequence have been joined with the sequence of the formate dehydrogenase to successfully assemble a
double-stranded fully sealed DNA sequence of the recombinant fusion protein.

**Fusion Protein Expression**

The transformed *E. coli* BL21 (DE3) cells containing the plasmid were cultured at 37°C in 5 ml of LB media supplemented with kanamycin (50 µg/ml) for an overnight. Subsequently, the culture was inoculated into 300 ml of fresh LB broth including kanamycin (50 µg/ml) and incubated at 37°C under shaking. At an optical density (OD₆₀₀) of 0.5–0.6, the flasks were subjected to a cold-shock stage for 30 min. Afterwards, the overexpression of recombinant fusion protein was induced by the addition of 0.1 mM of IPTG and the culture was grown at 20°C for an overnight. Cells were harvested by centrifugation (4,500 rpm, 20 min, 4°C).

**Fusion Protein Purification**

Cells were first resuspended in loading buffer (50 mM potassium phosphate buffer pH 7.5, 300 mM NaCl, 30 mM imidazole) and the resulting suspension subjected to sonication to disrupt cells (5 s pulse on and 10 s off for 8 min). Cell debris was removed by centrifugation (14,500 rpm for 45 min at 4°C) to yield a cell-free extract which was filtered (0.45 µm pore size). The affinity chromatography was performed using an AKTA Start system, equilibrated with the loading buffer. The crude extract was loaded (0.5 ml/min flow rate) onto a 5 ml His-trap Ni-afinity column (GE Healthcare) collecting the flow-through and washed with the loading buffer until the non-specific proteins were completely eluted. The collected solution was then applied to the column a second time to ensure that all the proteins were effectively bound, and the column was washed again. After that, an isocratic wash step with only 10% of elution buffer (50 mM potassium phosphate buffer pH 7.5, 300 mM NaCl, 300 mM imidazole) was performed to remove any non-specific protein still bound onto the column. Finally, recombinant proteins were eluted with 100% of elution buffer and dialyzed at 4°C under mild stirring, against storage buffer (50 mM potassium phosphate buffer pH 7.5). The dialysis was performed for an overnight replacing the buffer at least 2 times.

The purity of the obtained proteins was tested by 12% (w/v) SDS-PAGE and the concentration was quantified by Bradford assay.

**Fusion Protein Characterization**

One unit of activity was determined as the amount of enzyme needed to produce or consume 1 µmol of NADH per minute at 25°C. The activity assay was performed in a 96-well plate and detected by UV absorbance at 340 nm (ε₃₄₀ = 6.22 mM⁻¹ cm⁻¹). Each protein domain was evaluated singularly (GluDH: 10 mM α-ketoglutaric acid disodium salt and 200 mM ammonium chloride for reductive amination, 40 mM L-glutamic acid monosodium salt for oxidative deamination, in potassium phosphate buffer 50 mM pH 7.5; FDH: 100 mM sodium formate in potassium phosphate 50 mM pH 7.5). The assay was started by the addition of the cofactor in the reaction environment (0.2 mM of NADH or 1.0 mM of NAD⁺ as final concentration). The specific activity was calculated in U/mg considering the overall concentration of the fusion protein.

The kinetic constants were found in triplicate by measuring the dependence of initial rate on substrate concentration at 25°C in potassium phosphate buffer 50 mM (pH 8.0 for GluDH and pH 7.5 for FDH). For oxidative deamination reaction of GluDH, glutamic acid concentration was varied from 0.1 to 100 mM in the presence of 1 mM NAD⁺. For the reverse reaction, α-ketoglutaric acid concentration was ranging between 0.05 and 20 mM at a fixed concentration of 200 mM ammonium chloride; ammonium chloride was varied from 1 to 200 mM at 10 mM α-ketoglutaric acid. In both cases, the concentration of NADH was 0.35 mM. The kinetic constants for formic acid in the reaction of FDH were found by using a range between 1 and 100 mM formate concentration at 1 mM NAD⁺. The cofactor affinity was also evaluated by changing the concentration of NADH from 0.01 to 0.4 mM (10 mM α-ketoglutaric acid, 200 mM ammonium chloride) and 0.1–1.0 mM for NAD⁺ (40 mM glutamic acid for GluDH and 100 mM formic acid for FDH). Reactions were made by adding an appropriate amount of enzyme to the substrate solution and incubating the mixture for at least 10 min to allow the enzyme to adjust to the environment. The measurement started after mixing the cofactor and following the change in absorbance at 340 nm.

