Enzymology:
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First Evidence for Substrate Channeling between Proline Catabolic Enzymes

A VALIDATION OF DOMAIN FUSION ANALYSIS FOR PREDICTING PROTEIN-PROTEIN INTERACTIONS*

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Background: PRODH and P5CDH from Thermus thermophilus are monofunctional enzymes in proline catabolism.

Results: Steady-state kinetics and intermediate trapping data show the PRODH and P5CDH reactions are coupled by a channeling step.

Conclusion: Substrate channeling in monofunctional enzymes is achieved via weak interactions.

Significance: Evidence for substrate channeling between monofunctional proline catabolic enzymes is shown and confirms the Rosetta Stone hypothesis.

Proline dehydrogenase (PRODH) and Δ¹-pyrroline-5-carboxylate (P5C) dehydrogenase (P5CDH) catalyze the four-electron oxidation of proline to glutamate via the intermediates P5C and L-glutamate-γ-semialdehyde (GSA). In Gram-negative bacteria, PRODH and P5CDH are fused together in the bifunctional enzyme proline utilization A (PutA) whereas in other organisms PRODH and P5CDH are expressed as separate monofunctional enzymes. Substrate channeling has previously been shown for bifunctional PutA enzymes, but whether the monofunctional enzymes utilize an analogous channeling mechanism has not been examined. Here, we report the first evidence of substrate channeling in a PRODH-P5CDH two-enzyme pair. Kinetic data for the coupled reaction of PRODH and P5CDH from Thermus thermophilus are consistent with a substrate channeling mechanism, as the approach to steady-state formation of NADH does not fit a non-channeling two-enzyme model. Furthermore, inactive P5CDH mutants inhibit NADH production and increase trapping of the P5C intermediate in coupled assays of wild-type PRODH-P5CDH enzyme pairs, indicating that the mutants disrupt PRODH-P5CDH channeling interactions. A dissociation constant of 3 μM was estimated for a putative PRODH-P5CDH complex by surface plasmon resonance (SPR). Interestingly, P5CDH binding to PRODH was only observed when PRODH was immobilized with the top face of its (βα)₈ barrel exposed. Using the known x-ray crystal structures of PRODH and P5CDH from T. thermophilus, a model was built for a proposed PRODH-P5CDH enzyme channeling complex. The structural model predicts that the core channeling pathway of bifunctional PutA enzymes is conserved in monofunctional PRODH-P5CDH enzyme pairs.

The oxidative conversion of proline to glutamate, known as the proline catabolic pathway, involves two enzymes, a flavin-dependent proline dehydrogenase (PRODH) and a NAD⁺-dependent Δ¹-pyrroline-5-carboxylate (P5C) dehydrogenase (P5CDH) (Fig. 1). The intervening step involves the hydrolysis of P5C to L-glutamate-γ-semialdehyde (GSA). The proline catabolic pathway helps organisms respond to changes in the nutritional environment by initiating the breakdown of proline as a source for nitrogen, carbon, and energy (1–6). In most organisms, PRODH and P5CDH are expressed as separate enzymes. In Gram-negative bacteria, however, PRODH and P5CDH are fused into one polypeptide known as proline utilization A (PutA) (7).

A unique aspect of proline catabolism is the discovery of a substrate channeling mechanism for P5C/GSA transfer between the PRODH and P5CDH domains of PutA. Crystal structures of PutA enzymes from Bradyrhizobium japonicum (8) and Geobacter sulfurreducens (9) revealed that the two active sites are separated by a linear distance of 40–50 Å and connected by a curved tunnel that traverses ~70 Å. Kinetic data for these enzymes are consistent with a substrate channeling mechanism (8, 9). Blockage of the tunnel in B. japonicum PutA by mutagenesis abrogates channeling, which indicates that the tunnel functions as the conduit for the intermediate P5C/GSA (10). Furthermore, a substrate channeling mechanism is supported by kinetic studies of PutA from Escherichia coli (11) and Salmonella typhimurium (12) indicating that a general feature of PutA enzymes is direct transfer of P5C between the PRODH and P5CDH active sites.

The abbreviations used are: PRODH, proline dehydrogenase; BIAM, N-(biotinoyl)-N’-(iodoacetyl)ethylenediamine; CoO₂ anesthesia; DrP5CDH, Δ¹-pyrroline-5-carboxylate dehydrogenase from D. radiodurans; GSA, L-glutamate-γ-semialdehyde; o-AB, o-aminobenzaldehyde; P5C, Δ¹-pyrroline-5-carboxylate; P5CDH, Δ¹-pyrroline-5-carboxylate dehydrogenase; PutA, proline utilization A; Put2p, Δ¹-pyrroline-5-carboxylate dehydrogenase from S. cerevisiae; SPR, surface plasmon resonance; TtPRODH, proline dehydrogenase from T. thermophilus; TtP5CDH, Δ¹-pyrroline-5-carboxylate dehydrogenase from T. thermophilus.

