Simple and efficient expression of *Agaricus meleagris* pyranose dehydrogenase in *Pichia pastoris*

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**Abstract** Pyranose dehydrogenase (PDH) is a fungal flavin-dependent sugar oxidoreductase that is highly interesting for applications in organic synthesis or electrochemistry. The low expression levels of the filamentous fungus *Agaricus meleagris* as well as the demand for engineered PDH make heterologous expression necessary. Recently, *Aspergillus* species were described to efficiently secrete recombinant PDH. Here, we evaluate recombinant protein production with expression hosts more suitable for genetic engineering. Expression in *Escherichia coli* resulted in no soluble or active PDH. Heterologous expression in the methylotrophic yeast *Pichia pastoris* was investigated using two different signal sequences as well as a codon-optimized sequence. A 96-well plate activity screening for transformants of all constructs was established and the best expressing clone was used for large-scale production in 50-L scale, which gave a volumetric yield of 223 mg L$^{-1}$ PDH or 1,330 UL$^{-1}$ d$^{-1}$ in space–time yield. Purification yielded 13.4 g of pure enzyme representing 95.8% of the initial activity. The hyperglycosylated recombinant enzyme had a 20% lower specific activity than the native enzyme; however, the kinetic properties were essentially identical. This study demonstrates the successful expression of PDH in the eukaryotic host organism *P. pastoris* paving the way for protein engineering. Additionally, the feasibility of large-scale production of the enzyme with this expression system together with a simplified purification scheme for easy high-yield purification is shown.

**Keywords** Pyranose dehydrogenase · *Pichia pastoris* · Heterologous expression · *Agaricus* · Large scale · Microscale

**Introduction**

Pyranose dehydrogenase (PDH; pyranose/acceptor oxidoreductase, EC 1.1.99.29) is a flavin-dependent sugar oxidoreductase (Völc et al. 1997, 2001; Kujawa et al. 2007; Sygmund et al. 2008; Morris et al. 1999). It is an extracellular, glycosylated enzyme mainly found in a relatively narrow group of litter-decomposing fungi belonging to the Agaricaeae (Völc et al. 2001). The biological role of pyranose dehydrogenase in fungi still remains unclear although the enzyme is thought to participate in lignocellulose degradation. The functions suggested for PDH are the reduction of quinones or radical intermediates formed during lignocellulose degradation preventing their repolymerization (Ander and Marzullo 1997) as well as the production of Fenton’s reagent via reduction of metal ions (Kerem et al. 1999) which may attack cellulose, xylan, and lignin.

Pyranose dehydrogenase catalyzes the oxidation of free, nonphosphorylated sugars and the transfer of resulting electrons to a number of (substituted) quinones and...
complexed metal ions rather than to molecular oxygen. It exhibits extremely broad sugar substrate specificity, while in contrast only a rather limited number of electron acceptors are used by the enzyme. Unlike other sugar oxidoreductases like cellobiose dehydrogenase or pyranose 2-oxidase, which exclusively oxidize sugars at the C-1 and C-2, respectively, PDH is able to oxidize at C-1 to C-4 with C-2 and/or C-3 predominating, depending on the structure of the sugar (mono-, di-, oligo-saccharide, or glycoside) and the source of the enzyme.

Because of these unique properties, PDH is of interest for a number of technological applications, recently reviewed by Peterbauer and Volc (2010). Among these is the conversion of carbohydrates to interesting products for the food industry, e.g., the conversion of D-galactose to D-tagatose via 2-keto-D-galactose (Giffhorn 2000; Giffhorn et al. 2000). Plain di/tricarbonyl sugars as the main reaction products of PDH could serve as basis for the synthesis of amino sugar moieties of homologues to amino glycoside antibiotics (Reitz and Baxter 1990). The relaxed substrate specificity of PDH makes it an attractive target for further improvements of its reactivity with derivatives of already established substrates such as aromatic glycosides, which may provide new attractive building blocks for synthesis of carbohydrate-based drugs.

A different, very promising application for PDH is its use as anodic biocatalyst in enzymatic biofuel cells, which transform chemical to electrical energy via electrochemical reactions (Barton et al. 2004; Bullen et al. 2006). PDH was successfully “wired” to electrodes using osmium redox polymers and was shown to be suitable for this application (Tasca et al. 2007; Zafar et al. 2010). Apart from being able to provide a second pair of electrons per sugar molecule if double oxidation occurs, PDH’s inactivity with molecular oxygen can be considered an advantage. Unlike in biofuel cells, employing glucose oxidase or pyranose oxidase no harmful H2O2 is being formed as a byproduct, avoiding potential oxidative damage to the enzyme or the redox polymer, thereby prolonging the endurance of the biofuel cell. A possibility to increase the power output of a PDH biofuel cell would be to lower the redox potential of the enzyme. This, as well as increasing the activity of PDH with technologically important substrates, could be targeted by protein engineering.