The stability and activity assays at different temperatures and pH were executed in triplicate by incubating the biocatalyst solution at different temperatures (pH 7.5) or at chosen pH values (temperature 25°C) and withdrawing samples at regular times for proceeding with specific activity assays of the fusion protein in parallel with the original GluDH and FDH. The activity of the enzyme was also checked using the previous protocol at several pH and temperature conditions. The reagents were prepared with the chosen pH or incubated at the chosen temperature before proceeding with the activity tests. The results were fitted in a graph showing the trend of the stability/activity of both glutamate and formate DH at chosen temperature or pH values.

**Size Exclusion Chromatography**

The purified fusion protein was applied to a gel filtration column (Superdex 200 10/300 GL, GE Healthcare), which was equilibrated with buffer (50 mM Tris/HCl, 150 mM NaCl, pH 7.5) and mounted onto an ÄKTA Pure instrument (GE Healthcare). The flow rate was set at 0.5 ml/min. A calibration curve was made by plotting the elution times of the following protein standards: carbonic anhydrase (29 KDa), albumin (66 KDa), alcohol dehydrogenase from yeast (150 KDa), β-amylase from sweet potato (200 KDa), apoferritin from horse spleen (443 KDa), thyroglobulin bovine (669 KDa). Blue dextran (2,000 KDa) was employed to determine the void volume and to check column packing. The elution of the standard proteins and the sample was followed by absorbance readings at 280 nm. The molecular weight of the fusion protein was calculated through the elution time fitted in the equation obtained from the calibration curve.

**Expression and Purification of Wild-Type Enzymes**

The CsGluDH-pRSETb and the CbFDH-pET28b were separately used to transform *E. coli* BL21 STAR, which were then streaked
onto a LB agar plate containing the antibiotic (100 µg/ml ampicillin for GluDH and 50 µg/ml kanamycin for FDH). A colony from each plate was inserted in 300 ml autoinduction media containing the proper antibiotic and incubated at 37°C for 8 h. Afterwards, the Erlenmeyer flasks were incubated on ice for 30 min (cold shock) and left at 30°C (GluDH) or 25°C (FDH) for the overnight growth. Cells were harvested by centrifugation and pellet were resuspended with loading buffer to proceed with the purification (same buffers as fusion protein). The cells suspensions were sonicated and centrifuged to obtain cell extracts that were filtered and loaded onto a 1 ml His-trap FF crude Ni-affinity column on the ÄKTA start for the purification. The purity of the protein solutions was then checked by SDS-PAGE and quantified by Bradford assay. The activity was measured as described for the fusion protein.

**Enzyme Immobilization**

The chosen support was left in incubation with the enzyme solubilized in the appropriate buffer. Typically, 1 g of support was added to 1 ml of a solution containing the desired amount of protein and incubated at room temperature under mild agitation. The protein load is defined as the amount of protein that has been immobilized per Gram of carrier (mg/g).

Small amounts of supernatant were taken over time to detect enzymatic activity, that was compared to the activity of the initial enzyme solution that was offered to the carrier (control sample). To better confirm the full immobilization, the protein concentration was then checked by Bradford assay and SDS-PAGE. When the maximum achievable yield was obtained, the support was washed.

The activity assay was performed by adding 5–10 mg of immobilized enzyme to a 2 ml reaction mixture containing substrates and cofactor and monitoring it using the same procedure as for the activity assay of the free enzyme. The aim is to determine the expressed activity in U/g (units of enzyme catalyzing the formation of 1 µmol of product per minute per Gram of carrier (mg/g)).

