Directed Evolution of a Thermostable Phosphite Dehydrogenase for NAD(P)H Regeneration

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NAD(P)H-dependent oxidoreductases are valuable tools for synthesis of chiral compounds. The expense of the cofactors, however, requires in situ cofactor regeneration for preparative applications. We have attempted to develop an enzymatic system based on phosphite dehydrogenase (PTDH) from Pseudomonas stutzeri to regenerate the reduced nicotinamide cofactors NADH and NADPH. Here we report the use of directed evolution to address one of the main limitations with the wild-type PTDH enzyme, its low stability. After three rounds of random mutagenesis and high-throughput screening, 12 thermostabilizing amino acid substitutions were identified. These 12 mutations were combined by site-directed mutagenesis, resulting in a mutant whose $T_{1/2}$ is 20°C higher and half-life of thermal inactivation at 45°C is >7,000-fold greater than that of the parent PTDH. The engineered PTDH has a half-life at 50°C that is 2.4-fold greater than the Candida boidinii formate dehydrogenase, an enzyme widely used for NADH regeneration. In addition, its catalytic efficiency is slightly higher than that of the parent PTDH. Various mechanisms of thermostabilization were identified using molecular modeling. The improved stability and effectiveness of the final mutant were shown using the industrially important bioconversion of trimethylpyruvate to t-tert-leucine. The engineered PTDH will be useful in NAD(P)H regeneration for industrial biocatalysis.

The potential of enzyme-catalyzed transformations in the pharmaceutical and fine-chemical industries has long been considered to have great promise (7, 10, 23). Despite the development of rapid and efficient methods for creating designer biocatalysts, inherent limitations in the chemistry that these enzymes perform still persist. The majority of biocatalysts used in industry are hydrolytic in action (~65%) and perform rather simple chemistry (4). More complicated enzymatic reactions often require one or more costly cofactors, which, when added in stoichiometric quantities, make the process not economically feasible. Oxidoreductases, for example, catalyze a myriad of regio-, chemo-, and stereospecific reactions on a variety of functional groups, but often require either NADH or NADPH as a cofactor (8, 11).

Various in situ regeneration methods have been developed to allow the use of catalytic quantities of NAD(P)⁺ and NAD(P)H (20). The most successful and widely used enzymatic NADH regeneration system is based on the Candida boidinii formate dehydrogenase (FDH) (1). We have attempted to develop an NADH regeneration process using the enzyme phosphite dehydrogenase (PTDH) from Pseudomonas stutzeri (2). This enzyme catalyzes the nearly irreversible oxidation of phosphite to phosphate with the concomitant reduction of NAD⁺ to NADH. The kinetic and practical advantages of using PTDH for NADH regeneration have been recently reported (22).

In addition to NADH regeneration, the enzyme PTDH also has considerable potential in regenerating NADPH. The cost of NADPH is significantly higher than that of NADH, and no widely used NADPH regeneration system is currently available. To date, the most useful enzyme for NADPH regeneration is a mutant Pseudomonas sp. strain 101 FDH available from Jülich Fine Chemicals (18). Although wild-type PTDH has a preference for NAD⁺ over NADP⁺ by about 100-fold, a mutant PTDH has been created by rational design with relaxed specificity toward both nicotinamide cofactors (26). The catalytic efficiency of the PTDH double mutant with NADP⁺ ($k_{cat}/K_m$NADP⁺) is 33-fold higher than that of Pseudomonas sp. strain 101 FDH with a comparable turnover number.

As a biocatalyst for industrial applications, one of the main limitations of the wild-type PTDH is its low stability. Indeed, the wild-type enzyme shows fairly rapid inactivation under relatively mild temperatures (2). Thus, we sought to use directed evolution to improve its thermostability. Directed evolution has become a powerful tool for improving enzyme function and has been successfully used to improve the thermal stability of many different enzymes (9, 15, 19, 27). Here we report the directed evolution of a highly thermostable PTDH mutant for NAD(P)H regeneration. Three rounds of error-prone PCR and high-throughput screening coupled with site-directed mutagenesis yielded a PTDH mutant with higher thermostability than the most thermostable Candida boidinii FDH reported previously (17).