In the past few years, rational and semirational design approaches have shown to be very useful in improving enzyme properties (Lutz 2010). Crucial for success is expression of site-directed mutants or saturation mutagenesis libraries of the target gene with high yields in a genetically easy to manipulate expression host (bacteria and yeasts).

The gene of PDH1 from Agaricus meleagris was the first identified and isolated PDH-encoding gene (Kittl et al. 2008) and was recently expressed in Aspergillus spp. (Pisanelli et al. 2010). Hereby, we describe for the first time the successful heterologous expression of a PDH gene in the methylotrophic yeast Pichia pastoris and the scalability of production and purification of the enzyme on larger scale.

Materials and methods

Chemicals and microorganisms

All chemicals were of the highest purity grade available and were purchased from Sigma (St. Louis, MO, USA) unless stated otherwise. Restriction endonucleases, T4 DNA ligase and DNA-modifying enzymes were obtained from Fermentas unless stated otherwise and were used as recommended by the manufacturer. Nucleic acid amplifications were done employing Phusion High-Fidelity DNA Polymerase (New England Biolabs, Ipswich, MA, USA), dNTP mix, oligonucleotide primers (Invitrogen, Carlsbad, CA, USA), and a GeneAmp PCR System 2700 thermocycler (Applied Biosystems, Foster City, CA, USA). The pdh1 gene coding for pyranose dehydrogenase from the basidiomycete A. meleagris (GenBank accession number AY753307) has been previously isolated and cloned into the pCR Blunt II TOPO vector (Kittl et al. 2008). Escherichia coli strain DH5α-T1 was used for subcloning and BL21 DE3 for expression (Invitrogen). P. pastoris strain CBS 7435 was used for expression. Ferrocenium hexafluorophosphate and the substituted quinones were from Aldrich (Steinheim, Germany). Phenyl-Sepharose Fast Flow resin was purchased from Amersham Pharmacia Biotech (Uppsala, Sweden) while DEAE Sepharose Fast Flow resin was from GE Healthcare (Chafont St. Giles, UK). PNGaseF was purchased from NewEnglandBiolabs.

Construction of a pdh1 expression vector for E. coli

The pdh1 gene from A. meleagris was amplified from the published plasmid pdh1 in pCR Blunt II TOPO vector (Kittl et al. 2008) using the primers 5pdhN-Nde and 3pdhN-Iai and ligated into the equally treated expression vector for E. coli (Kittl et al. 2008). The polymerase chain reaction (PCR) fragment was cut with the restriction enzymes NdeI and HindIII and ligated into the equally treated expression vector pET21a. The resulting plasmid pET21apdh1 was transformed into electrocompetent E. coli BL21 DE3 for expression under control of the T7 promoter.

Expression of pdh1 in E. coli

E. coli cells carrying the pET21apdh1 plasmid were grown in TB medium (12 gL−1 peptone from casein, 24 gL−1...
yeast extract and 4 mL L\(^{-1}\) glycerol in 100 mM potassium phosphate buffer, pH 7.5) and induced with two concentrations of IPTG (0.01, 0.10 mM) as well as 0.5% lactose at a 100-mL scale in baffled flasks at 140 rpm and 25 °C overnight. Ten milliliter of the cultures were centrifuged at 4,000×\(g\) for 10 min at 4 °C, the cell pellets were resuspended in breaking buffer (50 mM potassium phosphate buffer pH 6.5 supplemented with 300 \(\mu\)M PMSF) and broken using a French press. After ultracentrifugation (30,000×\(g\), 30 min, 4 °C) the crude extract was tested for enzyme activity and the pellet was analyzed for insoluble PDH. Sodium dodecyl sulfate polyacrylamide gel electrophoresis (SDS-PAGE) was performed on commercial 8–25% acrylamide gels with silver staining of protein bands. Electrophoresis and the staining procedure were done with the PhastSystem (Pharmacia) according to manufacturer’s recommendations.

Codon optimization of \(pdh1\) for expression in \(P.\) pastoris

The open reading frame of the \(pdh1\) gene from \(A.\) meleagris and the \(Saccharomyces\) \(cerevisiae\) \(\alpha\)-factor sequence were codon-optimized with the program GeneDesigner from DNA2.0 (https://www.dna20.com/genedesigner2) employing the high-methanol codon usage of \(P.\) pastoris (Abad et al. 2010), which was calculated based on the genes for alcohol oxidase1 (AOX1), DAS, FLD1, and HbHNL using the DOS program GCUA (http://gcua.schoedl.de). Additionally, mRNA 3′-end processing signals and restriction enzyme recognition sequences (\(EcoRI, NotI, BglII, SacI\)) were eliminated. The resulting sequences were checked with the program Leto1.0 from Entelechon (www.entelechon.com) for avoiding strong stem-loop structures of the mRNA that were identified by GeneBee (http://www.genebee.msu.ru/). The synthetic gene was ordered from GenScript Corporation (Piscataway, NJ, USA) with the native leader sequence and the flanking restriction sites \(EcoRI\) and \(NotI\). For sequences, see Electronic supplementary material.