After washing, the support was incubated with 5 ml of metal buffer (1 M NaCl, 5 mg/ml CoCl2 in 50 mM phosphate buffer pH 6) for 2 h. The support was then washed and incubated with the protein solution in phosphate buffer 50 mM pH 8.5. The support was then washed with 3 ml of desorption buffer (50 mM EDTA, 500 mM NaCl in 50 mM phosphate buffer pH 7) to remove the metal and rinsed with deionized water. The support was left in 4 ml blocking buffer (3 M glycerol in 20 mM phosphate buffer pH 8.5) for an overnight and then washed (Mateo et al., 2007).

For the covalent immobilization on epoxy groups by interaction with ethylenediamine (EDA), 1 g of the epoxy resin was incubated with 6 ml of 2% (v/v) EDA in 100 mM NaHCO3 buffer pH 8.5 for 2 h. The support was then washed (water, 1 M NaCl and again with water). The protein solution was prepared in 5 mM phosphate buffer pH 7. After being washed, the immobilize enzyme was incubated with 1 ml of 5 mM NaHCO3 buffer pH 9 for at least 2 h. Afterwards, the support was washed and left in 4 ml blocking buffer (prepared as previously described) for an overnight (Trobo-Maseda et al., 2020).

For the coating of epoxy and glyoxyl resin with polyethyleneimine (PEI), a solution of 10 mg/ml of PEI 60 kDa was prepared in 100 mM NaHCO3 buffer pH 10 and 10 ml were incubated with 1 g of support for an overnight. After that, the epoxy resin was washed and stored, while the glyoxyl resin was incubated with 1 mg of NaBH4 in 10 ml of 100 mM NaHCO3 buffer pH 9 for at least 2 h. After washing, the support was washed and left in 4 ml blocking buffer (prepared as previously described) for an overnight (Trobo-Maseda et al., 2006; Velasco-Lozano et al., 2017).

For the qualitative assay with picrylsulfonic acid, an adapted version of a previously reported protocol was used (Guisán, 1988). 200 µl of 100 mM NaHCO3 buffer pH 10 were added to 20 mg of resin. Then, 20 µl of picrylsulfonic acid were inserted and left in incubation for 10 min. The support was filtered, washed 3 times with 500 µl of saturated NaCl and 3 times with 500 µl of 100 mM NaHCO3 buffer pH 10.

**Immobilization of the Fusion Protein**

The EP400/SS resin was firstly activated with glyoxyl groups and then coated with PEI 60 KDa. Several protein loadings were tested: 1, 5 and 10 mg per Gram of resin.

**Immobilization of the Wild-Type Enzymes**

Different strategies of immobilization were chosen, using the resins ReliSorb EP400/SS (Resindion S.R.L.) and Lifetech ECR8204F, ECR8304F, ECR8285 (Purolite Ltd.). The resins EP400/SS and ECR8204F were activated to obtain aldehyde groups. A part of them was employed for the covalent immobilization on glyoxyl groups, while the other part was functionalized with PEI, 60 KDa for the respective immobilization. The same epoxy resins were also modified with iminodiaceitic acid and cobalt, with EDA or with PEI for the relative immobilizations. The biocatalysts were also immobilized directly to Lifetech ECR8304F and ECR8285 (protein eluted in 5 mM phosphate buffer pH 7). All the carriers presenting amino groups were qualitatively analyzed using picrylsulfonic acid as reagent.
The immobilization yield and the recovered activity were calculated, by following the same conditions as for the activity assays (oxidative deamination direction for GluDH).

For the co-immobilization, ReliSorb EP400/SS was firstly treated to get glyoxyl groups and after that, it was coated with PEI 60 KDa. The two enzymes were incubated with the carrier both simultaneously and stepwise adding firstly GluDH and then FDH, and vice versa, to compare any difference in recovered activity. Different amounts of enzymes as well as different ratios were loaded.

**Operational Stability of the Immobilized Fusion Protein**

The immobilized protein was tested in terms of operational stability by performing consecutive biotransformations and checking the decrease of relative rate of conversion for each cycle of reaction.

The biotransformations were prepared in 1 ml total volume containing 50 mM α-ketoglutaric acid disodium salt, 200 mM ammonium chloride, 200 mM sodium formate and 1 mM NADH in potassium phosphate buffer 50 mM pH 7.5, adding 10–40 mg of immobilized biocatalyst. The reaction was left in incubation at 37°C under agitation for 30 min, then the supernatant was taken for sample preparation and subsequent GC-FID analysis.

