Orotidine 5′-Monophosphate Decarboxylase: The Operation of Active Site Chains Within and Across Protein Subunits

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ABSTRACT: The D37 and T100′ side chains of orotidine 5′-monophosphate decarboxylase (OMPDC) interact with the C-3′ and C-2′ ribosyl hydroxyl groups, respectively, of the bound substrate. We compare the intra-subunit interactions of D37 with the inter-subunit interactions of T100′ by determining the effects of the D37G, D37A, T100′G, and T100′A substitutions on the following: (a) $k_{cat}$ and $K_m$ values for the OMPDC-catalyzed decarboxylations of OMP and 5-fluoroornotidine 5′-monophosphate (FOMP) and (b) the stability of dimeric OMPDC relative to the monomer. The D37G and T100′A substitutions resulted in 2 kcal mol$^{-1}$ increases in $\Delta G^\ddagger$ for $k_{cat}/K_m$ for the decarboxylation of OMP, while the D37A and T100′G substitutions resulted in larger 4 and 5 kcal mol$^{-1}$ increases, respectively, in $\Delta G^\ddagger$. The D37G and T100′A substitutions both resulted in smaller 2 kcal mol$^{-1}$ decreases in $\Delta G^\ddagger$ for the decarboxylation of FOMP compared to that of OMP. These results show that the D37G and T100′A substitutions affect the barrier to the chemical decarboxylation step while the D37A and T100′G substitutions also affect the barrier to a slow, ligand-driven enzyme conformational change. Substrate binding induces the movement of an $\alpha$-helix (G′98–S′106) toward the substrate C-2′ ribosyl hydroxyl bond at the main subunit. The T100′G substitution destabilizes the enzyme dimer by 3.5 kcal mol$^{-1}$ compared to the monomer, which is consistent with the known destabilization of $\alpha$-helices by the internal Gly side chains [Serrano, L., et al. (1992) Nature, 356, 453–455]. We propose that the T100′G substitution weakens the $\alpha$-helical contacts at the dimer interface, which results in a decrease in the dimer stability and an increase in the barrier to the ligand-driven conformational change.

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Orotidine 5′-monophosphate decarboxylase (OMPDC) is a dimeric enzyme composed of two identical monomers. The enzyme catalyzes the decarboxylation of OMP to form UMP through a vinyl carbanion reaction intermediate (Scheme 1). The enzyme provides an enormous 31 kcal mol$^{-1}$ stabilization of the decarboxylation transition state. This stabilization has been partitioned into roughly equal contributions from protein interactions with the following three substrate fragments (Scheme 2): the nonreacting phosphodianion and ribosyl groups and the reacting pyrimidine ring. The binding interactions of the phosphodianion and ribosyl substrate pieces have been partitioned into interactions that are expressed at the Michaelis complex and interactions that only develop when approaching the reaction transition state.

Scheme 3 shows the mechanism that provides the specificity for the binding interactions of the phosphodianion and the ribosyl hydroxyls at the decarboxylation transition state. The substrate binding interactions are utilized in order to drive an energetically unfavorable protein conformational change from the inactive open form of OMPDC ($E_O$) to the active Michaelis complex ($E_M$), where OMP is locked in a structured protein cage. The full intrinsic substrate binding energy, which is expressed by $(K_m)_\text{int}$ for the binding of OMP to $E_O$ is greater than the observed binding energy by the amount of binding energy that is utilized to drive the enzyme conformational change ($((K_m)_\text{int}/K_m = K_C \ll 1)$. The full intrinsic substrate binding energy is expressed at the rate determining transition state so that the value of $k_{cat}/K_m$ for the enzyme turnover is the same as that for a second hypothetical reaction that proceeds through the closed enzyme complex $E_C$–OMP without the requirement for an uphill enzyme conformational change.