Construction of \(pdh1\) expression vectors for \(P.\) pastoris

The \(pdh1\) cDNA and the synthetic gene were amplified with the primer pairs 5pdhN-Asc/3pdhN-Not and 5pdhS-Eco/3pdhS-Not, respectively. Both PCR products were digested with the respective restriction enzymes and ligated into the equally treated vector pPpT2 (Abad et al. 2010; Ruth et al. 2010). Replacement of the \(\alpha\)-factor-derived sequence with that of the native signal sequence was done by overlap extension PCR. To this end, the whole \(\alpha\)-factor pre-pro leader sequence was amplified with the primers 5alpha-Eco and 3alpha for fusion with the synthetic gene and with 5alphaAsc and 3alpha for fusion with the native gene. These PCR products were finally fused to the corresponding sequences encoding the mature proteins that were amplified with 5pdhS-alpha/3pdhS-Not or 5pdhN-alpha/3pdhN-Not. The resulting PCR products were cut with the respective restriction enzymes and ligated into the equally digested shuttle vector pPpT2. An overview of the constructs is presented in Table 2. After transformation of the constructs into chemically competent \(E.\) coli DH5\(\alpha\)-T1 cells, the plasmids were proliferated, linearized with the respective restriction enzymes, and transformed into electrocompetent \(P.\) pastoris. Electrocompetent \(Pichia\) cells were prepared and transformed following the condensed protocol of Lin-Cereghino et al. (2005). For selection, Zeocin was used in a final concentration of 100 \(\mu\)g mL\(^{-1}\). The resulting colonies were picked and grown in 96-well deep well plates.

Microscale cultivation in deep well plates

Microscale cultivation and expression in 96-well deep well plates was done according to Weis et al. (2004) with some

### Table 1 Oligonucleotide primer sequences (restriction sites are underlined)

| Name         | Sequence 5′-3′ |
|--------------|---------------|
| 5pdhN-Nde    | TGCAATATGGCTATCACGTACCAACACC |
| 3pdhN-Hind   | CGAATAAGCTTATTAACCTTTTGCAAGATGCTGGTTCGAGTGACC |
| 5pdhN-Asc    | TCAAGGCGCGCCCGGAACACGTGAACGTCGTCGAGTGACC |
| 3pdhN-Not    | AGATGCCGCGCGTTATTTAACTCTTGTATCAACG |
| 5pdhS-Eco    | ATTCGAGAATTCGAGAAGCTATCTATCAACG |
| 3pdhS-Not    | AGATGCCGCGCGTTATATTG |
| 5alpha-Eco   | AAAGAATTCGAAGACGTGATAGATCCC |
| 5alpha-Asc   | AAAAGGCAGCGCGGAAAGACGTAGATCCC |
| 3alpha       | AGCTTCGGCTCTCTCTTCTCG |
| 5pdhN-alpha  | CGGAGAAGAGAGGCGGAAAGCTATCAACG |
| 5pdhS-alpha  | CGGAGAAGAGAGGCGGAAAGCTATCAACG |

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modifications. Cells were grown in 250 μL BMD1 (1.34% yeast nitrogen base, 4×10⁻⁵% biotin, 1% glucose, 200 mM potassium phosphate, pH 6.0) at 28 °C, 320 rpm, and 80% humidity for approximately 60 h to reach the stationary growth phase. BMM2 medium (250 μL; 1.34% yeast nitrogen base, 4×10⁻⁵% biotin, 1% methanol, and 200 mM potassium phosphate, pH 6.0) was added so that it gave a final concentration of 0.5% methanol. About 70, 82, and 108 h after the start of the cultivation, 50 μL BMM10 (BMM2 with 5% methanol) were added to keep up induction. After a total of 130 h, the cultivation was stopped. The deep-well plates were centrifuged at 4,000 rpm and 4 °C for 10 min and the supernatants were used for enzyme activity assays in microtiter plates.

Large-scale production of recPDH1 in P. pastoris

A fed-batch cultivation was carried out in a 60-L computer-controlled stirred tank reactor (Applikon, Schiedam, The Netherlands) with an initial volume of 40 L fermentation basal salts medium (21 mL L⁻¹ H₂PO₄ (85%); 0.93 g L⁻¹ CaSO₄·2H₂O; 14.9 g L⁻¹ MgSO₄·7H₂O; 18.2 g L⁻¹ K₂SO₄; 4.13 g L⁻¹ KOH; 4% (v/v) glycerol) supplemented with 4.35 mL L⁻¹ PTM1 trace salts (Invitrogen) and 10 mL Antifoam 204 (Sigma, St. Louis, MO, USA).