MATERIALS AND METHODS

Reagents. Escherichia coli BW25141 and plasmid pLA2 were kindly provided by William Metcalf at the University of Illinois (Urbana, IL) (6). The pRW2 plasmid was created from pLA2 as described elsewhere (26). Escherichia coli BL21(DE3) and plasmid pET15b were purchased from Novagen (Madison, WI). Taq DNA polymerase was obtained from Promega (Madison, WI) and cloned Pfu Turbo DNA polymerase was obtained from Stratagene (La Jolla, CA). The DNA-modifying enzymes NdeI, BamHI, PciI, and T4 DNA ligase were pur-
chased from New England Biolabs (Beverly, MA). PCR-grade deoxynucleotide triphosphates and DNAse I were obtained from Roche Applied Sciences (Indi-
anapolis, IN). l-Aspartic acid was purchased from Fluka (St. Louis, MO). Ampicillin, kanamycin, lysozyme, nitro blue tetrazolium (NBT), phenazine methosulfate, and NAD+ were purchased from Sigma (St. Louis, MO). Phos-
phoroc acid was obtained from Aldrich (Milwaukee, WI). Other salts and reagents were purchased from either Sigma-Aldrich or Fisher Scientific (Pitts-
burgh, PA). Oligonucleotide primers were obtained from Integrated DNA Tech-
nologies (Coralville, IA). The QiAquick spin plasmid mini-prep kit, QIAEX II gel purification kit, and QiAquick PCR purification kit were purchased from QIAGEN (Valencia, CA). BD Talon metal affinity resin was purchased from BD Biosciences Clontech (Palo Alto, CA). Amicon Ultra-15 centrifugal filter devices were purchased from Fisher Scientific (Pittsburgh, PA). Purified formate dehydrogenase (FDH) from Candida boidinii was purchased from Jülich Fine Chemicals (Jülich, Germany).

Random mutagenesis and library creation. Random mutagenesis was carried out by error-prone PCR as described elsewhere (29). Plasmid pH2 containing the parent gene was used as the template for the first generation for random mutagenesis. For the 1.0- kb PTDH parent target gene, 0.2 mM MnCl2 was used to obtain the desired level of mutagenesis rate (~1 to 2 amino acid substitutions). Forward 5′-TTTTGGATCCGGGAATCTCAGATGCTTTG-3′ and reverse 5′-CCGAGAAGGCGTTACCGGTATCATC-3′ primers were used to amplify the gene. PCR-mutated genes were digested with NdeI and PciI and ligated into the plasmid pH2 digested with the same two restriction enzymes. Ligation reactions (10 μl total volume) contained ~100 ng inserts, ~100 ng vector, 1X T4 DNA ligase buffer, and 0.5 U T4 DNA ligase, and were incubated at 16°C for 16 h. The resulting plasmids were transformed into newly prepared electrocom-
potent E. coli WM1788 cells, which were then plated on Luria-Bertani (LB) agar plates containing ampicillin and kanamycin.

Thermostability screening. For identification of thermostable PTDH variants, a 96-well plate thermostability screening assay was developed based on a colori-
metric NBT-phenazine methosulfate assay (14). Library colonies were picked with sterile toothpicks and used to inoculate 100 μl of LB medium containing 50 μg/ml kanamycin in 96-well plates. The plates were incubated at 37°C for 5 h, and protein expression was induced with arabinose (final concentration ~10 mM) before assaying. Typically, inactivation was monitored until >80% of the activity was lost. Half-lives of thermal inactivation were calculated using the equation \[ T_{1/2} = \frac{\ln(2)}{k_{\text{inact}}} \]
where \( k_{\text{inact}} \) is the inactivation rate constant obtained from the slope by plotting log (residual activity/initial activity) versus time.