We are working to define the roles of the active site amino acid side chains in OMPDC-catalyzed decarboxylation (Figure 1). We first examined the dianion gripper side chains Q215, Y217, and R235 as well as S154, which connects the phosphodianion gripper loop (P202–V220) to the pyrimidine umbrella loop (A151–T165). These side chain interactions provide a significant fraction of the driving force that activates the ligand-driven change in the enzyme conformation (Scheme 3), which results in the immobilization of two flexible...
In this paper, we focus on the interactions of the D37 and T100′ side chains with the C-3′ and C-2′ hydroxyls, respectively, of the substrate ribofuranosyl ring, where the D37 interaction is within a single enzyme subunit while the T100′ interaction spans the two subunits (Figure 1). Wolfenden and co-workers previously characterized the effects of D37A and T100′A substitutions on the kinetic parameters for the OMPDC-catalyzed decarboxylations of OMP and 2′-deoxyOMP. We revisit this problem and report the effects of a more extensive range of D37G, D37A, T100′A, and T100′G variants on the kinetic parameters for the OMPDC-catalyzed decarboxylations of OMP and 5′-fluoroorotidine 5′-monophosphate (FOMP) and, for the first time, examine the effect of these substitutions on the association constant $K_a$ for the formation of the active dimer of OMPDC from the inactive monomer. The D37G substitution is more conservative than that of D37A with respect to the effect on the activation barrier for the OMPDC-catalyzed decarboxylation. By contrast, the T100′A substitution is more conservative than that of T100′G by the same criteria, while the T100′G substitution results in an unusually large decrease in the stability of the active OMPDC dimer relative to the inactive monomer.

**METHODS**

**Materials.** OMP and FOMP were prepared enzymatically by literature procedures. 3-(N-Morpholino)-propanesulfonic acid (MOPS) and imidazole were purchased from Sigma (St. Louis, MO). Sodium hydroxide (1.0 N), hydrochloric acid (1.0 N), sodium chloride, and Amicon centrifugal filter units with a 10K molecular weight cutoff (MWCO) were purchased from Fisher (Hampton, NH). Nickel chloride hexahydrate was a generous gift from Prof. Andrew Murkin (University at Buffalo, Buffalo, NY). Chelating Sepharose Fast-Flow and Q-Sepharose were purchased from GE Healthcare (Marlborough, MA). Water was purified using a Milli-Q Academic purification system from EMD Millipore (Burlington, MA). All other commercial chemicals were reagent grade or better and were used without further purification.

**Preparation of Wild Type and Variant Yeast Orotidine 5′-Monophosphate Decarboxylases.** The plasmid pScODC-15b containing the gene encoding wild type orotidine 5′-monophosphate decarboxylase from *Saccharomyces cerevisiae* with an N-terminal His$_6$ tag was available from earlier studies. The protein sequence differs from the published sequence for wild type OMPDC by the following substitutions: S2H, C155S, A160S, and N267D. The C155S substitution results in a more stable protein but does not affect the kinetic parameters or overall structure of the enzyme. Site-directed mutagenesis on pScODC-15b was carried out using the QuikChange II kit from Stratagene (San Diego, CA). The following primers, with the bold-face altered codons underlined, were used to prepare the new variant enzymes.

protein loops by interactions with the substrate dianion and the pyrimidine ring.18,21
from wild type OMPDC: T100A, GCT GAC ATT GGT AAT GCA GTC AAA TTG CAG TAC TCT GC; T100G, GCT GAC ATT GGT AAT GGA GTC AAA TTG CAG TAC TCT GCG GG; D37A, C TTT TG TGT TGC TCA TTG GCA GTT CGT ACC ACC AAG GAA TTA CTG G; and D37D, C TTG TGT TGC TCA TTG GCA GTT CGT ACC ACC AAG GAA TTA CTG G.

These variants of OMPDC were overexpressed after the transformation of *Escherichia coli* BL21 (DE3) with the appropriate plasmid. The proteins were purified as described previously. The N-terminal His6 tag was removed in the final step by treatment with thrombin, and the tag was separated from the protein by purification over a column of Q-Sepharose.

Kinetic Parameters for the Decarboxylation of OMP and FOMP. The decarboxylation of OMP was monitored spectrophotometrically by following the decrease in absorbance at 279 nm (0–0.12 mM OMP, Δε = 2400 M cm⁻¹), 290 nm (0.12–0.48 mM OMP, Δε = 1620 M cm⁻¹), and 295 nm (0.48–1.9 mM OMP, Δε = 840 M cm⁻¹) as described in previous work. The decarboxylation of FOMP was monitored spectrophotometrically by following the decrease in absorbance at 282 nm (0.02–0.30 mM FOMP, Δε = 1380 M cm⁻¹), 290 nm (0.30–0.5 mM FOMP, Δε = 1090 M cm⁻¹), 295 nm (0.6 mM FOMP, Δε = 805 M cm⁻¹), and 300 nm (0.9 mM FOMP, Δε = 490 M cm⁻¹) as described in previous work.