Temperature, dissolved oxygen (DO), and pH were monitored online by the Bio Controller ADI 1030 (Applikon). Oxygen concentration was regulated to a minimum of 4% and kept constant by supplying steril-filtered air and adjusting stirrer velocity. Temperature was set to 30 °C and the pH value was regulated to 5 and kept constant by addition of ammonium hydroxide solution. Fifteen 1-L baffled shaking flasks containing 300 mL YPD-Zeocin medium each (10 g L⁻¹ yeast extract, 20 g L⁻¹ peptone from casein, 4 g L⁻¹ glucose, and 25 mg L⁻¹ Zeocin) were inoculated with P. pastoris strain n-PDH_natCC11 and grown overnight at 30 °C and 150 rpm. This preculture was used for inoculation of the fermenter (10% v/v).

When the initially added glycerol was consumed completely (indicated by a DO spike after 13.5 h), a glycerol feed (50% glycerol supplemented with 12 mL L⁻¹ PTM1 trace salts) was started. Recombinant protein production was induced after 40 h of fermentation by feeding methanol (100% methanol supplemented with 12 mL L⁻¹ PTM1 trace salts). The feed rate was adjusted at constant air flow rate and stirrer speed to a DO of 4%. After 180 h of cultivation, the fermenter was harvested, cells were separated from the culture medium by centrifugation at 4,000×g, 4 °C for 10 min (Sorvall Evolution RC, Kendro Laboratory Products, Newtown, CT, USA) and the supernatant was recovered as a crude enzyme preparation.

Samples of the fermentation broth were taken periodically; wet biomass was determined by centrifugation (Eppendorf Centrifuge 5424, Hamburg, Germany) for 10 min at 14,000×g and weighing of the pellet. PDH activity was measured with the standard assay and soluble protein concentration was determined according to Bradford.

Purification of recombinant PDH1

recPDH1 was purified at room temperature by hydrophobic interaction chromatography and anion exchange chromatography. (NH₄)₂SO₄ was added to the crude extract to 30% saturation, clarified by centrifugation and the solution was loaded onto a Phenyl-Sepharose Fast Flow column (8,500 mL; flow rate 330 mL min⁻¹) equilibrated with 50 mM phosphate buffer (pH 6.5, 1.5 M (NH₄)₂SO₄). After washing the column with 1 column volume (CV) of the same buffer, recPDH1 was eluted with a linear gradient of starting buffer to 50 mM phosphate buffer (pH 6.5) in 1 CV and fractions were collected.

Fractions with high PDH activity were concentrated and desalted to a conductivity equal to or less than 2 mS cm⁻¹ using cross-flow filtration (Microza ultrafiltration module, Pall Austria Filter GmbH, Vienna, Austria). A pool of the fractions containing highest activity was loaded onto a DEAE Sepharose Fast Flow anion exchange chromatography column (500 mL) equilibrated with 50 mM Bis Tris buffer (pH 6) and washed with 1 CV of this buffer. For elution, a linear salt gradient (0–1 M NaCl in 50 mM Bis Tris, pH 6) was run in 10 CV at 10 mL min⁻¹.

### Table 2

| Construct          | Leader | Gene  | Number of screened transformants | Maximum activity (U mL⁻¹) |
|--------------------|--------|-------|----------------------------------|--------------------------|
| α-PDH Nat          | α-Factor | Native | 176                              | nd                       |
| n-PDH Nat          | Native  | Native | 352                              | 0.57                     |
| α-PDH Syn          | α-Factor | Synthetic | 84                              | nd                       |
| n-PDH Syn          | Native  | Synthetic | 252                              | 0.36                     |

*nd* Not detectable
Fractions with the highest activities were pooled, concentrated (Vivaflow 50 module, Sartorius AG, Göttingen, Germany), aliquoted, and stored at −30 °C.

Enzyme assay

Pyranose dehydrogenase activity was determined by following spectrophotometrically the D-glucose-dependent reduction of the ferrocenium ion (Fc⁺) to ferrocene at 300 nm (ε₃₀₀=4.3 mM⁻¹ cm⁻¹) for 3 min at 30 °C (Kujawa et al. 2007). One unit of PDH activity was defined as the amount of enzyme required for the reduction of 2 μmol of Fc⁺ ion per minute.

Molecular properties

The protein concentration was determined by the method of Bradford using a prefabricated assay from Bio-Rad Laboratories (Hercules, CA, USA) and bovine serum albumin as the standard. SDS-PAGE was carried out using Mini-PROTEAN TGX precast gels with a denaturing gradient of 4–20%. Protein bands were visualized by staining with Bio-Safe Coomassie; Dual Color Precision Plus Protein Standard was used for mass determination. All procedures were done according to the manufacturer’s recommendations (Bio-Rad).