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A potential rationale for channeling P5C/GSA may be to limit reactions with other molecules and regulate its signaling activity. P5C/GSA is appreciably labile and has been shown to react with metabolites (13, 14), inhibit enzymes (15–18), and affect signaling pathways (19–22). High levels of P5C/GSA were shown to generate inactive adducts with pyridoxal phosphate, leading to lower amounts of functional vitamin B6 in patients (13). P5C has also been shown to act as a signaling molecule in eukaryotes resulting in apoptosis (19, 20). Very recently, P5C was proposed to act as a signaling metabolite linking proline metabolism with lipid oxidation in \textit{Caenorhabditis elegans} (22). A more compelling reason for channeling P5C/GSA, however, may be to avoid futile cycling (11, 12, 23, 24). Cycling between proline catabolism and proline biosynthesis would be energetically costly for the cell. Thus, substrate channeling may be a necessary mechanism for maintaining efficient proline catabolic flux.

A major question in proline catabolism is whether PRODH and P5CDH enzymes that are not covalently linked utilize a substrate channeling mechanism similar to PutA. According to the Rosetta Stone hypothesis, which predicts protein-protein interactions based on gene fusion events (25, 26), individual PRODH and P5CDH enzymes are expected to interact, possibly in a manner analogous to PutA. To date, however, no detailed studies have been performed to test substrate channeling between monofunctional PRODH and P5CDH enzymes.

To address this gap, we chose to characterize the coupled reaction of PRODH and P5CDH enzymes from \textit{Thermus thermophilus}. The x-ray crystal structures of \textit{T. thermophilus} PRODH (TtPRODH) and P5CDH (TtP5CDH) have been solved providing a solid framework for investigating substrate channeling with these enzymes (27, 28). TtPRODH has a \((\alpha_8 \beta_8)\) barrel structure which is conserved in the PRODH domain of PutA (8, 9, 29). TtP5CDH shares the fundamental aldehyde dehydrogenase (ALDH) dimeric structure of the ALDH superfamily (28), which is found in PutA and P5CDH from eukaryotes (30, 31). Thus, TtPRODH and TtP5CDH have domain structures that are conserved from prokaryotes to higher eukaryotes.

Here, we present a variety of kinetic strategies for evaluating substrate channeling between TtPRODH and TtP5CDH. The sum of the kinetic evidence supports a substrate channeling mechanism for the TtPRODH-TtP5CDH coupled reaction. TtPRODH and TtP5CDH interactions were also explored by surface plasmon resonance and used to help model a putative TtPRODH-TtP5CDH bienzyme complex. The structural model suggests that TtPRODH-TtP5CDH forms a channeling pathway that is conserved with covalently linked PRODH-P5CDH domains in PutA.

**EXPERIMENTAL PROCEDURES**

**Materials**—All chemicals and buffers were obtained from Sigma-Aldrich or Thermo Fisher Scientific Inc. unless otherwise noted. (DL)-P5C (50/50 mixture) was synthesized by the method of Williams and Frank and stored in 1 M HCl at 4 °C and was neutralized the day of experiments on ice by titrating with 6 M NaOH (32). Assays containing exogenously added (DL)-P5C contained \(-150 \text{ mM NaCl from the pH neutralization.} N-(\text{Biotinyl})-N’-(\text{iodoacetyl})\text{ethylenediamine} (\text{BIAM}) and dimethyl sulfoxide were obtained from Molecular Probes. The streptavidin (SA) sensor chip and PD-10 desalting columns were purchased from GE Healthcare. Active site and surface mutants for this study were generated using the GeneTailor mutagenesis kit (Invitrogen). All mutations were confirmed by DNA sequencing. All experiments were conducted in nanopure water.