Thermostable phosphite dehydrogenase. Thermostable phosphite dehydrogenase (PTDH) overexpression and purification. Protein purification of the parent and mutant PT DHs were carried out using a protocol described elsewhere (26) with some modifications. Small-scale spin columns containing approximately 0.5 ml of BD Talon resin were used to purify multiple enzymes in parallel. The columns were equilibrated in start buffer A (0.5 M NaCl, 20% glycerol, and 20 mM Tris-HCl, pH 7.6). The following method was used to purify PTDH (with His, tag) ~10 ml of clarified supernatant (~1 to 5 g of cell paste) loaded onto the column, column washed 10 times with start buffer A, column washed five times with start buffer A plus 10 mM imidazole, and proteins eluted with elution buffer (0.5 M imidazole, 0.5 M NaCl, 20% glycerol, and 20 mM Tris-HCl, pH 7.6). Elution fractions were collected and each fraction was checked for activity using NBT-phenazine methosulfate assay. Pooled active fractions were concen-
trated using an Amicon Ultra-15 filter device and washed four times with 20 ml of 50 mM morpholinepropanesulfonic acid (MOPS) buffer (pH 7.25). The en-
zyme was then stored at a concentration of >2 mg/ml in 20% glycerol at ~80°C.

Enzyme kinetics. The kinetic rate constants for the parent and mutant PT DHs were determined as described elsewhere (26). Enzyme concentration was deter-
minded by measuring A280 (ε = 30 mM−1 cm−1). The data represent an average of all statistically relevant data with a standard deviation of less than 10%.

Half-lives of thermal inactivation. Purified enzymes (0.2 mg/ml) were incu-
bated in an MJ Research PTC-200 thermocycler (±0.3°C) to study enzyme inactivation. PTDH enzymes were incubated in 50 mM MOPS buffer (pH 7.25), whereas Candida boidinii FDH was incubated in 100 mM potassium phosphate buffer (pH 7.5). Aliquots were taken at specific time points and placed on ice before assay. Typically, inactivation was monitored until >80% of the activity was lost. Half-lives of thermal inactivation were calculated using the equation \[ T_{1/2} = \frac{\ln(2)}{k_{\text{inact}}} \]
where \( k_{\text{inact}} \) is the inactivation rate constant obtained from the slope by plotting log (residual activity/initial activity) versus time.

 Batch production of L-seryl-leucine. Small-scale reactions were conducted to demonstrate the improved stability and effectiveness of the 12x thermostable PTDH mutant. The final volume of each batch reaction was 300 μl. Each sample contained 300 mM ammonium trimethylpyruvate, 400 mM diammonium phos-
phate (400 mM ammonium formate in the case of FDH), 0.4 mM NAD+, 5.26 U/ml Bacillus cereus leucine dehydrogenase, and 100 μg of PT DH or FDH. Each reaction was mixed gently and incubated at 40°C. At fixed time intervals, small aliquots were removed from the reaction and immediately frozen at −20°C. The concentration of L-seryl-leucine was monitored using a high-pressure liquid chro-
matograph (Shimadzu) equipped with an evaporative light-scattering detector. Substrates and products were separated on an Alltech C-18 Penta column with an isocratic elution of 94% water, 4.5% acetonitrile, and 1% acetic acid. The concentration of each sample was determined by using a standard curve prepared with authentic L-seryl-leucine.

RESULTS

Generation of thermostable PT DH mutants. The thermo-
stability of PTDH was improved by combining error-prone PCR and site-directed mutagenesis approaches. Error-prone PCR was used to introduce an average of 1 to 2 amino acid substitutions per PT DH gene. This low mutation rate made it possible to identify the contribution of individual amino acid substitutions on enzyme thermostability. Three rounds of er-
ror-prone PCR mutagenesis were used to generate libraries of PT DH variants. After each round of error-prone PCR, site-
directed mutagenesis was used to combine thermostabilizing mutations into one template and the resulting template served
as the parent for the next round of error-prone PCR mutagenesis.