**Biotransformations**

Enough free or immobilized biocatalyst was added to a 2 ml tube containing 1 ml of reaction mixture (50/300/400 mM α-ketoglutaric acid disodium salt, 200/400/300 mM ammonium chloride, 200/400 mM sodium formate and 1 mM NADH in 50 mM potassium phosphate buffer pH 7.5). A control without enzyme was run in parallel. The mixture was then incubated at 37°C under agitation (150 RPM) and aliquots were taken over time.

**Analytical Method**

Samples were prepared and derivatized following the available protocol (Walsh et al., 2014) employing methyl chloroformate as reagent, in combination with methanol and pyridine. The reaction components (derivatized glutamic acid and derivatized α-ketoglutaric acid) were extracted in chloroform and submitted to the GC-FID analysis (temperature of inlet: 250°C, FID detector: 275°C). The column oven temperature was initially held at 75°C for 1 min, then increased to 200°C with a ramp of 25°C per minute and a final hold of 4 min, using nitrogen as gas carrier with a flow of 6.5 ml/min and CP-Chirasil-Dex CB 25m x 250 µm x 0.25 µm as GC column.

The production of glutamic acid and the consumption of α-ketoglutaric acid were monitored and compared between free and immobilized biocatalytic reaction to assess the efficiency of the system.

**RESULTS AND DISCUSSION**

**Rational Design of the Genetic Construction**

The genetic construction of the recombinant fusion protein has been developed using the original plasmid encoding the FDH from *Candida boidinii* (pET28b-FDH) as a template, which contains an N-terminal His-tag (Schütte et al., 1976; Padrosa et al., 2021). The gene of the GluDH from *Clostridium symbiosum* was inserted downstream of FDH DNA sequence (with removal of the FDH stop codon), to enable the correct folding of the GluDH C-terminal domain which loops into the internal cavity of the hexameric complex when the quaternary structure is formed (Rice et al., 1985; Syed et al., 1991; Baker et al., 1992). The construct (N-His₆-tag)-FDH-linker-GluDH is shown in Figure 2.

A 9 aa flexible linker was inserted between the two protein domains to facilitate folding (Chen et al., 2013). Linkers are in fact ubiquitously observed in naturally occurring multidomain proteins with the function of maintaining the necessary distance to reduce steric hindrance and/or permit favorable domain-domain interaction between the two protein moieties (Yu et al., 2015).

The linker has been designed to include two flanking restriction sites to facilitate direct modification of the genetic sequence (length of the linker, flexibility, amino acids involved, etc.) and versatility of the construct (see Materials and Methods).

Computational studies have been performed to probe the rationale behind the gene design, before proceeding with the experimental development of the fusion protein. The PDB file resulting from the in-silico analyses supported the insertion of such linker to facilitate the correct assembly of each subunit. As shown in Figure 3, the six subunits are favored to assemble with the hexameric GluDH located in the center (green color) of the quaternary structure. The fused FDH subunits are exposed to the bulk to form three assembled dimers (pink color) enabling proper folding without any major distortion.

The FDH-GluDH fused gene was successfully created via Gibson assembly and confirmed by sequence analysis.

**Protein Expression and Purification**

Several conditions to maximize overexpression have been tested (data not shown). LB media induced with 0.1 mM of IPTG at 20°C
The two domains exhibited very similar activity to the wild-type free proteins with an average of 86.4% of GluDH and an 80% of FDH activity. This outcome is consistent with a previous report where the FDH moiety retained less parental activity than partner enzyme (Zhang et al., 2017). Likewise, around an 80% of retained activity was attained by the FDH of that fusion protein, which was presenting the flexible linker (GSSSS)$_2$, very similar to the linker of the current study.