The initial velocity, v (M s⁻¹), for the reaction of ≤10% of the substrate at 25 °C, pH 7.1 (30 mM MOPS), and I = 0.105 (NaCl) was determined as the slope of a linear plot of the absorbance vs time for reactions at several different values of [OMP] or [FOMP]. The decarboxylation of OMP that was catalyzed by the T100’ G variant was monitored at 25 °C after mixing 80 μL of a 400 μM stock solution of enzyme variant with 0.920 μL of an assay solution that contained OMP in order to give solutions of 32 μM OMPDC at pH 7.1 (30 mM MOPS) and I = 0.105 (NaCl). Similar procedures were followed at 25 °C in order to monitor the following: (a) the T100’ A ([E] = 270 nM), D37A ([E] = 210 nM), and D37G ([E] = 110 nM) variant-catalyzed decarboxylations of FOMP at pH 7.1 (30 mM MOPS) and I = 0.105 (NaCl) at 279 nm and (b) the D37A ([E] = 60 nM) and D37G ([E] = 56 nM) enzyme variant-catalyzed decarboxylations of FOMP at pH 7.1 (30 mM MOPS) and I = 0.105 (NaCl) at 279 nm. The values of kcat and Km for the variant-catalyzed decarboxylation were determined from the nonlinear least-squares fits of plots of v/[E] (s⁻¹) vs either [OMP] or [FOMP] to the Michaelis–Menten equation.

Apparent first-order rate constants kobs (s⁻¹) for the T100’ A ([E] = 130 nM, monitored at 282 nm) and T100’ G ([E] = 35 μM, monitored at 292 nm) variants of the OMPDC-catalyzed decarboxylations of FOMP at 25 °C, pH 7.1 (30 mM MOPS), and I = 0.105 (NaCl) were determined from the fit to an exponential decay over at least 5 reaction half-life times at [FOMP] ≤ 0.1Km. The second-order rate constant, kcat/Km (M⁻¹ s⁻¹), for the variant OMPDC-catalyzed decarboxylation of FOMP was calculated from the relationship kcat/Km = kobs/[E].

Equilibrium Constants for Converting OMPDC Monomers to the Dimer. The initial velocity, v (M s⁻¹), for the OMPDC-catalyzed decarboxylation of OMP was determined after ≥12-fold dilutions of a stock enzyme solution (0.5–6 μM, wild type OMPDC; 200–400 μM, T100’ G variant; 2.6 μM, T100’ A variant; or 5 μM, D37G variant) in order to give solutions at 25 °C, pH 7.1 (30 mM MOPS), and I = 0.105 (NaCl). The value of [v/E]obs determined over ≤10% of the reaction of total OMP increases with increasing values of [E] = [EM] + 2[ED] when dimeric OMPDC (ED) is in equilibrium with significant concentrations of the monomeric enzyme EM. The association constant Kₐ is defined as eq 1 and Scheme 4 for the conversion of monomeric OMPDC (EM) to the dimer (ED).

\[
K_a = \frac{[ED]}{[EM]^2} = \frac{\frac{v}{[E]_{obs}}}{f_D} = \frac{\frac{v}{[E]_{max}}}{1 + \frac{1 - \sqrt{1 + 8K_a[E]}}{4K_a[E]}}
\]