Enzymatic deglycosylation

Deglycosylation of a 15-μg sample of PDH was performed with peptide/N-glycosidase F, which cleaves the bond between N-acetyl-glucosamine. All procedures of the enzymatic deglycosylation of PDH with PNGaseF under denaturing conditions were done according to the manufacturer’s recommendations.

Kinetic properties

Apparent kinetic constants for D-glucose, D-maltose, L-arabinose, and lactose were measured using the standard ferrocenium assay as described earlier. Kinetic constants for the electron acceptors 1,4-benzoquinone, 2-chloro-1,4-benzoquinone, and Fc⁺ hexafluorophosphate were determined with 25 mM glucose as electron donor. The pH dependence of PDH activity with different electron acceptors was determined with the following buffers: 100 mM citrate (pH 2.5–6.0), 100 mM phosphate (pH 6.0–8.0), and 100 mM borate (pH 8.0–9.0). All kinetic constants were calculated by nonlinear least-squares regression, fitting the observed data to the Henri–Michaelis–Menten equation. Turnover numbers (kcat) and catalytic efficiencies (kcat/Km) were calculated using the molecular mass.

Results

Expression of pdh1 in E. coli

The E. coli expression plasmid pET21apdh1 was constructed by cloning the nucleotide sequence of mature PDH1 into the pET21a expression vector under control of the T7 promoter. Transformed E. coli BL21 (DE3) cultures carrying the plasmid were grown under different inducing conditions. Cultures were harvested 16 h after induction with 0.01 and 0.1 mM IPTG as well as with 0.5% lactose, reaching an OD₆₀₀ of roughly 11. Harvested cells were resuspended in lysis buffer (three times the weight of the wet biomass of the sample) and disrupted using a French press. Disruption efficiencies were compared by measuring total protein concentration of the cleared lysate and varied between 1.1 and 2.0 mg/mL (corresponding to 4.4–8.0 mg of protein per gram of cell paste). Crude extracts were tested for PDH activity using the standard enzyme assay. Unspecific background of a crude extract from an equally treated culture of E. coli BL21 (DE3) cells carrying the empty pET21a vector was subtracted from the measurements. No significant PDH activity was measured in any of the cultures. SDS-PAGE analysis of the total soluble proteins of the crude extract showed no band at the expected size (62 kDa) of PDH (data not shown). The insoluble part of the crude extract was tested for the possible existence of inclusion bodies of PDH on an SDS-PAGE gel (data not shown). The majority of insoluble proteins were roughly the size of purified native PDH.

Expression of pdh1 in P. pastoris

Several constructs for extracellular expression of pdh1 in P. pastoris were designed, all of them under control of

![Fig. 1](https://example.com/fig1.png) Large-scale fermentation of P. pastoris n-PDH natCC11. Circles wet biomass, squares specific activity, diamonds volumetric activity. The measurements were done in duplicates and the error was always less than 5%
the inducible AOX1 promoter. For extracellular expression with the native signal sequence, the full-length open reading frames of \textit{pdh1} and the gene sequence optimized for expression in \textit{P. pastoris} (\textit{synpdh1}) were cloned into the vector pPpT2 (Abad et al. 2010; Ruthe et al. 2010). To test the efficiency of the native leader sequence, it was exchanged to a codon optimized α-factor pre–pro leader peptide of \textit{S. cerevisiae} by overlap extension PCR.

All resulting constructs were transformed into \textit{P. pastoris} strain CBS 7435, cultivated in 96-well deep well plates, and screening for PDH activity (Table 2). On day 6 of the cultivation with approximately 72 h of induction by methanol, supernatants were tested for PDH activity. Constructs with the α-factor leader sequence showed no significant PDH activity. In contrast, constructs with the native leader sequence resulted in \textit{P. pastoris} transformants secreting active recombinant PDH1 (\textit{recPDH1}).

Overall, no significant difference in average activity could be observed when comparing the expression levels of all tested native with synthetic gene constructs. The standard deviation within one group of constructs was always approximately 10%, which shows the uniformity of the 96-well deep well plate screening and proves its suitability for high-throughput screening. To confirm the results from the first round of screening, a rescreening experiment with multiple parallel determinations was performed. The three synthetic gene transformants selected for the rescreen yielded an average volumetric PDH activity of around 0.31 U mL\(^{-1}\). Three native gene transformants showed a slightly lower average PDH activity of 0.28 U mL\(^{-1}\) but clone n-PDH\_natCC11 had a volumetric activity of 0.47 U mL\(^{-1}\). This clone was selected for large-scale fermentation.