The first-generation mutant library was constructed using a mutant PTDH that we recently engineered (25). This PTDH mutant differs from wild-type PTDH at five amino acid positions (D13E, M26I, E175A, E332N, and C336D). These mutations improved the enzyme solubility and activity, but showed little effect on the enzyme thermostability. At 45°C, the \( t_{1/2} \) of wild-type PTDH is 1.4 \( /H_11006 \) 0.3 min, whereas the \( t_{1/2} \) of the parent is 1.2 \( /H_11006 \) 0.2 min. After creating the first-generation mutant library using error-prone PCR, approximately 3,200 clones were screened for increased thermostability at 43°C. Five variants were identified and confirmed to have longer half-lives than the parent at 45°C (Table 1). Sequencing these variants revealed the following amino acid substitutions: 1-1A9 (Q137R), 1-8D6 (R275Q), 1-11A6 (Q215L), 1-23C7 (Q132R), and 1-25E8 (I150F) (Fig. 1). Site-directed mutagenesis was used to combine these thermostabilizing mutations onto the parental template to create 4x (Q137R, I150F, Q215L, R275Q) and 5x (Q132R, Q137R, I150F, Q215L, R275Q) mutants.

The gene encoding the 4x mutant was chosen as a template for generating the second-generation mutant library using error-prone PCR (Table 2). The 4x mutant was chosen over the 5x mutant as the template for second-generation random mutagenesis based on a higher catalytic activity and longer half-life at 45°C. Approximately 6,000 clones from the 4x library were screened for increased thermostability at 43°C. Three clones were found to have longer half-lives than the parent at 45°C. After sequencing these three variants, the following additional amino acid substitutions were identified: 2-2C10 (D162N, V315A), 2-9B6 (A319E), and 2-17C2 (L276Q) (Fig. 1). Clone 2-2C10 had two amino acid substitutions. Both mutations were incorporated individually onto the 4x template using site-directed mutagenesis to determine the effect of each mutation separately on enzyme thermostability. Single mutant V315A shows a threefold increase in thermostability compared to the 4x mutant, whereas single mutant D162N shows a 1.8-fold decrease in thermostability. Using site-directed mutagenesis, the substitutions L276Q, A319E, and V315A were combined on the 4x template to create a 7x mutant.

The 7x mutant was used as the parent for the third round of random mutagenesis. From this library, ~10,000 clones were screened for increased thermostability at 62°C. Five variants were identified and confirmed to have longer half-lives at 45°C than the 7x mutant. Sequencing these variants revealed the following changes: 3-34G8 (Q132R), 3-57E11 (V71I), 3-89D3 (E130K), 3-110G7 (I313L) and 3-135H2 (A325V) (Fig. 1). It is noteworthy that Q132R appeared twice, in the first and third-generations. This gave further evidence that Q132R is indeed a thermostabilizing mutation, and thus it was incorporated into the final template. Site-directed mutagenesis was used to incorporate all third round mutations into the 7x template to create the 12x mutant.

**Thermostabilities of the evolved PTDHs.** Table 1 lists the \( t_{1/2} \) and \( T_{50} \) values of the parent and the evolved thermostable PTDHs accumulated during three generations of directed evolution. At 45°C, the \( t_{1/2} \) of the parent is 1.2 min and its \( T_{50} \) is 39°C. The first-generation variants have half-lives of thermal

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**TABLE 1. Thermostability data for the parent and mutant PTDHs**

<table>
<thead>
<tr>
<th>Generation</th>
<th>Variant</th>
<th>( t_{1/2} ) (min, 45°C)</th>
<th>Fold Improvement</th>
<th>( T_{50} ) (°C)</th>
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<td>1</td>
<td>39.0</td>
</tr>
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<td>3.2</td>
<td>41.9</td>
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<td>1-8D6</td>
<td>4.6</td>
<td>3.8</td>
<td>40.7</td>
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<tr>
<td></td>
<td>1-11A6</td>
<td>8.7</td>
<td>7.3</td>
<td>42.5</td>
</tr>
<tr>
<td></td>
<td>1-23C7</td>
<td>2.3</td>
<td>1.9</td>
<td>40.0</td>
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<tr>
<td></td>
<td>1-25E8</td>
<td>7.0</td>
<td>5.8</td>
<td>42.2</td>
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<td>4x</td>
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<td>2-9B6</td>
<td>567</td>
<td>473</td>
<td>53.8</td>
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<td>2-17C2</td>
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<td>7x</td>
<td>1,421</td>
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<td>2,350</td>
<td>1,958</td>
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<td></td>
<td>3-89D3</td>
<td>2,000</td>
<td>1,667</td>
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<td></td>
<td>3-110G7</td>
<td>1,765</td>
<td>1,471</td>
<td>57.5</td>
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<tr>
<td></td>
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<td>2,315</td>
<td>1,929</td>
<td>57.5</td>
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<tr>
<td></td>
<td>12x</td>
<td>8,440</td>
<td>7,033</td>
<td>59.3</td>
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Average of all statistically relevant data (standard deviation <10%). Wild type PTDH has a \( t_{1/2} \) of 1.4 ± 0.3 min at 45°C. All experiments were performed in 50 mM MOPS (pH 7.25).