### RESULTS

The kinetic parameters for the variant OMPDC-catalyzed decarboxylations of OMP were determined at high [OMPDC], where the enzyme exists mainly as the dimer, so that v/[E] (s⁻¹) was independent of [E]. The following are the Michaelis–Menten plots of v/[E] (s⁻¹) against [OMP]: Figure 2 shows the decarboxylation of OMP catalyzed by the 32 μM T100’ G variant and Figures S1A, S1B, and S1C show the decarboxylations of OMP catalyzed by the 110 nM D37G variant, the 210 nM D37A variant, and the 270 nM T100’ A variant, respectively. The values of kcat and Km for the variant OMPDC-catalyzed decarboxylations of OMP, determined from the fits of these plots to the Michaelis–Menten equation, are reported in Table 1. The values of kcat and Km for the D37G (56 nM, Figure S2A) and D37A (60 nM, Figure S2B) variant-catalyzed decarboxylations of FOMP, determined from the fit of plots of v/[E] (s⁻¹) against [FOMP] to the Michaelis–Menten equation, are reported in Table 1.
Figure 2. Dependence of $v/[E]$ for the decarboxylation of OMP catalyzed by the T100′ variant of OMPDC for reactions at 25 °C, pH 7.1 (30 mM MOPS), $I = 0.105$ (NaCl), and 32 μM OMPDC (solid circles). The solid triangles show the limiting values of $v/[E]$ determined for reactions catalyzed by high [OMPDC] at 60 μM (Figure 3A) and 230 μM OMP (Figure 3B).

for >5 reaction half-life times. The apparent first-order rate constants $k_{obs} = 0.57 \pm 0.01$ s$^{-1}$ (average of 3 runs) and $k_{obs} = 0.25 \pm 0.01$ s$^{-1}$ (average of 2 runs) were determined from the fits of the exponential decay curves of absorbance vs time for the decarboxylations catalyzed by the T100′ ([$E$] = 130 nM) and T100′G ([$E$] = 35 μM) variants of OMPDC, respectively.16,18 The second-order rate constants $k_{cat}/K_m ^{G}$ (M$^{-1}$ s$^{-1}$) reported in Table 1 for the variant OMPDC-catalyzed decarboxylations of FOMP were calculated from the relationship $k_{cat}/K_m = k_{obs}/[E]$.

Association Constants for the Dimerization of OMPDC. Figure 3A shows the effect of increasing [E] on $[v/[E]]_{obs}$ for decarboxylation catalyzed by the T100′G variant of OMPDC at 25 °C, pH 7.1 (30 mM MOPS), $I = 0.105$ (NaCl), and 60 μM OMP, where [E] is the total concentration of the OMPDC monomer and $[v/[E]]_{max}$ is the limiting value observed when OMPDC exists essentially exclusively as the active dimer. The increase in $[v/[E]]_{obs}$ with increasing [E] is due to a shift in the position of the dimerization equilibrium ($K_d$, Scheme 4) toward the active dimer. Figure 3B shows the data for the decarboxylation of 230 μM OMP catalyzed by the T100′ G variant. The solid lines for Figure 3A and B show the nonlinear least-squares fits of the experimental data to eq 2 (derived for Scheme 4) using values of $K_d = 1.69 \times 10^5$ and $1.61 \times 10^5$ M$^{-1}$, respectively, and $[v/[E]]_{max} = 0.086 \pm 0.007$ and 0.53 ± 0.05 s$^{-1}$, respectively. The limiting values of $[v/[E]]_{max}$ determined for the decarboxylation of 60 and 230 μM OMP (Figure 3A and B) are shown by the solid triangles on the Michaelis–Menten plots (Figure 2) for the decarboxylation of OMP catalyzed by the T100′ G variant.

Figure 4A–C shows the effect of increasing [E] on $[v/[E]]_{obs}$ for the decarboxylation catalyzed by wild type OMPDC ([$OMP$] = 34 μM), the T100′A variant ([$OMP$] = 99 μM), and the D37G variant ([$OMP$] = 52 μM), respectively, at 25 °C, pH 7.1 (30 mM MOPS), and $I = 0.105$ (NaCl). The solid lines for Figure 4 show the nonlinear least-squares fits of the experimental data to eq 2 (derived from Scheme 4) using values of $K_d$ reported in Table 2 and values of $[v/[E]]_{max} = 23 \pm 2, 2.4 \pm 0.1$, and 3.1 ± 0.6 s$^{-1}$ for the reactions catalyzed by the wild type and T100′A and D37G variants of OMPDC, respectively. The limiting values of $[v/[E]]_{max}$ determined for these variant enzyme-catalyzed decarboxylation reactions are...
shown as solid triangles on the following Michaelis–Menten plots: Figure S1A for the D37G variant ([OMP] = 52 μM) and Figure S1C for the T100′A variant ([OMP] = 99 μM).