Scale up of \textit{recPDH1} production

\textit{P. pastoris} clone n-PDH\_natCC11, expressing extracellular \textit{recPDH1}, was cultivated on a 50-L scale in a fed-batch fermentation for a total of 181 h (Fig. 1). The initial glycerol batch phase lasted for 13.5 h and its end, i.e., the exhaustion of the carbon source, was indicated by a sharp increase of the dissolved oxygen level. At this time point, a glycerol feed was started to further increase the cell density. After 40 h of cultivation, when the wet biomass had reached 111 g L\(^{-1}\), the glycerol feed was stopped and induction with methanol started by continuously supplying this inducer. Levels of wet biomass reached 159 g L\(^{-1}\) during the induction phase and the concentration of soluble protein in the culture supernatant increased from 70 to 440 mg L\(^{-1}\). Volumetric PDH activity in the culture supernatant reached a value of 10,040 U L\(^{-1}\) at the time of harvest (corresponding to ~230 mg of \textit{recPDH1} per liter) with a specific activity of 22.9 U mg\(^{-1}\) total protein.

Purification of recombinant PDH1

\textit{recPDH1} was purified by a two-step chromatographic procedure as outlined in Table 3. Hydrophobic interaction chromatography was followed by an anion exchange chromatography step resulting in two pools containing active PDH. Pool 1 represented the center of the elution

| Table 3 Purification of recombinant PDH1 from \textit{A. meleagris} | Total activity (U) | Total protein (g) | Volume (L) | Specific activity (U mg\(^{-1}\)) | Purification (-fold) | Activity yield (%) |
|---------------------------------------------------------------|-------------------|-----------------|----------|-------------------------------|-----------------|---------------|
| Culture supernatant                                           | 517,000           | 22.6            | 51.5     | 22.9                          | 1.0             | 100           |
| Phenyl Sepharose FF                                           | 675,000\(^a\)     | 19              | 15.4     | 35.5                          | 1.55            | 130.6\(^a\)  |
| DEAE Sepharose FF pool 1\(^b\)                               | 48,100            | 1.1             | 0.98     | 43.7                          | 1.90            | 10.7          |
| DEAE Sepharose FF pool 2                                     | 440,000           | 12.3            | 1        | 35.8                          | 1.57            | 85.1          |

\(^a\) PDH activity is higher when ammonium sulfate is present  
\(^b\) Determination of kinetic constants was done with protein from this fraction
peak and contained a homogenous enzyme preparation as judged by SDS-PAGE. Only 10% of the starting activity could be found in this amber-colored enzyme preparation with a specific activity of 43.6 U mg\(^{-1}\) representing a 1.9-fold purification from the culture supernatant. Protein from this pool was used for all further analyses. Since the emphasis of this purification procedure was put on high purification yields, aiming at possible subsequent applications of the enzyme, all other fractions containing PDH with relatively high specific activity were combined in pool 2 resulting in a recovery of 85% with a specific activity of 35.9 U mg\(^{-1}\). In total, 13.4 g of more than 80% pure recPDH1 were obtained.

Catalytic properties

The pH/activity profiles of recPDH1 were measured for the electron acceptors 1,4-benzoquinone, Fc\(^+\) hexafluorophosphate, and 2-chloro-1,4-benzoquinone, using glucose as electron donor (Fig. 2). recPDH1 showed a virtually identical pH behavior compared to the wild-type protein with benzoquinones having their pH optima in the acidic range whereas activity with Fc\(^+\) continually increased with pH in the range of pH 3–9. Apparent kinetic constants for these electron acceptor substrates were determined at those pH values that were previously used for wild-type PDH1 (Table 4). In addition, apparent kinetic constants were determined for selected sugar substrates and compared to those of the wild-type enzyme from \textit{A. meleagris} (Table 5). Catalytic properties of recPDH1 expressed in \textit{P. pastoris} are almost identical to those of the wild-type enzyme (Sygmund et al. 2008).

Molecular properties

The molecular mass of recombinant PDH1 overexpressed in \textit{P. pastoris} was determined to be ~93 kDa by SDS-PAGE (Fig. 3). Deglycosylation under denaturing conditions using PNGase F reduced the mass of the protein to 64 kDa. This mass corresponds very well to the theoretical molecular mass of 61,967 Da based on the cDNA sequence of mature PDH1. Based on these results, the level of glycosylation of recPDH1 was calculated to be approximately 30%.