The predicted assembly suggested by the molecular modelling has been therefore validated experimentally. The subunits of the two domains were clearly assembled without any major distortion, meaning that the GluDH possessed a hexameric quaternary structure as the wild-type, while the FDH presented a dimeric configuration. The size exclusion chromatography further reinforced the accuracy of the molecular model, showing a predominant hexameric and octameric form with traces of assembled 2 and 12 monomers (see paragraph 3 in the Supplementary Material for more details). The kinetic constants for all substrates were determined and the results are shown in Table 2. The assays of GluDH were performed at pH 8.0 in order to provide a better evaluation of the results of this work with respect to previously reported values (Sharkey and Engel, 2009), as far as the WT GluDH. The calculated results were found to be comparable. On the contrary, the specific activity was measured at pH 7.5 as compromise between stability/activity of both domains of the fusion protein. The different pH numbers were the reason why the values presented in Table 1 for GluDH were dissimilar from the values of the same enzyme shown in Table 2.

The resulting $K_M$ values were comparable between the fusion protein and the wild-type enzymes; the affinity to the substrates has not been affected where the fusion of the two proteins could have destabilized the quaternary structure. The FDH domain showed a lower $K_M$ value towards NAD$^+$ compared with that of the free FDH, while the GluDH had a decreased affinity for the NADH after the fusion. The same alteration of the cofactor kinetic parameters was found in the LeuDH-FDH fusion protein as well as in the KR fused with the mutant formate dehydrogenase MycFDH (Sührer et al., 2014; Zhang et al., 2017). It has been suggested that the different affinity values towards the cofactor were caused by moiety-moiety interactions or the proximity of two cofactor binding domains in the fusion enzymes (Zhang et al., 2017).

The maximum velocities ($V_{max}$) of the fusion protein were consistently half of those of GluDH, and 3 to 5 times lower than FDH, but this simply reflects the fact that the weight of the fusion protein is approximately twice that of the wild-type enzymes, therefore also the rate of conversion is conserved. The bifunctional protein retained the overall catalytic properties of the individual enzymes, as seen for some fused enzymes. In other cases,

| Characterization |
|------------------|
| The specific activity of the fusion protein was determined for both catalytic domains (GluDH and FDH) and compared with the activity of the single enzymes. The monomeric molecular mass of the free enzymes (GluDH 53 and FDH 43 KDa) is about half than the fusion protein (93 KDa), therefore the results are reported in Table 1 normalized for concentration of each individual enzyme (see paragraph 2 in the Supplementary Material for calculations). |

**Table 1| Specific activity of the two protein domains compared to the normalized specific activity of the wild-type FDH and GluDH (shown only as reference values without the standard deviations).**

| Formate dehydrogenase | Glutamate dehydrogenase |
|------------------------|-------------------------|
| **Amination**          | **Deamination**         |
| Fusion protein          | 1.2 ± 0.2 [U/mg]        | 135 ± 6 [U/mg]     |
| Wild-type enzyme        | 1.5                     | 17.0 ± 0.5 [U/mg]  |
|                        |                         | 156                 |
|                        |                         | 19.7                |
TABLE 2 | Kinetic constants for the fusion protein, the glutamate- and the formate dehydrogenase.

| Fusion protein–GluDH       | Wild-type CsGluDH   |
|-----------------------------|--------------------|
|                            |                   |
| **Fusion protein–GluDH**    | **Wild-type CsGluDH** |
|                            |                   |
| KM [mM]                    | KM [mM]          |
| V max [mmol min\(^{-1}\) mg\(^{-1}\)] | V max [mmol min\(^{-1}\) mg\(^{-1}\)] |
| L-glutamate                | 2.9 ± 0.8         | 3.0 ± 1.1      |
|                           | 0.04 ± 0.01       | 0.08 ± 0.01    |
| α-ketoglutarate            | 0.41 ± 0.22       | 0.48 ± 0.31    |
|                           | 0.07 ± 0.05       | 0.13 ± 0.1     |
| Ammonia                    | 32.5 ± 10.0       | 32.3 ± 10.0    |
|                           | 0.18 ± 0.06       | 0.36 ± 0.15    |
| NADH                       | 0.20 ± 0.08       | 0.10 ± 0.06    |
|                           | 0.06 ± 0.05       | 0.12 ± 0.1     |
| NAD\(^+\)                  | 0.17 ± 0.04       | 0.17 ± 0.02    |
|                           | 0.02 ± 0.01       | 0.05 ± 0.02    |
| **Fusion protein–FDH**     |                   |
|                            |                   |
| KM [mM]                    |                   |
| V max [mmol min\(^{-1}\) mg\(^{-1}\)] |                   |
| Formate                    | 6.0 ± 1.0         | 5.0 ± 1.0      |
|                           | 0.0010 ± 0.0003   | 0.0003 ± 0.001 |
| NAD\(^+\)                  | 0.088 ± 0.014     | 0.168 ± 0.030  |
|                           | 0.0010 ± 0.0004   | 0.0005 ± 0.001 |

