Rate and Equilibrium Constants for an Enzyme Conformational Change during Catalysis by Orotidine 5′-Monophosphate Decarboxylase

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ABSTRACT: The caged complex between orotidine 5′-monophosphate decarboxylase (ScOMPDC) and 5′-fluoroorotidine 5′-monophosphate (FOMP) undergoes decarboxylation ~300 times faster than the caged complex between ScOMPDC and the physiological substrate, orotidine 5′-monophosphate (OMP). Consequently, the enzyme conformational changes required to lock FOMP at a protein cage and release product 5′-fluorouridine 5′-monophosphate (FUMP) are kinetically significant steps. The caged form of ScOMPDC is stabilized by interactions between the side chains from Gln215, Tyr217, and Arg235 and the substrate phosphodianion. The control of these interactions over the barrier to the binding of FOMP and the release of FUMP was probed by determining the effect of all combinations of single, double, and triple Q215A, Y217F, and R235A mutations on \( k_{\text{cat}}/K_m \) and \( k_{\text{cat}} \) for turnover of FOMP by wild-type ScOMPDC; its values are limited by the rates of substrate binding and product release, respectively. The Q215A and Y217F mutations each result in an increase in \( k_{\text{cat}}/K_m \) and a decrease in \( k_{\text{cat}} \) due to a weakening of the protein–phosphodianion interactions that favor fast product release and slow substrate binding. The Q215A/R235A mutation causes a large decrease in the kinetic parameters for ScOMPDC-catalyzed decarboxylation of OMP, which are limited by the rate of the decarboxylation step, but much smaller decreases in the kinetic parameters for ScOMPDC-catalyzed decarboxylation of FOMP, which are limited by the rate of enzyme conformational changes. By contrast, the Y217A mutation results in large decreases in \( k_{\text{cat}}/K_m \) for ScOMPDC-catalyzed decarboxylation of both OMP and FOMP, because of the comparable effects of this mutation on rate-determining decarboxylation of enzyme-bound OMP and on the rate-determining enzyme conformational change for decarboxylation of FOMP. We propose that \( k_{\text{cat}} = 8.2 \text{ s}^{-1} \) for decarboxylation of FOMP by the Y217A mutant is equal to the rate constant for cage formation from the complex between FOMP and the open enzyme, that the tyrosyl phenol group stabilizes the closed form of ScOMPDC by hydrogen bonding to the substrate phosphodianion, and that the phenyl group of Y217 and F217 facilitates formation of the transition state for the rate-limiting conformational change. An analysis of kinetic data for mutant enzyme-catalyzed decarboxylation of OMP and FOMP provides estimates for the rate and equilibrium constants for the conformational change that traps FOMP at the enzyme active site.

Orotidine 5′-monophosphate decarboxylase from Saccharomyces cerevisiae (ScOMPDC) provides a large 31 kcal/mol stabilization of the transition state for decarboxylation of OMP to form uridine 5′-monophosphate [UMP (Scheme 1)], through a UMP carbanion intermediate, and shows an extraordinary specificity in binding this transition state with an affinity much higher than that of OMP, whose ground-state Michaelis complex is stabilized by only 8 kcal/mol. This rate acceleration is achieved by sequestering OMP in a structured protein cage, which provides for optimal stabilizing interactions with the decarboxylation transition state. The protein cage is formed by an energetically demanding conformational change, driven by the binding interactions with the substrate phosphodianion and other nonreacting substrate fragments, which function to mold two flexible protein loops into an active enzyme (Figure 1).
decarboxylation of OMP and the phosphodianion-truncated substrate 1-(β-D-erythrofuranosyl)orotic acid (EO).\textsuperscript{4,19} The single mutations result in ≤2.5-fold decreases in $k_{\text{cat}}/K_m$ for the catalyzed reaction of EO.\textsuperscript{4} By contrast, these mutations result in similar large decreases in $k_{\text{cat}}/K_m$ for ScOMPDC-catalyzed decarboxylation of OMP,\textsuperscript{19} and in the third-order rate constant for activation of ScOMPDC-catalyzed decarboxylation of EO by phosphite dianion.\textsuperscript{4} These results show that interactions of ScOMPDC with the phosphodianion of OMP, or with phosphite dianion have the sole function of activating the enzyme for catalysis of decarboxylation at the distant orotate ring, through stabilization of a catalytically active caged substrate complex.\textsuperscript{4,10,11,19} This model also provides a simple rationalization for the activation by phosphite dianion of triosephosphate isomerase\textsuperscript{20−23} and L-glycerol 3-phosphate dehydrogenase-catalyzed reactions\textsuperscript{24−26} of their phosphodianion-truncated substrate.