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**FIG. 1.** Lineage and protein-level mutations in the evolved thermostable PTDH variants. Corresponding DNA-level mutations are shown in parentheses. The substitution shown in italics represents a mutation that does not increase PTDH thermostability. EP-PCR, error-prone PCR; SDM, site-directed mutagenesis; *, previous directed evolution work to improve activity and solubility (25).
The evolved 12x mutant has similar \( k_{\text{cat}} \) that of the parent at 45°C and its \( T_{50} \) is 15.5°C higher. The half life of the 7x mutant is increased by 130 to 170 times that of the parent and their \( T_{50} \) are \( \sim 14°C \) higher.

A further round of random mutagenesis resulted in three variants that have half-lives of thermal inactivation 360 to 170 times longer than the parent at 45°C and their \( T_{50} \) are 14.8 to 15.5°C higher. The half life of the 7x mutant is \( \sim 1,200 \) times that of the parent at 45°C and its \( T_{50} \) is increased by 17.9°C. The third-generation variants have half-lives of thermal inactivation \( \sim 1,250 \) to 2,000 times longer than the parent at 45°C and their \( T_{50} \) are 18.5 to 19.3°C higher. The 12x mutant has a half-life at 45°C that is approximately 7,000 times longer than the parent.

**Activities of the evolved PTDHs.** The kinetic parameters of the parent and evolved PTDHs toward the substrates NAD, NADP and phosphite are listed in Table 2. All five first-generation variants show similar \( k_{\text{cat}} \) and \( K_{m,\text{NAD}} \), compared to the parent. First-generation variants 1-8D6 and 1-25E8 exhibit a slight increase in \( K_{m,\text{Pt-H}} \). The 4x and 5x mutants both have lower activities than the parent, and the 4x mutant exhibits about a 2.5-fold increase in \( K_{m,\text{Pt-H}} \). All three second-generation variants and the 7x mutant have similar kinetic parameters to the 4x mutant. All five third-generation variants are slightly more active than the 7x mutant, while their \( K_{m,\text{NAD}} \) remains unchanged and their \( K_{m,\text{Pt-H}} \) increased slightly. The final 12x mutant retains approximately 75% of the parent enzyme's turnover number and has a slightly higher catalytic efficiency \( (k_{\text{cat}}/K_{m,\text{NAD}}) \). The 12x mutant exhibits a \( K_{m,\text{Pt-H}} \) similar to that of the parent. The kinetic parameters of the 12x mutant toward nicotinamide cofactor NADP⁺ were also determined. The evolved 12x mutant has similar \( k_{\text{cat}} \) and \( K_{m,\text{NADP}^+} \) compared to the parent: 12x mutant, \( k_{\text{cat}} = 80 \pm 6 \text{ min}^{-1} \), \( K_{m,\text{NADP}^+} = 49 \pm 5 \text{ mM}^{-1} \text{ min}^{-1} \) versus the parent, \( k_{\text{cat}} = 110 \pm 1 \text{ min}^{-1} \), \( K_{m,\text{NADP}^+} = 65 \pm 8 \text{ mM}^{-1} \text{ min}^{-1} \) (25).

The activity-temperature profile for the parent and the combined mutants is shown in Fig. 2. For each successive round (parent \( \rightarrow 4x \rightarrow 7x \rightarrow 12x \)), the activity-temperature profile broadens and the specific activities of PTDH enzymes increase with increasing temperature until the enzyme denatures. The temperature optimum, \( T_{\text{opt}} \), of the 12x mutant is 59°C, which is \( \sim 20°C \) higher than that of the parent, which is in good agreement with the observed increases in \( T_{50} \) (Table 1).