However, such as for some ADH-CHMO fusion proteins, the catalytic efficiency was dramatically diminished, possibly due to the orientation of the two domains or the fusion itself that hindered the association between the subunits to form the oligomers. It is in fact possible that one protein domain covers the oligomerization interface of its fusion partner, preventing the multimeric assembly, essential for the activity (Aalbers and Fraaije, 2017). In the case of the GluDH-FDH fusion protein, the domains were able to interact properly without any structural impact.

The fusion protein presented the same profiles as the wild-type enzymes for both pH and temperature activity/stability (details in paragraph two of ESI). Furthermore, the FDH moiety acquired a cooperative behavior that is typical of the GluDH but does not occur in the WT FDH. The fusion, therefore, led to a tight connection between the two enzymes where one domain is dependent to the other for structural modifications induced by temperature shift, for example.

Enzyme Immobilization

To further probe the applicability of this new assembly, the fusion protein was immobilized on a solid support. Preliminary immobilization tests on the individual WT enzymes, based on a trial-and-error approach, showed that all the immobilization strategies based on covalent bonds were exhibiting almost no recovered activity (Supplementary Figures S11, 12). Attempts of immobilization on the resins ECR8204F, ECR8304F, ECR8285 from Purolite did not result in any improvement, giving very low activity with either covalent, hydrophobic, or ionic interactions. Instead, the best option for both WT GluDH and WT FDH was the immobilization on ReliSorb EP400/SS (Resindion) through ionic bonds with polyethyleneimine (PEI) polymer, that covered all the surface of the support. In this case, the WT GluDH showed 85% recovered activity, while the WT FDH exhibited a recovered activity of 100% (at 1 mg/g of loaded protein) with a complete immobilization yield. Therefore, the same support (EP400/SS) and chemical strategy (ionic interactions with PEI) have been applied for the co-immobilization of the WT enzymes, both as simultaneous addition of the two protein solutions and as sequential approaches by first immobilizing one of the two enzymes and then the second one, and vice versa. No decrease from the previously found 85 and 100% recovered activity was detected when the immobilization was performed sequentially, firstly with the binding of the GluDH followed by the addition of the FDH.

Recently, a python-based GUI-application called CapiPy has been developed to assist in protein immobilization, which supplies information to rationalize the choice of immobilization strategy (Roura Padrosa et al., 2021).

CapiPy offered an initial evaluation of the amino acid residues available on the surface of the enzyme through a detailed analysis of the hexameric structure, whose model was given by minimizing a manually assembled hexameric structure followed by minimization with openMM (Eastman et al., 2017). The final quaternary structure is shown in Figure 4, where it is evident that negative charged groups (red) are spread throughout the surface of the hexamer, offering many points of interactions with positively charged groups. Therefore, the functionalization of a support with amino groups would offer easy interaction with the aspartate and glutamate clusters of the fusion protein.

The polymethylacrylate epoxy resin ReliSorb EP400/SS has been selected as a support, and it provides a highly porous structure. The material of the resin offers a robust support to the enzyme, while the porosity gives many points of anchoring. Polyethyleneimine (PEI) has been used to coat the carrier, to enhance the enzyme binding through the high number of amino groups introduced and create a more hydrophilic environment.

The enzyme was efficiently bound through ionic interactions between the amino groups of PEI and the exposed carboxyl groups of the enzyme (aspartate and the glutamate residues). This favors the retention of enzyme flexibility which is essential for its activity since it undergoes a conformational change from an “active” to an “inactive” form, where the six subunits have the capacity for cooperative interaction (Syed and Engel, 1990; Wang and Engel, 1995).