**DISCUSSION**

**Variant Enzyme-Catalyzed Decarboxylation of OMP and FOMP.** The values of $k_{cat}/K_m$ for the wild type and variant OMPDC-catalyzed decarboxylations of OMP at 25 °C, pH 7.1 (30 mM MOPS), and $I = 0.105$ (NaCl) reported in Table 1 are ca. 10-fold smaller than the values reported by Wolfenden and co-workers for the decarboxylation in solutions that contain no NaCl. Similar effects have been noted in earlier works. The 4.0 and 2.3 kcal mol$^{-1}$ differences in the activation barriers $\Delta G^\ddagger$ for the decarboxylations catalyzed by the D37A and T100′A variants, respectively, compared to that of wild type OMPDC (Table 1) are similar to the 3.4 and 2.4 kcal mol$^{-1}$ (Table 1) differences from earlier work. Finally, the value for the association constant $K_m = 6 \times 10^{-3}$ M$^{-1}$ reported in Table 2 is 15-fold larger than $K_m = 4 \times 10^{-4}$ M$^{-1}$ for the dimerization of OMPDC at pH 7.2, 0.01 M MOPS buffer, and no NaCl, which was determined by a different method. A similar increase in the value of $K_m$ with an increase in NaCl concentration was noted in ref 24.

X-ray crystal structures for the complexes formed between wild type yeast OMPDC and the tightly bound inhibitors show the D37 side chain from the main subunit hydrogen bonded to the C-3′ ribosyl-hydroxyl, and the T100′ side chain from the second subunit hydrogen bonded to the C-2′-hydroxyl. The contribution of these interactions to the stabilization of the decarboxylation transition state can be estimated by examining the enzyme variants that eliminate the interaction. Table 1 highlights the difficulties associated with this analysis. A comparison of the effects of the Gly and Ala substitutions on $\Delta G^\ddagger$ for the wild type OMPDC-catalyzed decarboxylation of OMP (Table 1) shows that Gly is the more conservative substitution at D37 (ΔΔ$\mu$G = 2.1 and 4.0 kcal mol$^{-1}$ for the D37G and D37A variants, respectively) while Ala is the more conservative substitution at T100′ (ΔΔ$\mu$G = 5.0 and 2.3 kcal mol$^{-1}$ for the T100′G and T100′A variants, respectively). The results are consistent with the conclusion that the D37 and T100′ side chains each contribute ca. 2 kcal mol$^{-1}$ to the total 31 kcal mol$^{-1}$ transition state stabilization.

The decarboxylation of enzyme-bound OMP ($K_m$ (Scheme 4) for the conversion of monomeric wild type and variant OMPDC to the dimer form at 25 °C, pH 7.1 (30 mM MOPS), and $I = 0.105$ (NaCl))

| enzyme | $K_m$ (M$^{-1}$) | $\Delta G^\ddagger$ (kcal mol$^{-1}$)$^{a}$ |
|---------|-----------------|----------------------------------|
| wild type | (5.9 ± 0.6) × 10$^{-6}$ | 3.5 ± 0.2 |
| T100′G | (1.6 ± 0.6) × 10$^{-6}$ | <0.1 ± 0.1 |
| T100′A | (5.7 ± 0.6) × 10$^{-6}$ | 0.9 ± 0.4 |
| D37G | (1.4 ± 0.8) × 10$^{-6}$ | |

The effect of amino acid substitution on $\Delta G^\ddagger$ for the association of OMPDC monomers in order to form the dimer. $^{a}$Determined from the nonlinear least-squares fit of data from Figure 4A to eq 2 (derived from Scheme 4). $^{b}$The quoted uncertainty is the standard deviation from the fitted correlation. $^{c}$The average of the values of $K_m$ determined for the T100′G variant-catalyzed decarboxylation of 60 μM (Figure 3A) and 230 μM (Figure 3B) OMP. $^{d}$Determined from the nonlinear least-squares fit of data from Figure 4B to eq 2 (derived from Scheme 4). $^{e}$Determined from the nonlinear least-squares fit of data from Figure 4C to eq 2 (derived from Scheme 4).

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**Figure 3.** Effect of increasing concentrations of OMPDC on $[v/ [E]]_0$ for the decarboxylation of OMP catalyzed by the T100′G variant at 25 °C, pH 7.1 (30 mM MOPS), and $I = 0.105$ (NaCl). (A) The T100′G variant OMPDC-catalyzed decarboxylation of 60 μM OMP. (B) The T100′G variant-catalyzed decarboxylation of 230 μM OMP.