Discussion

Production of PDH activity in its natural source, \textit{A. meleagris}, is extremely tedious and time consuming as these cultivations have to be performed in resting shallow cultures for up to 6 weeks. Recently, the PDH1-encoding gene from \textit{A. meleagris} was successfully expressed in

### Table 4 Apparent kinetic constants of recPDH1 from \textit{A. meleagris} for selected electron acceptor substrates

| Substrate                  | Recombinant PDH1 | Native AmPDH\(^a\) |
|----------------------------|------------------|---------------------|
|                            | \(k_{cat}\) (s\(^{-1}\)) | \(K_m\) (mM) | \(k_{cat}/K_m\) (mM\(^{-1}\) s\(^{-1}\)) | \(k_{cat}\) (s\(^{-1}\)) | \(K_m\) (mM) | \(k_{cat}/K_m\) (mM\(^{-1}\) s\(^{-1}\)) |
| 1,4-Benzooquinone          | 4.0              | 65.4±5.5           | 1.38±0.28 | 54.4             | 76.0±2.3\(^b\) | 1.82±0.11\(^b\) | 57.5\(^b\) |
| 2-Chloro-1,4-benzoquinone  | 5.0              | 12.9±1.2           | 0.62±0.11 | 20.9             | 15.1±1.0       | 0.55±0.11       | 25.5       |
| Ferrocenium hexafluorophosphate | 8.5              | 130±11             | 0.16±0.04 | 812              | 104±8          | 0.13±0.03       | 802       |

Kinetic data were determined at 30 \(^\circ\)C with 25 mM D-glucose as the electron donor
\(^a\) Data from Sygmund et al. (2008)
\(^b\) Measurements were done at pH 3.0

### Table 5 Apparent kinetic constants of recPDH1 from \textit{A. meleagris} for selected electron donor substrates

| Substrate | Recombinant PDH1 | Native AmPDH\(^a\) |
|-----------|------------------|---------------------|
|           | \(k_{cat}\) (s\(^{-1}\)) | \(K_m\) (mM) | \(k_{cat}/K_m\) (mM\(^{-1}\) s\(^{-1}\)) | \(k_{cat}\) (s\(^{-1}\)) | \(K_m\) (mM) | \(k_{cat}/K_m\) (mM\(^{-1}\) s\(^{-1}\)) |
| D-Glucose | 37.8±1.1         | 0.69±0.09          | 54.8  | 45.9±0.3       | 0.82±0.03       | 57.5   |
| D-Galactose | 47.3±3.0        | 1.07±0.13          | 44.2  | 48.5±0.6       | 1.05±0.05       | 46.2   |
| D-Xylose   | 62.4±1.6         | 1.92±0.14          | 32.5  | 43.4±0.8       | 1.93±0.17       | 22.9   |
| L-Arabinose | 34.7±1.6        | 0.49±0.05          | 70.8  | 37.2±0.6       | 0.54±0.08       | 62.1   |
| Lactose    | 41.0±8.1         | 128±11.9           | 0.32  | 39.6±0.8       | 134±6.3        | 0.29    |

Kinetic data were determined at 30 \(^\circ\)C with 0.4 mM ferricenium as the electron acceptor
\(^a\) Data from Sygmund et al. (2008)
was stained with Coomassie blue. The gel was stained with Coomassie blue.

**Fig. 3** SDS PAGE analysis of glycosylated and deglycosylated recombinant pyranose dehydrogenase expressed in *P. pastoris* and PDH1 purified from *A. meleagris*. 1 PDH1 from *A. meleagris*, 2 deglycosylated PDH1 from *A. meleagris*, 3 molecular mass marker (Precision Plus Protein Standards, Bio-Rad), 4 recPDH1 expressed in *P. pastoris* (purification pool 1), 5 deglycosylated recPDH1 expressed in *P. pastoris* (purification pool 1); 6 molecular mass marker.

Aspergillus nidulans and Aspergillus niger. By using this approach, the final volumetric activity as well as the space–time yield could be increased significantly with *A. nidulans* as expression host (Pisanelli et al. 2010) compared to production by the natural producer. However, time-consuming transformation protocols and complicated genetic manipulation of *Aspergillus* spp. severely hamper any protein-engineering attempts. Therefore, other microbial expression systems that are better amenable to protein engineering approaches were tested for their ability to produce soluble and active PDH1. Both the prokaryotic host *E. coli* as well as the eukaryotic host *P. pastoris* were selected for expression studies.

The *E. coli* expression construct was designed for expression of PDH1 without N-terminal signal peptide. To reduce the possible formation of insoluble recombinant protein cultivation, temperature was lowered to 25 °C and the concentration of the inducer IPTG was varied. Still, PDH1 could only be found as insoluble protein fraction without any enzymatic activity under all tested conditions. One possible reason for the failure of expressing PDH in a prokaryotic system could be the need of glycosylation for correct protein folding as observed previously with other enzymes (Weis et al. 2007). On the other hand, there are examples where random mutagenesis was used to achieve functional expression of fungal proteins in *E. coli* (Sun et al. 2001). A similar strategy could possibly be used for successful expression of PDH1 in *E. coli*. Another key factor that seems to have major impact on PDH expression is the 25-amino acid long signal peptide. As reported earlier, expression levels of PDH1 in *A. nidulans* were much higher compared to the glaA signal sequence when its native signal sequence was used (Pisanelli et al. 2010). The same effect was observed in *P. pastoris*, and here it was even more pronounced. No PDH activity could be detected when using the α-factor pre–pro signal sequence although identical constructs containing the native signal sequence showed relatively high expression levels. A similar effect, although less pronounced, was observed for the expression of other proteins in *P. pastoris* before (Colao et al. 2006). A possible reason could be the formation of secondary structures in the mRNA when the α-factor is fused to the *pdh1* gene. However, analysis of the mRNA sequences of the four tested constructs using the program GeneBee (Brodsky et al. 1995) showed that constructs containing the α-factor leader sequence neither have a lower calculated total free energy nor do they form stronger stem–loop structures compared to the sequences with the native signal sequence. The only significant difference with regard to stem–loop formation was between the synthetic gene and the noncodon-optimized sequence. This effect can be explained by the removal of regions capable of forming secondary structures during gene optimization.