**Purification of Enzymes**—TtPRODH wild-type and mutants (R288M/R289M, S9C, and S88C), TtP5CDH wild-type and mutant C322A, and \textit{Deinococcus radiodurans} P5CDH (DrP5CDH) mutant C325A were expressed from a pKA8H expression vector while the \textit{Saccharomyces cerevisiae} P5CDH (Put2p) mutant C351A was expressed from a pET14b expression vector (pET14b-PUT2). All enzymes were expressed and purified with a N-terminal His tag as previously described in \textit{E. coli} BL21(DE3) pLysS cells (27, 31, 33). Assays with non-His tag P5CDH enzymes confirmed that the His tag does not affect the Kirsch assays used for testing substrate channeling. The final concentration of purified protein was determined using the bicinchoninic acid method (Thermo Scientific Pierce) using bovine serum albumin (BSA) as a standard and is expressed as concentration of monomer unless stated otherwise (34).

**PRODH and P5CDH Kinetic Assays**—TtPRODH activity was measured at 25 °C using Coenzyme Q\(_{1}\) (CoQ\(_{1}\)) as the terminal electron acceptor. Kinetic parameters for proline and CoQ\(_{1}\) were determined in assays using TtPRODH (0.5 μM) in 50 mM phosphate buffer (pH 8.0) and varying proline (0–100 μM) at a fixed concentration of CoQ\(_{1}\) (150 μM) or varying CoQ\(_{1}\) (0–300 μM) at a fixed proline concentration (150 μM). Reaction progress curves were monitored by following reduction of CoQ\(_{1}\) at 275 nm (\(ε = 13700 \text{ M}^{-1} \text{cm}^{-1}\)) using a 0.15 cm path length on a Hi-Tech Scientific SF-61DX2 stopped-flow instrument (TgK Scientific) at 25 °C (11). The kinetic parameters \(K_m\) and \(k_{cat}\) were estimated by fitting initial velocities to the Michaelis-Menten equation (35, 36).

P5CDH activity was measured by monitoring NADH formation at 340 nm (\(ε = 6200 \text{ M}^{-1} \text{cm}^{-1}\)) at 25 °C. Assays were performed using 0.5 μM P5CDH, 0.2 mM NAD\(^+\), 50 mM potas-
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Coupled TtPRODH-TtP5CDH Activity Assays—The optimal ratio of TtPRODH to TtP5CDH was determined first by varying TtPRODH (0.1 μM to 10 μM) and holding TtP5CDH fixed at 0.5 μM. Assays were performed with proline (1 mM), CoQ1 (0.1 mM), and NAD⁺ (0.2 mM) in 50 mM potassium phosphate (pH 7.5) and 25 mM NaCl. Equal volumes of substrate and enzyme solutions were mixed on a SF-61DX2 stopped-flow spectrophotometer and the reaction was monitored by following NAD⁺ reduction at 340 nm (ε₃₄₀₋₅₉₅ = 2900 M⁻¹ cm⁻¹) (39). Coupled TtPRODH-TtP5CDH activity assays were performed as described above except in the presence of 1 mM o-AB. The maximum amount of P5C trapped by o-AB was determined in assays without NAD⁺. The effect of inactive mutants TtP5CDH C322A and Put2p C351A on P5C trapping was determined by assays using increasing concentrations of the inactive P5CDH mutants (0–37.5 μM, monomer). The ratio of inactive P5CDH mutant to wild-type TtP5CDH ranged from 0 to 75. Assays were performed in a 96 well plate (200 μl total assay volume) and initiated by adding a mixture of substrate mixture with final concentrations after mixing of 0.5 μM TtPRODH, 0.5 μM TtP5CDH, 0–37.5 μM inactive P5CDH mutant, 1 mM proline, 0.2 mM NAD⁺, 0.2 mM CoQ1, and 1 mM o-AB in 50 mM potassium phosphate buffer (pH 7.5, 25 mM NaCl). P5C-o-AB complex formation was followed at 443 nm for 20 min on a Powerwave XS 96-well plate reader (Biotek).