**Comparison with Candida boidinii formate dehydrogenase.** The half-lives of thermal inactivation of the 12x PTDH mutant and *Candida boidinii* FDH were measured at 50°C. Both enzymes exhibit first-order inactivation kinetics as shown in Fig. 3. At 50°C, the half-lives of thermal inactivation of the 12x PTDH mutant and FDH are 705 and 299 min, respectively. The \( t_{1/2} \) of FDH at 50°C is in good agreement with values reported elsewhere for the wild-type and recombinant FDH \( (t_{1/2} \geq 300 \text{ min at } 50°C \text{ in } 100 \text{ mM potassium phosphate buffer, pH 7.5}) \) (17). Slusarczyk and coworkers improved the oxidative stability of the wild-type FDH by replacing two cysteine resi-

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**Table 2. Kinetic parameters of the parent and evolved PTDH variants in 50 mM MOPS (pH 7.25) at 25°C**

<table>
<thead>
<tr>
<th>Generation</th>
<th>Variant</th>
<th>( k_{\text{cat}} ) (min⁻¹)</th>
<th>( K_{m,\text{NAD}} ) (μM, NAD)</th>
<th>( k_{\text{cat}}/K_{m,\text{NAD}} ) (μM⁻¹ min⁻¹)</th>
<th>( K_{m,\text{Pt-H}} ) (μM, Pt-H)</th>
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<td>4.0</td>
<td>57</td>
</tr>
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<td>66</td>
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<td>48</td>
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<td>78</td>
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</table>

* Average of all statistically relevant data (standard deviation <10%).

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**Figure 2. Activity-temperature profiles of the parent (○), 4x mutant (△), 7x mutant (●), and 12x mutant (○).**
dyes using site-directed mutagenesis, but this resulted in several mutants with significantly lower thermostability (17). Thus, the 12x mutant is substantially more thermostable than these FDH variants.

**NADH regeneration for l-tert-leucine production.** The 12x thermostable PTDH mutant was tested by coupling it with leucine dehydrogenase from *Bacillus cereus* for the production of *l*-tert-leucine in small-scale batch reactions. PTDH converts NADH into NADPH with the conversion of phosphite to phosphate. The regenerated NADH is then used by leucine dehydrogenase to convert trimethylpyruvate and ammonia to *l*-tert-leucine. The 12x mutant was compared to wild-type PTDH and commercially available FDH from *C. boidinii* to demonstrate its improved stability and effectiveness. Figure 4 shows the production of *l*-tert-leucine over 300 min with the three different regeneration enzymes. The wild-type PTDH precipitates rapidly within the first 30 min of the reaction, whereas the 12x mutant retains its stability and reaches 100% conversion. The 12x mutant also has a ~2-fold faster reaction rate than FDH under these conditions.

**DISCUSSION**

**Evolutionary strategy.** The evolutionary strategy implemented in this work consisted of random mutagenesis and high-throughput screening followed by site-directed mutagenesis to incorporate the best mutations into the parental template for the next round of directed evolution (Fig. 1). By introducing an average of only 1 to 2 amino acid substitutions per PTDH gene, it was possible to identify the contribution of individual amino acid substitutions on enzyme thermostability. The effects of thermostabilizing mutations are additive as found in many other enzymes (24, 28). Combined mutants (4x, 5x, 7x, and 12x) were significantly more thermostable than any single mutant in a given round. Similar to other directed evolution efforts, we found that the thermostability of PTDH could be increased significantly while retaining 75% of the enzyme’s catalytic activity at lower temperatures (5, 29).