**Figure 4.** Effect of increasing concentrations of OMPDC on $[v/ [E]]_0$ for the decarboxylation of OMP catalyzed by the wild type and variant enzymes at 25 °C, pH 7.1 (30 mM MOPS), and $I = 0.105$ (NaCl). The solid line shows the fit of the experimental data to eq 2 where [E] is the concentration of the OMPDC monomers. (A) The wild type OMPDC-catalyzed decarboxylation reaction with 34 μM OMP. (B) The T100′A variant-catalyzed decarboxylation reaction of 99 μM OMP. (C) The D37G variant-catalyzed decarboxylation reaction of 52 μM OMP.
partly rate determining for the decarboxylation of OMP.24 The 5-F of FOMP provides a ca. 4 kcal mol\(^{-1}\) stabilization of the vinyl carbanion-like transition state for \(k_{\text{chem}}\) for the OMPDC-catalyzed decarboxylation\(^{16,31}\) but leads to only 1.1- and 6-fold increases in \(k_{\text{cat}}/K_m\) and \(k_{\text{cat}}\), respectively, for the decarboxylation of FOMP. This result shows that the large effect of the strongly electron-withdrawing 5-F group on \(k_{\text{chem}}\) is only weakly expressed at the virtual transition state for \(k_i\) (which is rate determining for \(k_{\text{cat}}/K_m\)) or \(k_{\text{cat}}\) (which is rate determining for \(k_{\text{cat}}\)) for the OMPDC-catalyzed decarboxylation of FOMP.\(^{16}\)

Amino acid substitutions have different effects on the values of \(\Delta G^\dagger\) for the OMPDC-catalyzed decarboxylations of OMP and FOMP when the substitutions cause different changes to the barriers to \(k_{\text{chem}}\) and \((k_i, k_{\text{cat}})\), respectively, that control the activation barriers for the OMPDC-catalyzed decarboxylation of these substrates.\(^{16-18}\) For example, the conservative T100\(\alpha\) and D37G substitutions result in only small 0.5–0.6 kcal mol\(^{-1}\) increases in \(\Delta G^\dagger\) to \(k_{\text{cat}}/K_m\) for the decarboxylation of FOMP and larger 2.1–2.3 kcal mol\(^{-1}\) increases in the barrier to the decarboxylation of OMP (Table 1). These results show that the T100\(\alpha\) and D37G substitutions cause only small changes in the barriers to \(k_i\) and that most of their effect (\(\approx 2\) kcal mol\(^{-1}\)) is on the barrier to \(k_{\text{chem}}\) for the rate determining decarboxylation of the enzyme-bound substrate.

By comparison, the T100\(\alpha\) and D37A substitutions result in 5.0 and 4.0 kcal mol\(^{-1}\) increases, respectively, in the activation barrier \(\Delta G^\dagger\) to \(k_{\text{cat}}/K_m\) for the enzyme-catalyzed decarboxylation of OMP, and in similar 4.4 and 3.1 kcal mol\(^{-1}\) increases in the barrier for the decarboxylation of FOMP (Table 1). This provides strong evidence that the effects of these substitutions are due largely to changes in the barrier to \(k_i\), which is partly rate determining for the OMPDC-catalyzed decarboxylation of OMP and strongly rate determining for the decarboxylation of FOMP.\(^{16}\)

The small range of values of \(k_{\text{cat}}\) determined for the decarboxylation of OMP catalyzed by the four variants (0.5–4.2 s\(^{-1}\)) shows that these substitutions result in similar increases in \(\Delta G^\dagger\) (\(k_{\text{chem}}\)) for the decarboxylation of enzyme-bound OMP. We concluded that the D37A/G and T100\(\alpha\)/A/G substitutions each result in significant increases in \(\Delta G^\dagger\) for the decarboxylation of OMP, and that the larger effects of the D37A and T100\(\alpha\)/G substitutions on the barrier to \(k_{\text{cat}}/K_m\) for the decarboxylation of OMP (Table 1) are due to their additional effects on the barrier to the enzyme conformational change (\(k_i\)), which is partly rate determining for \(k_{\text{cat}}/K_m\). The D37G and D37A substitutions result in small 2-fold changes in the value of \(k_{\text{cat}}\) for the decarboxylation of FOMP. This shows that the substitutions result in only small changes in the barrier to \(k_{\text{cat}}\), which is rate determining for \(k_{\text{cat}}\).