In this paper, we consider the kinetic barriers for conversion of the open form of ScOMPDC to the caged complex to substrate (Figure 1), and for conversion of the caged product complex to the open enzyme. Rate constants for conformational changes that convert enzymes from an inactive to an active form may be determined directly as the appropriate

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**Scheme 1**

\[
\begin{align*}
\text{OMP} & \rightarrow \text{UMP} \\
2\text{O}_2\text{P} & \rightarrow \text{H}_2\text{O}
\end{align*}
\]

Figure 1. Space filling models of ScOMPDC from yeast.\textsuperscript{18} The structure on the left is the open unliganded form of ScOMPDC (PDB entry 1DQW), and the structure on the right shows the complex with 6-hydroxyuridine S'-monophosphate (PDB entry 1DQX). Two colored flexible loops close to form the active site cage: Pro202−Val220, on the left-hand side of each structure, interact with the substrate dianion, and Glu152−Thr165, on the right-hand side, interact with the pyrimidine ring. The guanidine side chain of R235, at the base of the left-hand loop, is colored green.

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**Scheme 2**

\[
\begin{align*}
\text{Y} & = \text{CH}_3\text{OPO}_{2}^- \\
X & = \text{H: OMP} \\
X & = \text{F: FOMP} \\
\end{align*}
\]

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Figure 2. X-ray crystal structure (PDB entry 1DQX) of yeast ScOMPDC in a complex with 6-hydroxyuridine S'-monophosphate.\textsuperscript{18} This structure shows the important interactions of Gln215, Tyr217, and Arg235 side chains from the phosphodianion gripper loop with the ligand phosphodianion. Reproduced from ref 4. Copyright 2012 American Chemical Society.
enzyme kinetic parameter, when the conformational change is rate-determining for catalysis. Substrate binding is already partly rate-determining for ScOMPDC-catalyzed decarboxylation of OMP,\textsuperscript{9,27} while the S-F substituent of 5-fluorouridine 5'-monophosphate (FOMP) is exceptional in providing strong stabilization of transition states for ScOMPDC-catalyzed reactions.\textsuperscript{27,28} For example, the S-F results in a 3400-fold increase in $k_{cat}/K_m$ for ScOMPDC-catalyzed decarboxylation of truncated substrate FOMP.\textsuperscript{27,28} Now, if enzyme conformational changes are partly rate-determining for ScOMPDC-catalyzed decarboxylation of OMP, then these steps should be strongly rate-determining for chemically rapid ScOMPDC-catalyzed decarboxylation of FOMP.\textsuperscript{27,28}

The S-F at FOMP results in only small 1.1- and 6-fold increases in $k_{cat}/K_m$ and $k_{cat}$ respectively, for decarboxylation of OMP (Scheme 2),\textsuperscript{29} so that the S-F substituent effect is only weakly expressed at the virtual transition states that govern the values of these kinetic parameters.\textsuperscript{31} Instead, the value of $k_{cat}/K_m$ is mainly limited by the rate of formation of the Michaelis complex, and the value of $k_{cat}$ is mainly limited by the rate of product release.\textsuperscript{27} The values of the kinetic parameter for decarboxylation of FOMP are not strongly affected by changes in solvent viscosity $\eta$, so that substrate binding and product release are not strictly diffusion-controlled reactions.\textsuperscript{27} It was proposed that $k_{cat}/K_m$ is instead limited by the rate of the protein conformational changes that follow substrate binding.\textsuperscript{27}

Roughly, a 390/1.1 = 350-fold increase (Scheme 2) in the observed rate constant of $k_{cat}/K_m$ for decarboxylation of FOMP, relative to that for decarboxylation of OMP, is required to effect similar rate-determining decarboxylation steps for the catalyzed reactions of both OMP and FOMP.