A second possible explanation for the failure of expressing PDH1 with the α-factor could be inefficient processing of this signal sequence in combination with the PDH protein sequence. Three proteolytic cleavage steps are necessary to fully process the α-factor pre–pro leader sequence, and especially the efficiency of the last step can be influenced by the amino acids surrounding the cleavage site (Goda et al. 2000; Almeida et al. 2001). Isoleucine and tyrosine at position 2 and 4 of the mature protein can cause steric hindrance to the access of proteases, and thus affect processing of the protein by the Kex2 or the Ste13 protease. Another possible cause for inefficient processing of the protease cleavage sites could be glycosylation close to the *N*-terminus of the protein (Weiss et al. 1998). One possibility to solve this problem is to remove the Glu-Ala repeats by cloning the target gene flush with the Kex2 cleavage site. Another approach could be random mutagenesis in the entire secretion leader, which has been recently reported as a suitable strategy for the functional expression of a foreign laccase in yeast (Maté et al. 2010). Expression levels of both constructs with the native signal sequence in 96-well deep well cultivations were similar and in the range of 0.3 U mL⁻¹. Because of this relatively high and uniform expression level the, 96-well deep well cultivations can be used to screen for mutated enzymes produced by semirational design approaches. The use of the optimized synthetic gene did not increase the expression level; indeed, the highest volumetric activity was achieved by a transformant, n-PDH_natCC11, with native *pdh1*.
sequence. A large-scale laboratory fermentation of clone n-PDH\_natCC11 was performed to obtain large amounts of homogenous protein for protein crystallography and fast kinetic studies as well as to demonstrate the feasibility of industrial scale production of PDH for biotechnological applications. The 50-L fermentation resulted in a volumetric activity of 10,040 UL\(^{-1}\) or 1,330 UL\(^{-1}\) d\(^{-1}\) in space–time yield. This is a 28-fold improvement of the space–time yield compared to the wild-type producer and a 1.6-fold improvement compared to \textit{A. nidulans} expression yields (Pisanelli et al. 2010). The cultivation yielded 223 mg L\(^{-1}\) of recombinant protein, which corresponds to 52% of total extracellular protein. Because of the high specific activity (22.9 Umg\(^{-1}\)) of PDH in the culture supernatant, the previously established purification protocol (Sygmund et al. 2008) was changed to a two-step procedure with focus on fast high-yield purification. This represents another major advantage over expression in \textit{Aspergillus} spp. where the presence of a large number of secreted proteins considerably reduces purification yields. The purification resulted in an apparently pure protein preparation with a specific activity of 43.7 Umg\(^{-1}\) that was used for the subsequent biochemical characterization of the enzyme. To show that recombinant PDH1 can be purified in very high yields with this simple purification protocol for biotechnological applications, where a purity of the target protein of 80% is sufficient, other fractions with high specific activity were pooled resulting in a recovery of 85% of the initial activity and 12.3 g of PDH1. For applications where higher purity is required, the activity yield of 10.7% of pool 1 might be too low and then an additional purification step would be necessary. The specific activity of recombinant PDH1 obtained with \textit{P. pastoris} is about 20% lower compared to wild-type PDH1. A possible reason for the reduced specific activity might be the higher degree of glycosylation of recPDH1 expressed in \textit{P. pastoris} (30% compared to 7%), or inactive, not correctly folded protein that could not be separated from the active form. In contrast to \textit{S. cerevisiae}, hyperglycosylation is not common for proteins expressed in \textit{P. pastoris}. The high degree of glycosylation of PDH1 is a hint that the transit from the endoplasmic reticulum to the Golgi apparatus might be a bottleneck for the expression. The kinetic properties are essentially identical to those measured for the wild-type enzyme isolated from its natural source \textit{A. meleagris}.

This study demonstrates the successful expression of PDH in the eukaryotic host organism \textit{P. pastoris} and thus offers the possibility to easily produce and screen for genetically engineered PDH variants paving the way for future protein engineering. Additionally the feasibility of large-scale production of the enzyme using this expression system together with a simplified purification scheme for easy high-yield purification is shown.

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