The T100\(\alpha\) side chain lies close to the N-terminal end of the \(\alpha\)-helix (G98–S106, Figure 1).\(^{2}\) Substrate binding induces the movement of this \(\alpha\)-helix toward the ribosyl ring at the main subunit and gives rise to an interaction between the T100\(\alpha\) side chain and the substrate C-2’ ribosyl hydroxyl. Internal Gly side chains are known to destabilize \(\alpha\)-helices relative to Ala,\(^{32,33}\) which is consistent with the proposal that the larger barrier to \(k_i\) for the T100\(\alpha\)/G variant compared to the T100\(\alpha\)/A variant is associated with the effect of the T100\(\alpha\)/G substitution on the helix stability.

![Figure 5](https://dx.doi.org/10.1021/acs.biochem.0c00241)

Figure 5. A pancake representation of the interactions between the amino acid side chains of OMPDC and the bound substrate OMP (PDB 1DQX, but with OMP inserted into the position of the 6-hydroxyuridine 5’-monophosphate inhibitor). The D96’ and T100’ side chains from the second enzyme subunit are shaded red. The D37’ side chain is hydrogen bonded to the C-3’ ribosyl hydroxyl and interacts with the substrate phosphodianion through an intervening water molecule. The interactions of the substrate phosphodianion with the Q215, Y217, and R235 side chains develop during the conformational change from the open unliganded enzyme E0 to the closed Michaelis complex E–C–OMP (Scheme 3).

D37 and the substrate phosphodianion. A major driving force for the large substrate-induced enzyme conformational change is the formation of interactions between the phosphodianion of the substrate and the Q215, Y217, and R235 side chains. We suggest that the D37A and D37G substitutions cause different perturbations in the packing of the water that bridges D37 and the phosphodianion at wild type OMPDC, which are manifested as differences in the effects of these substitutions on \(k_i\) for the enzyme conformational change (Scheme 5).

**Effects of Substitutions on the Stability of Dimeric OMPDC.** OMPDC is a homodimer with an active site that extends across the dimer interface. Most of the side chains that interact with the phosphodianion, ribosyl, and pyrimidine fragments of bound OMP (Figure 5) are located at a single subunit monomer; however, the T100’ and D96’ side chains are located at the second subunit (Figure 5). The dissociation of the dimer to the monomer gives rise to a protein that shows no activity for the decarboxylation of OMP\(^{24}\) so that the enzyme activity \(v/[E]\) is proportional to the fraction of OMPDC present as the dimer (\(f_{\text{dimer}}\) eq 2).


Table 2 reports the values of the association constant $k_{su}$ for the dimerization of OMPDC monomers determined from the fits to eq 2 using data from Figures 3 and 4. The T100′A substitution results in little change in the value of $k_{su}$ determined for wild type OMPDC, while the T100′G substitution results in a 3.5 kcal mol$^{-1}$ destabilization of the active OMPDC dimer relative to the inactive monomer. These results are consistent with a destabilization of the dimeric form of the T100′G variant due to the structural perturbation of the G′98–S′106 α-helix, which lies at the dimer interface. The 300-fold larger $k_{su}$ value for the dissociation of OMP from the T100′G variant compared to wild type OMPDC (Table 1) is also consistent with the large effect of this substitution on protein structure, which results in a decrease in the observed OMP binding energy. The smaller 15-fold effect of the T100′G substitution on the value of $k_{su}$ for the decarboxylation of OMP (Table 1) suggests that OMP binds selectively to the small fraction of OMPDC with the wild type structure for the G′98–S′106 α-helix and that ca. 3 kcal mol$^{-1}$ of the intrinsic OMP binding energy is utilized in order to restore this helix to the wild type conformation at the Michaelis complex to OMP. Additionally, the transition state for the decarboxylation of OMP bound to the T100′G variant is destabilized by 2 kcal mol$^{-1}$ by the loss of the hydrogen bond to the C-2′ ribosyl hydroxyl.