SPR Analysis of TtPRODH-TtP5CDH Interactions—Cysteine residues were introduced on opposite sides of TtPRODH at surface residues Ser-9 and Ala-88. TtPRODH mutants S9C and A88C were then purified as described above. Purified TtPRODH mutants S8C and A88C were produced in E. coli and purified as described above. Purified TtPRODH mutants S8C and A88C required the presence of the reducing agent Tris[3-hydroxypropyl]phosphine to prevent precipitation during purification. The specific activities of TtPRODH mutants S8C and A88C relative to wild-type TtPRODH were 74 and 60%, respectively. Purified TtPRODH mutants S8C and A88C were labeled with biotin at the incorporated cysteine residue using the thiol-reactive BIAM reagent. A 5 mM stock of BIAM was prepared by dissolving in dimethyl sulfoxide. Under anaerobic conditions, purified TtPRODH mutants S8C and A88C and BIAM were mixed at final concentrations of 51 μM (TtPRODH) and 715 μM (BIAM). The labeling reaction was allowed to proceed overnight at 4 °C until the next morning when 25 mM β-mercaptoethanol was added to quench unreacted BIAM. Biotin-labeled TtPRODH was then separated from quenched BIAM by passing the reaction mixture through a PD-10 desalting column. The concentration of total protein was measured using the Pierce 660 nm Protein Assay Reagent (Pierce). After the biotinylation procedure, the specific activity of both mutants dropped to 44% relative to wild-type TtPRODH. Biotin labeling of TtPRODH was confirmed by Western blot analysis using streptavidin-conjugated horseradish peroxidase (Pierce). ECL prime reagents (GE Biomedical, Piscataway, NJ) were used for detection. The biotinylated TtPRODH was purified by immobilized BSA and the concentration determined using the Pierce 660 nm Protein Assay Reagent (Pierce).
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Healthcare) were used for detection and visualization of biotin-labeled TtPRODH.

The interaction between TtPRODH and TtP5CDH was studied using a BIAcore 3000 (BIACORE AB, Uppsala, Sweden) instrument maintained in the Nanomaterials Characterization Core Facility at the University of Nebraska Medical Center. A streptavidin (SA) sensor chip was first washed with HBS-EP buffer (10 mM HEPES, pH 7.4, 0.15 M NaCl, 3.4 mM EDTA, and 0.05% surfactant P20) for 6 min (20 μl/min flow rate). Prior to immobilization, the SA sensor chip was activated with three consecutive 40 μl injections of activating buffer (1 M NaCl and 50 mM NaOH). Biotinylated TtPRODH mutants S9C and A88C were then diluted in HBS-EP to a final concentration of 250 μM and injected at a flow rate of 30 μl/min followed by a 5 min dissociation phase at a flow rate of 30 μl/min.

The binding of TtP5CDH to immobilized TtPRODH S9C and A88C were estimated by injecting increasing concentrations of TtP5CDH (0.5–7.5 μM, monomer) pre-equilibrated in HBS-EP buffer. From lowest to highest concentration, wild-type TtP5CDH was injected for 2 min (30 μl/min) followed by a 5 min dissociation phase at a flow rate of 30 μl/min with HBS-EP buffer. The SA-TtPRODH binding surface was regenerated using 90 μl of 2 M NaCl (30 μl/min) and equilibrated with HBS-EP buffer (30 μl/min, 5 min) after each injection of TtP5CDH. The resulting sensorgrams were globally fitted to 1:1 Langmuir binding model using BIA-evaluation software 4.1. The experiments were carried out at 25 °C. All buffers were filtered and degassed prior to injection on SA sensor chip. Signals from the reference flow cells were subtracted from the sensorgrams.

RESULTS

Inhibition of TtP5CDH by Proline—X-ray crystal structures of TtP5CDH (28) have shown that both glutamate (PDB ID:2BHQ) and proline (PDB ID:2J40) bind to the active site of TtP5CDH with identical orientations of the carboxylate group resulting in favorable hydrogen bonding interactions with Ser-321 and backbone amide nitrogen atoms. Proline inhibition of TtP5CDH, however, has not been characterized. Thus, prior to examining the coupled TtPRODH-TtP5CDH reaction, proline inhibition of the TtP5CDH reaction was determined.

The steady-state kinetic parameters determined for TtP5CDH were \( K_m = 43 \pm 1 \mu M \) l-P5C and \( k_{cat} = 0.52 \pm 0.03 \text{ s}^{-1} \). When varying proline, the decrease in P5CDH activity fit well to a competitive inhibition model with an inhibition constant \( (K_i) \) of 3.9 mM proline (Fig. 2A). A double reciprocal plot of 1/V versus 1/[S] with varying proline concentrations is also shown (Fig. 2B) and is consistent with proline as a competitive inhibitor of P5CDH with respect to P5C/GSA.

Transient Time of Coupled TtPRODH-TtP5CDH Reaction—The ratio of TtPRODH:TtP5CDH was initially optimized to ensure that the coupled TtPRODH-TtP5CDH reaction was not limited by the concentration of either enzyme and to prevent surplus formation of P5C by excess TtPRODH. To avoid proline inhibition of TtP5CDH, we characterized coupled TtPRODH-TtP5CDH activity using 1 mM proline which is \( < K_{(pro)} \) (3.9 mM). Fig. 3 shows the dependence of NADH formation on the concentration of TtPRODH while keeping the substrates and TtP5CDH fixed. With TtP5CDH fixed at 0.5 μM, the NADH formation rate plateaus above 1 μM TtPRODH. Thus, an equimolar mixture of 0.5 μM TtPRODH and 0.5 μM TtP5CDH was considered to be appropriate for testing substrate channeling.