**Structural analysis.** A three-dimensional homology model of the wild-type PTDH was created as described elsewhere (26), and the 12 thermostabilizing mutations were mapped into this model (Fig. 5). None of these mutations occurred near the three-active site residues (Arg237, Glu266, and His292), although one mutation (I150F) did occur in the Rossmann fold involved with NAD(P) binding. A mutation (R275Q) occurred in a loop region directly after the active site Glu266 in the sequence 267-DWARADRPR-275. It has been suggested that the three arginines in this region could be involved in binding substrate phosphate (26). Substituting arginine for glutamine at position 275 does not appear to affect the binding of phosphate according to the kinetic data (Table 2). Therefore, while the other two arginines may still be involved in binding phosphate, it appears that Arg275 is not essential to phosphate binding. The mutations A319E and A325V occur in the highly flexible C-terminal region.

Based on this structural model, the molecular basis for improved thermostability was further probed. The thermostabilizing mutations are all distributed over the surface of the enzyme except for mutations V71I and I150F. This finding underscores the importance of protein surface on stability as found in many other enzymes with improved thermostability (3, 29). It is consistent with the notion that the initial steps in protein unfolding during the irreversible thermal denaturation primarily involve surface-located parts of the protein (3). The two nonsurface mutations are located in β-sheets and both are buried in extremely hydrophobic regions within the protein. The isoleucine to phenylalanine substitution (I150F) in β5 is located in the Rossmann fold. Surprisingly, the incorporation of the large hydrophobic phenylalanine residue into the GxxGxGxxG (where x represents any type of amino acids) nucleotide binding motif does not seem to affect the enzyme’s ability to bind the nicotinamide cofactor NAD+. The $K_m$,NAD+ values for the parent and the I150F mutation (1-25E8) are essentially the same (66 and 75 μM, respectively). More im-
portantly, based on the homology model, the substitution of Met49, Val57, Leu93, and Phe81. Only two mutations (I313L and V315A) are located in α-helices. These substitutions may increase stability by increasing helical propensity since Leu and Ala have nonbranched side chains compared to Ile and Val, respectively. Three mutation sites, E130K, Q132R, and Q137R, are located in the loop region between α6/β3. Mutations Q132R and Q137R introduce positively charged residues which may increase protein stability by creating a more favorable surface charge distribution, as found in the engineered thermostable cold shock proteins (2). Mutation E130K is located at the dimer interface and may increase enzyme stability by creating beneficial electrostatic interactions between charged residues of the two subunits (E130K is within ~8Å of D272 of the other subunit).

The three mutations Q215L, R275Q, and L276Q are located in the loop region between β10/α13. Although it is surprising that substituting an uncharged hydrophilic residue for a nonpolar hydrophilic residue (Q215L) at the protein surface increases protein stability, and this effect has been observed in other thermostability studies (12, 13). Mutation R275Q substitutes a positively charged residue for an uncharged residue which may increase protein stability by influencing the surface charge distribution. The mechanism by which L276Q enhances thermostability is not clear. The two mutations on the highly flexible C-terminal region, A319E and A325V, most likely increase protein thermostability by anchoring the C terminus and this would make the protein more resistant to unfolding.

It should be noted that the above-proposed mechanisms for thermostabilization have been identified in many naturally occurring thermophilic and hyperthermophilic proteins as well as engineered thermostable proteins (3, 21). Further confirmation of these mechanisms should be greatly facilitated by the determination of the crystal structure of the 12x mutant and additional mutagenesis and stability analysis. However, since there are so many different structural routes to improved thermostability and the contribution of each route to the total free energy of stabilization is quite small, it seems very difficult to predict the specific thermostabilizing mutations for a protein of interest based on its structure, let alone a protein without a crystal structure such as PTDH.

In conclusion, this study shows that the thermostability of phosphite dehydrogenase can be rapidly improved by directed evolution without a loss of its catalytic efficiency. Various stabilization mechanisms have been uncovered, many of which are difficult to predict a priori, thus underscoring the notion that directed evolution is a generally applicable and highly effective approach for protein stabilization (29). The improved stability and effectiveness of the thermostable PTDH mutant were demonstrated in the production of an important specialty chemical, t-tert-leucine. The 12x mutant was found to be a more effective regeneration catalyst than the most successful and widely used NADH-regenerating enzyme, C. boidinii FDH, under the conditions tested. The results of this work will hopefully enable us to reach our ultimate goal of developing a low-cost, highly efficient biocatalyst for NAD(P)H regeneration.

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