The Partitioning of the OMPDC Active Site between Two Enzyme Subunits. Figure 6A shows the X-ray crystal structure for the open form of yeast OMPDC, but with a hypothetical 6-azauridine S′-monophosphate (azaUMP) ligand at the position observed for the closed liganded enzyme. Figure 6B shows the X-ray crystal structure of the closed form of OMPDC determined for the azaUMP complex, but with only one of the two bound ligands shown. Our previous work highlighted the ligand-driven movement of the phosphodianion gripper loop (P202–V220) toward the pyrimidine umbrella loop (A151–T165) and the clamping interaction between the −CH$_2$OH and amide side chains of S154 and Q215, respectively, which forms a bridge between the two enzyme loops. We now note that the second subunit undergoes a large hinge motion upon ligand binding that is partly or entirely driven by interactions of the G′98–S′106 α-helix with the C-2′ ligand hydroxyl (the T100′ side chain) and the mobile pyrimidine umbrella loop (A151–T165) from the main subunit. In other words, the collective motion of two loops from the main subunit and the G′98–S′106 α-helix at the second subunit is driven by the formation of numerous stabilizing contacts with the bound substrate and intra-subunit interactions between the pyrimidine umbrella loop and the G′98–S′106 α-helix.

Substrate and allosteric binding sites are positioned at the subunit interfaces of phosphofructokinase; this has been linked to the allosteric activation and inhibition of the enzyme-catalyzed phosphorylation of fructose 6-phosphate by ATP in order to form 1,6-fructose diphosphate. We note that OMP binding to the “main” subunit of OMPDC induces the movement of the unoccupied subunit toward the closed enzyme conformation, and that this may result in an increase in the affinity for substrate binding to the second subunit. Such cooperativity in substrate binding has not been reported for OMPDC, but it is not clear that the data are of sufficient quality to rigorously demonstrate a similar affinity for the binding of the first and second substrate. The question of whether the large ligand-driven conformational change of OMPDC, which encompasses the two enzyme subunits, results in cooperativity in the substrate binding has important implications with respect to the mechanism of action of OMPDC and is deserving of further study.

CONCLUSIONS

The effects of the substitutions of D37 and T100′ on the activity of the OMPDC-catalyzed decarboxylations of OMP and FOMP reflect changes in the barriers to the formation of the decarboxylation transition state and to a rate determining enzyme conformational change. The D37A/G and T100′A/G mutations each resulted in a ca. 2 kcal mol$^{-1}$ increase in the reaction barrier, which was proposed to be equal to the stabilization provided by the lost hydrogen bonds to the C-3′ and C-2′ ribosyl hydroxyls, respectively. The D37A and T100′A substitutions resulted in an additional 2–3 kcal mol$^{-1}$ increase in the barrier to a slow enzyme conformational change. These results demonstrate the imperatives for OMPDC to minimize the barriers to both the formation of the decarboxylation transition state and the complex conformational change from the inactive open enzyme $E_{O}$ to the active closed Michaelis complex $E_{O}$OMP (Scheme 5). The T100′G substitution results in a 3.5 kcal mol$^{-1}$ destabilization of dimeric OMPDC relative to the OMPDC monomer. We attribute this to the destabilization of the G′98–S′106 α-helix, which sits at the dimer interface. We propose that an ordered α-helix is also required for the efficient catalysis of the decarboxylation, and that the barrier to restoring the native α-helical structure contributes to the activation barrier for the rate determining conformational change for the T100′G variant.

ASSOCIATED CONTENT

Supporting Information

The Supporting Information is available free of charge at https://pubs.acs.org/doi/10.1021/acs.biochem.0c00241.

Michaels–Menten plots of $v/[E]$ vs [OMP] for the decarboxylation of OMP catalyzed by the D37G, D37A, and T100′A variants of OMPDC (Figures S1A–S1C, respectively) and Michaelis–Menten plots of $v/[E]$ vs...
Accession Codes
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Abbreviations
OMPDC, orotidine 5′-monophosphate decarboxylase; OMP, orotidine 5′-monophosphate; UMP, uridine 5′-monophosphate; FOMP, 5-fluoroorotidine 5′-monophosphate; FUMP, 5-fluorouridine 5′-monophosphate; MOPS, 3-(N-morpholino)propanesulfonic acid; GlyGly, glycyl glycine

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