The reaction progress curve of NADH formation using an equimolar mixture of TtPRODH-TtP5CDH (0.5 μM:0.5 μM) was next examined and compared with that of a diffusion-limited two-enzyme reaction model (Eq. 2). An experimental transient time of 38.5 s was observed for an equimolar mixture of

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**FIGURE 2. Inhibition pattern of TtP5CDH with L-proline.** A, non-linear least squares fit to the competitive inhibition model (Eq. 1) with 0 (●), 2 (○), 4 (▼), 6 (△), 8 (■), 10 (□), and 15 mM (▲) proline and varied P5C. Best fit parameters were \( K_m = 0.43 \pm 0.02 \text{ s} \), \( K_i = 27.7 \pm 4.3 \mu M \), and \( K_i(pro) = 3.9 \pm 0.7 \text{ mM} \) proline. B, double reciprocal plot of the same data in panel A showing competitive inhibition pattern.
TtPRODH-TtP5CDH prior to achieving a steady-state velocity of 0.08 M/s NADH s⁻¹ (Fig. 4). The transient time for a non-channeling TtPRODH-TtP5CDH coupled reaction was then simulated using the kinetic constants of the individual reactions. The steady-state kinetic parameters determined for TtPRODH were $K_m$ of 8.2 ± 0.3 mM proline and 162 ± 8 μM CoQ₁, with an overall $k_{cat}$ of 17.0 ± 0.4 s⁻¹. Under the assay conditions for the coupled reaction, TtPRODH generates P5C/GSA at 0.08 M/s (v₁). The predicted transient time before reaching a linear rate of NADH formation is 164.2 s for a non-channeling TtPRODH-TtP5CDH coupled reaction. Thus, the transient time for the experimental reaction is suggestive of substrate channeling between TtPRODH and TtP5CDH.

One explanation for the shorter lag time of the TtPRODH-TtP5CDH-coupled reaction, however, could be allosteric activation of one or both enzymes (40). Upon adding up to 50-fold excess TtP5CDH, however, we observed no increase in TtPRODH activity. Likewise, no activation of TtP5CDH activity was observed with 50-fold excess of the TtPRODH mutant R288M/R289M. Thus, we found no evidence of allosteric activation between TtPRODH and TtP5CDH.

**P5C Trapping Shows Evidence of Channeling**—If the intermediate P5C/GSA is channeled between TtPRODH and TtP5CDH, then the amount of intermediate in the bulk solvent should be sensitive to TtP5CDH activity, which requires NAD⁺. The ability of excess o-AB (1 mM), which traps P5C as a dihydroquinazolinium compound that is detected at 443 nm, was examined using an equimolar mixture of TtPRODH (0.5 μM) and TtP5CDH (0.5 μM) with TtPRODH varied from 0.1 to 10 μM. Assays were performed in 50 mM potassium phosphate (pH 7.5, 25 mM NaCl), and NADH formation was monitored at 340 nm. Data points are the mean ± S.D. from three assays.

**FIGURE 3.** Dependence of the NADH formation rate of P5CDH on TtPRODH concentration. The reaction mixture contained 0.5 μM TtP5CDH, 1 mM proline, 0.2 mM NAD⁺, and 0.1 mM CoQ₁ with TtPRODH varied from 0.1 to 10 μM. Assays were performed in 50 mM potassium phosphate (pH 7.5, 25 mM NaCl), and NADH formation was monitored at 340 nm. Data points are the mean ± S.D. from three assays.

**FIGURE 4.** Experimental and simulated reaction progress curves of coupled TtPRODH and TtP5CDH activity. The solid line is the steady-state formation of NADH followed at 340 nm in an assay containing an equimolar mixture of TtPRODH (0.5 μM) and TtP5CDH (0.5 μM), 1 mM proline, 0.2 mM NAD⁺, and 0.1 mM CoQ₁ (pH 7.5). The dotted line is simulated NADH formation from a non-channeling free diffusion model (Eq. 2) using experimentally determined values of $v_1$ (0.08 μM/s), $K_m$ (43 μM), and $v_2$ (0.26 μM/s).