A complete immobilization yield was achieved, with no remaining protein in the supernatant. As indicated in Figure 5, a recovered activity of 50 and 100% for GluDH and FDH, respectively, was obtained with a loading of 1 mg of protein per Gram of support. Differently from the sequential co-immobilization of the WT enzymes, the two domains of the fusion protein have been simultaneously immobilized since their fusion did not allow for a sequential approach. Increasing the loading to 5 mg/g of support caused a slight decrease to 35% for GluDH and 85% for FDH in recovered activity. This was further reduced at 10 mg/g of loading (30% for GluDH and 60% for FDH) indicating a diffusion limitation issue.
α-Ketoglutaric Acid Biotransformations With Free and Immobilized Fusion Protein

The enzyme was trialed first for its potential application in the synthesis of glutamic acid at different concentrations of starting material (α-ketoglutarate) and sub-stoichiometric amounts of NADH (1:10, 1:50, 1:300). In this case, the ammonia and formate substrates are kept in excess with respect to the α-ketoglutaric acid (see Materials and Methods). The biotransformation was performed with the immobilized fusion protein, which was compared to the results attained from the biocatalytic reaction done with the free bifunctional protein. Initially, a 5 mg/g immobilized enzyme with 10 mM of α-ketoglutarate and 1 mM NADH yielded 72% molar conversion in 30 min, reaching completion in less than 1 hour. The same results have been seen for an equal amount of soluble protein in the same conditions of biotransformation.

At a 50 mM scale of α-ketoglutaric acid (1 mM NADH), the free protein and the immobilized form appeared to biocatalyse similarly the reaction, showing analogous conversion rates (Figure 6) with equal amounts of biocatalyst. As a matter of fact, both forms showed a complete conversion in less than 4 h, demonstrating the high efficiency of the system. No significant limitation due to the less efficient FDH was notable. Under these conditions, the operational stability of the immobilized protein was tested through repeated usage, showing that 50% of the activity was still retained after 10 cycles (Supplementary Figure S9).

Then, a large-scale biotransformation of 300 mM of α-ketoglutaric acid (Figure 7) with still just 1 mM of NADH...
was set up to test the cost-effectiveness and to push the self-sustainability of the system. Full conversion was obtained in less than 40 h using the immobilized enzyme, while the soluble biocatalyst achieved 70% conversion within the same timeframe. Some protein precipitation was noted in the reaction environment with the free form, causing a decrease of activity over time. The loss of stability was probably caused by the presence of high substrates concentration in the solution (300 mM α-ketoglutarate disodium salt, 400 mM ammonium chloride, 400 mM sodium formate and 1 mM NADH). However, the immobilized enzyme worked efficiently reaching a 99% conversion, suggesting that the immobilization effectively enhanced the stability of the bifunctional system over time even at higher substrates concentration.

**Bio-Removal of Ammonia as Proof of Concept**

The GluDH-FDH fused system could be envisaged as a tool to remove ammonia, for example in contaminated waters following spillages. To better evaluate the efficiency of the system in such potential application, different experiments where the ammonia must be fully depleted have been designed. The removal of ammonia was determined by measuring the consumption of the α-ketoglutarate (substrate provided in excess) combined with the rate of production of the glutamate, since the ammonia employed for the amination is stoichiometric with the two consumed/formed components.

Firstly, to compare the performance of GluDH alone and the GluDH in the fused assembly, with or without the assistance of the FDH, a low amount of ammonia (15 mM) was supplemented with an excess of α-ketoglutarate (30 mM) and the catalytic efficiency tested in the absence or presence of the formate substrate, with either stoichiometric or catalytic amounts of NADH. The reactions were monitored at 1, 2, and 24 h but, in all cases, after 1 h no further change in the ammonia elimination was observed. The fusion protein emerged to be more efficient (Table 3), achieving full depletion of ammonia only when the FDH is activated (entry 2). This result is particularly encouraging as at 15 mM, the concentration of ammonia is significantly below the calculated K_M value (Table 2). In contrast, the GluDH in both fused and wild-type form achieved only 80% conversion, in the absence of FDH activation, even when the NADH was provided in a stoichiometric amount (entry 1).