**FIGURE 5.** Trapping of P5C intermediate shows evidence of substrate channeling. A, solid line is P5C-o-AB production from an equimolar mixture of TtPRODH (0.5 μM) and TtP5CDH (0.5 μM) with 1 mM proline and 0.2 mM CoQ₁. The dotted line is the same reaction mixture but with 0.2 mM NAD⁺. Assays were performed in 50 mM potassium phosphate (pH 7.5) and 1 mM o-AB. Complex formation of P5C-o-AB is followed at 443 nm. B, steady-state P5C-o-AB formation rate from an equimolar mixture of TtPRODH (0.5 μM) and TtP5CDH (0.5 μM) as a function of increased ratio of inactive TtP5CDH C322A (filled circles) or Put2p C351A mutants (open circles) (mutant/wild-type TtP5CDH). Assays were performed in 50 mM potassium phosphate (pH 7.5) containing 1 mM proline, 0.2 mM NAD⁺, 0.2 mM CoQ₁, and 1 mM o-AB. Percent leakage was estimated by dividing the rate of P5C-o-AB formation by the maximum rate of P5C-o-AB formation determined by assays without NAD⁺ (dotted line).
In the presence of NAD$^+$ (0.2 mM), o-AB trapping of P5C is significantly reduced indicating that a large fraction of P5C is not solvent accessible during the TtPRODH-TtP5CDH coupled reaction. In these assays, the apparent fraction of P5C that is channeled is 80%. Trapping assays performed with the TtP5CDH C322A inactive mutant showed no change in P5C trapping with NAD$^+$. A second approach for testing P5C release into bulk solvent was developed using a strategy devised by Geck and Kirsch (38). In this strategy, inactive mutants are used as competitors of the putative channeling complex. For these assays, an inactive mutant of TtP5CDH was generated by mutating Cys-322, which is the essential active site nucleophile. If the coupled TtPRODH-TtP5CDH reaction involves a channeling complex, the inactive TtP5CDH C322A mutant is anticipated to increase P5C trapping by disrupting wild-type TtPRODH-TtP5CDH interactions. Fig. 5B shows that the amount of P5C trapped by o-AB in a TtPRODH-TtP5CDH coupled reaction is about 19% of the maximum possible. Thus, the leakage of P5C during the coupled TtPRODH-TtP5CDH reaction is estimated to be ~19%. Addition of the inactive TtP5CDH C322A mutant shows a concentration dependent increase in trapped P5C with 92% leakage estimated at 50-fold excess of the C322A mutant relative to wild-type TtP5CDH. This result indicates the C322A mutant causes a nearly 5-fold increase in P5C release into bulk solvent consistent with disrupting a TtPRODH-TtP5CDH channeling complex. We also tested the effect of an inactive P5CDH from another organism. Fig. 5B shows that an inactive P5CDH mutant from yeast S. cerevisiae, Put2p C351A, had a much smaller effect with an increase of only 33% in P5C trapping. Thus, the effect appears to be specific for the TtP5CDH C322A mutant.

Inactive Mutants Disrupt TtPRODH-TtP5CDH-coupled Reaction—Substrate channeling in the TtPRODH-TtP5CDH coupled reaction was next tested using the original strategy of Geck and Kirsch. In these assays, the effects of inactive TtP5CDH and TtPRODH mutants on the overall coupled reaction rate were determined. The inactive TtPRODH mutant R288M/R289M was generated by replacing two catalytic arginine residues (Arg-288 and Arg-289) that are essential in PRODH and PutA for proline binding via ionic interactions with the carboxylate moiety of proline (9, 30, 41, 42).

Fig. 6 shows that the inactive TtP5CDH C322A mutant significantly decreased the rate of the PRODH-P5CDH coupled reaction in a concentration dependent manner (Fig. 6). The rate of NADH formation decreased by 60% in the presence of 50-fold excess of TtP5CDH C322A (25 μM) relative to wild-type TtP5CDH (0.5 μM). To test the specificity of this effect, inactive P5CDH from other organisms were used in identical assays. An inactive mutant of P5CDH from D. radiodurans (DrP5CDH C325A) was added up to 50-fold excess relative to TtP5CDH. The maximum decrease in PRODH-P5CDH activity was 16% at a 10-fold excess of DrP5CDH C325A mutant. Also, no decrease in activity was observed with 50-fold excess of the Put2p inactive mutant C351A or 50-fold excess of BSA in the PRODH-P5CDH coupled assay (data not shown). Thus, the observed decrease in the PRODH-P5CDH activity appears to be specific for the TtP5CDH C322A mutant.

Adding the TtPRODH inactive mutant R288M/R289M to the PRODH-P5CDH reaction also decreased activity but to a lesser extent. A 50-fold excess of TtPRODH R288M/R289M (25 μM) relative to wild-type TtPRODH (0.5 μM) resulted in a 35% decrease in the rate of NADH formation (Fig. 6).