While the fusion protein fared very well with low concentration of ammonia, the robustness of the system was tested under significantly harsher conditions. A second set of reactions was set up with 300 mM ammonia together with a moderate excess of α-ketoglutarate (400 mM) and formate (400 mM), and 1 mM NADH. Surprisingly, only the immobilized fusion protein showed a >99% conversion in 48 h (Figure 8). The soluble fusion protein appeared to have a slower pace but achieved almost 80% in 72 h. To complete the comparative evaluation, the individual WT enzymes were also included in this experiment. Both a combination of free forms and a co-immobilized version (all calculated to be in the same ratio and concentration of the fusion protein) were trialed. Surprisingly, the combination of the wild-type soluble proteins reached only 20% of conversion, suggesting a stability issue of at least one of the two enzymes, reinforcing the hypothesis that the fused assembly infers overall a higher structural stability to the catalysts. This is much more apparent in high ionic strength conditions. The co-immobilized enzymes however, also showed improved performance, mimicking the behavior of the free fusion protein, confirming the general higher stability of immobilized systems.

Despite the general improved stability attained with the immobilization method, the conversion rates achieved in 48 h with the immobilized fusion protein and the co-immobilized WT enzymes are clearly divergent (>99 versus 65%, respectively) with 1.5-fold higher rate for the bifunctional protein. The performance in working conditions provides a suitable method of comparison of the catalytic efficiency of the two systems (with and without the fusion) where the new enzyme is more effective timewise.

Looking at the literature, also the PheDH-FDH fusion protein showed a 1.5-fold higher molecular conversion rate than the individual enzymes, due to the improved cofactor regeneration

### TABLE 3 | Percentage of conversion of 15 mM ammonia without (first line) and with (second line) NADH recycling.

| Entry | Ammonia | α-ketoglutarate | NADH | Formate | Conversion [%] |
|-------|---------|-----------------|------|---------|----------------|
|       |         |                 |      |         | Wild-type CsGluDH | GluDH domain |
| 1     | 15      | 30              | 15   | /       | 80%             | 80% |
| 2     | 15      | 30              | 1    | 100     | /               | 100% |
process, where the product of the first reaction can be quickly transferred to the second enzyme, which leads to a lower time of diffusion (Jiang and Fang, 2016).

Consequently, the bifunctional enzyme gained remarkable enhancements, especially in the performance, reducing the reaction duration and intensifying the overall conversion degree. The fusion protein results a superior biocatalyst than the soluble or co-immobilized wild-type enzymes under several points, from the time- and cost-effectiveness production (including transformation, expression, purification, immobilization steps) to the enhanced biocatalytic reaction.

CONCLUSION

The GluDH from Clostridium symbiosum and the FDH from Candida boidinii have been successfully fused to yield one active, stable, and self-sufficient protein. The new bifunctional enzyme has been created overcoming all the possible drawbacks that may appear during the production of a fusion protein. Moreover, the close proximity of the two biocatalysts, given by the direct linkage, offered a great enzymatic tool for the application in small and high scale conversion of ammonia and α-ketoglutarate. The fusion protein was also immobilized, providing a highly efficient biocatalyst, that can be reused for several cycles of reaction. In the future, the design and development of a protein possessing the activity of two different enzymes may surpass the already advantageous co-immobilization for a more industrially appealing solution, that fulfill the demand for sustainable processes. The novel fusion protein has proved to remove ammonia efficiently in experimental conditions, as proof of concept. The treatment of environmental samples and the coupling to other reaction systems where ammonia is the by-product are still potential applications by now. Nevertheless, this work provides a useful starting point for further studies and future applications.

DATA AVAILABILITY STATEMENT

The original contributions presented in the study are included in the article/Supplementary Material, further inquiries can be directed to the corresponding author.

AUTHOR CONTRIBUTIONS

FP conceptualized the idea and supervised the project, VM performed the experiments and the characterization studies, and wrote the initial draft; DRP performed the MD analysis, AIB-M designed and supervised the genetic experiments and the immobilization studies. All the authors discussed the results and reviewed the manuscript.

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SUPPLEMENTARY MATERIAL

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