Evidence for TtPRODH-TtP5CDH Binding—Prompted by the kinetic evidence of a substrate channeling mechanism for the coupled TtPRODH-TtP5CDH reaction, potential binding interactions between TtPRODH and TtP5CDH were investigated by SPR. For the SPR analysis, TtPRODH was immobilized on a Biacore streptavidin (SA) sensor chip in two different orientations. TtPRODH lacks cysteine residues making it convenient to specifically label engineered cysteine residues. Surface residues Ser-9 and Ala-88 were chosen for cysteine replacement. Ser-9 and Ala-88 are near the top and bottom of the PRODH β$\alpha_8$ barrel, respectively, enabling TtPRODH to be immobilized in two different orientations (Fig. 7A). Immobilization of TtPRODH S9C should leave the bottom face of the PRODH β$\alpha_8$ barrel exposed to solvent (Fig. 7A) whereas the top face of the PRODH β$\alpha_8$ barrel should be solvent accessible with TtPRODH A88C.

Fig. 7B shows that upon injecting TtP5CDH, a binding response was observed with immobilized TtPRODH A88C but not with immobilized TtPRODH S9C. This result suggests that TtP5CDH distinguishes between the two orientations of TtPRODH and specifically recognizes the configuration of immobilized TtPRODH A88C. It also indicates that the immobilized orientation of TtPRODH S9C conceals a surface on TtPRODH that is critical for TtP5CDH binding. No binding response was observed for DrP5CDH or BSA with the immobilized TtPRODH mutants (data not shown) further indicating a specific binding event between immobilized TtPRODH A88C and TtP5CDH.

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The binding kinetics of TtP5CDH to immobilized TtPRODH A88C were then evaluated (Fig. 7C). SPR analysis shows TtP5CDH binds to TtPRODH A88C with a dissociation constant ($K_D$) of 3 μM. The estimated $k_{on}$ and $k_{off}$ rate constants for TtP5CDH binding were $1.49 \times 10^5$ M$^{-1}$s$^{-1}$ and $4.51 \times 10^{-3}$ s$^{-1}$, respectively.

**DISCUSSION**

The channeling of metabolites is generally perceived to enhance metabolic flux thereby benefiting cellular fitness (43–45). Metabolic channeling has been proposed for various pathways such as fatty acid β-oxidation (46), tricarboxylic acid cycle (47), and purine biosynthesis (48, 49). Generating kinetic and structural evidence for channeling between two enzymes of consecutive steps in a pathway, however, is challenging. As a simple two-enzyme system, the proline catabolic pathway affords the opportunity to investigate mechanisms of substrate channeling and their importance *in vivo*. In addition, the two enzymes of proline catabolism are either free or fused together (i.e. PutA) enabling the comparison of two biological strategies for handling the same metabolite. The Rosetta Stone hypothesis predicts that if two enzyme pairs are fused together in one organism, then in other systems in which they are free, they will interact (25, 26).

The first observation of substrate channeling for the PRODH-P5CDH reaction was reported in PutA from *Salmonella typhimurium* by Surber and Maloy who concluded that the oxidation of proline to glutamate involves a leaky channel (12). In recent years, kinetic and structural data of PutA enzymes from *B. japonicum* and *G. sulfurreducens* have provided unprecedented molecular details of P5C/GSA channeling including the three-dimensional space and electronic environment of the intervening cavity between PRODH and P5CDH, potential exit and entry sites for water, and replacement of residues lining the cavity with bulkier side chains that block substrate channeling. In addition, a novel hysteresis mechanism was found for PutA from *E. coli* suggesting a channeling step in the overall PRODH-P5CDH reaction becomes activated in the first few catalytic turnovers. From these aforementioned studies, substrate channeling as a mechanism for the two-step conversion of proline into glutamate is now well established in PutA enzymes.

In this study, we present the first evidence of substrate channeling for a PRODH-P5CDH pair that are not covalently linked. Rate measurements for the coupled TtPRODH-TtP5CDH enzyme pair demonstrates that steady-state formation of NADH is achieved faster than the theoretical approach to steady-state by two individual enzymes. Although such rapid product formation can be explained by allosteric activation by one enzyme on the other (40), our results show that this is not the case for TtPRODH and TtP5CDH. The observed lag time is greater than that observed for PutA. The lag times reported for PutA enzymes thus far range from no apparent lag to 23 s for *E. coli* PutA (8, 9, 11). The longer lag time of the TtPRODH-
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TtP5CDH pair most likely reflects the dynamic nature of a potential channeling pathway between two free enzymes.

The experiments utilizing the catalytically inactive TtP5CDH C322A mutant were very useful for testing substrate channeling in the TtPRODH-TtP5CDH reaction. The inactive TtP5CDH mutant clearly disrupted the TtPRODH-TtP5CDH coupled reaction as observed by decreased NADH formation and increased trapping of the P5C intermediate. As noted by Yadid et al., assays involving trapping of the intermediate provide a rigorous test for channeling, and kinetic evidence for protection of a reactive intermediate implies a protein-protein complex (50). Thus, our kinetics data provide strong evidence that the TtPRODH-TtP5CDH reaction employs a channeling mechanism and imply a protein-protein complex.

It appears that TtPRODH and TtP5CDH form a weak transient complex. Such complexes are characterized by a $K_D$ in the micromolar range (51) and can be challenging to study using traditional biophysical methods. We found no evidence of a stable complex using gel filtration chromatography (data not shown) but were able to estimate a $K_D$ of $\sim 3 \mu M$ for TtPRODH-TtP5CDH using SPR. These results are consistent with TtPRODH and TtP5CDH forming a weak transient complex. Thus, although PRODH and P5CDH from *T. thermophilus* are not covalently linked as in PutA and do not form a tightly bound complex, substrate channeling is still achieved via dynamic interactions. We note that there is precedent for weak transient enzyme–enzyme complexes in other metabolic pathways, such as purine biosynthesis (48), the aspartate pathway of amino acid biosynthesis (52), branched chain amino acid catabolism (53), and the tricarboxylic acid cycle (47). In particular, the $K_D$ of 3 $\mu M$ that we measured for TtPRODH-TtP5CDH is very similar to that of 2.8 $\mu M$ measured in branched chain amino acid catabolism (53).

The advantage of a dynamic channeling system versus a fixed system such as PutA, may be the ability to regulate P5C/GSA levels and utilize P5C as a metabolite signaling molecule (19–22) or drive the proline-P5C redox cycle (54–56). If the consecutive enzymes were fixed together, the potential to regulate P5C would be limited. In organisms with free PRODH and P5CDH, additional regulation of the proline catabolic pathway is possible due to dynamic interactions. Thus, a stable and perfect channeling enzyme pair would not necessarily be a benefit for the cell.

Crystal structures of PutA (8, 9) provide a basis for building testable models of the TtPRODH-TtP5CDH complex. TtPRODH and TtP5CDH have the same folds as the respective domains of PutA. Therefore, a simple model of the channeling complex was made by superimposing a TtP5CDH dimer and two copies of the TtPRODH $\beta_8 \alpha_8$ barrel onto a dimer of *B. japonicum* PutA (PDB 3HAZ) (Fig. 8A). In this model, the top face of the TtPRODH $\beta_8 \alpha_8$ barrel forms the interface with TtP5CDH (Fig. 8B). This prediction is consistent with the SPR data showing that TtP5CDH specifically binds to the top face of the TtPRODH barrel but not to the bottom face (Fig. 7B). The size of the interface is also reasonable. According to a trend observed by Nooren and Thornton, a transient heterodimer with a $K_D$ $\sim 3 \mu M$ can be expected to have an interface contact area of $\sim 500 \AA^2$ (57). From our model, we calculated a contact area of 700 $\AA^2$ for the binding interface between TtPRODH and TtP5CDH (58), which is consistent with this trend. TtP5CDH also forms a trimmer-of-dimers hexamer at $> 1$ mg/ml ($>17 \mu M$ monomer). Application of the 3-fold symmetry operator of the TtP5CDH hexamer to the model in Fig. 8A generates a dodecamer model (Fig. 8C). Notice that in this model, the NAD$^+$ binding site is visible from the exterior indicating that the NADH and glutamate can exit into the bulk solvent rather than into the hexameric core region. Whether such a large assembly is present at the relatively low protein concentrations used in our assays (0.5 $\mu M$) is unknown, but if so, the population is likely low. These models will be tested by site-directed mutagenesis of the predicted interfaces.

In summary, the results here show that the two-step reaction catalyzed by PRODH and P5CDH from *T. thermophilus* involves a substrate channeling mechanism. This is the first evidence of substrate channeling between PRODH and P5CDH enzymes that are not fused together. This study also provides a useful approach for examining substrate channeling in PRODH...
and P5CDH enzymes from other organisms as well as other enzyme pairs that catalyze consecutive reactions.

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REFERENCES

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