A comparison of the effects of site-directed mutations on the kinetic parameters for ScOMPDC-catalyzed decarboxylation of OMP, which is limited by chemical decarboxylation, and on ScOMPDC-catalyzed decarboxylation of FOMP, which is limited by the rate of formation and breakdown of Michaelis complexes, provides insight into the effect of the mutations on the relative barriers to the chemical steps and the enzyme conformational changes.\textsuperscript{32} We previously reported the effect of single (Q215A, R235F, and Y217F), double (Q215A/Y217F, Q215A/R235A, and R235A/Y217F), and triple (Q215A/R235A/Y217F) mutations of amino acid residues that interact with the phosphodiadion of OMP (Figure 2) on the kinetic parameters for ScOMPDC-catalyzed decarboxylation of OMP.\textsuperscript{4,19} We report here the effect of these same mutations on the kinetic parameters for ScOMPDC-catalyzed decarboxylation of FOMP. We note that these mutations generally cause larger changes in the values of the kinetic parameters for decarboxylation of OMP than of FOMP. This is because of their larger effects on the barrier to the decarboxylation reaction, which limits the rate of decarboxylation OMP, compared with their effect on the barrier to the enzyme conformational change, which limits the rate of decarboxylation of FOMP.\textsuperscript{27} We also note interesting exceptions to this trend, which show that mutations of gripper side chains cause significant changes in the barriers to the protein conformational changes, which control the rate constants for formation and breakdown of complexes between ScOMPDC and FOMP or FUMP.
FUMP were obtained by fitting the absorbance date to the equation for an exponential decay. The value of $k_{\text{cat}}/K_m$ (M$^{-1}$s$^{-1}$) was then calculated from the relationship $k_{\text{cat}}/K_m = k_{\text{obsd}}/[E]$.

**RESULTS**

Panels A–C of Figure 3 show the fits of plots of $v_0/[E]$ versus [FOMP] to the Michaelis–Menten equation for decarboxylation of FOMP catalyzed by mutant forms of ScOMPDC in solutions that contain 30 mM MOPS (50% free base) at pH 7.1, 25 °C, and $I = 0.105$ (NaCl).37 By comparison, a $k_{\text{cat}}$ of 2.0 s$^{-1}$, a $K_m$ of 9.4 × 10$^{-5}$ M$^{-1}$, and a $k_{\text{cat}}/K_m$ of 21000 M$^{-1}$ s$^{-1}$ were reported for the reaction catalyzed by this mutant at an undefined low ionic strength.38 A decrease in ionic strength from $I = 0.105$ to $I = 0.020$ was shown in earlier work to result in a 9-fold decrease in $K_m$ and $k_{\text{cat}}/K_m$ for the decarboxylation of OMP catalyzed by the R235A mutant of ScOMPDC.29

**DISCUSSION**

An enzyme conformational change at the ternary E-FEO-HPO$_3^-$ complex is rate-determining for phosphite dianion activation of ScOMPDC-catalyzed decarboxylation of the fluorinated—truncated substrate FEO to form FEU.30 The 1.1-fold effect of the S-F at the whole substrate FOMP on $k_{\text{cat}}/K_m$ for wild-type ScOMPDC-catalyzed decarboxylation of OMP shows that there is minimal stabilization of this rate-determining transition state by polar/electrostatic interactions between the electronegative -F and transition-state negative charge at C-6. The simplest interpretation is that $k_{\text{cat}}/K_m$ is limited by the rate of diffusion-controlled formation of the Michaelis complex between ScOMPDC and FOMP. However, a study of the effect of changes in solvent viscosity ($\eta_o$) on kinetic parameter ($k_{\text{cat}}/K_m$)$_o$ for decarboxylation in water gave slopes of 0.37 and 0.64 for plots of ($k_{\text{cat}}/K_m$)$_o$/$(k_{\text{cat}}/K_m)_d$ for decarboxylation of OMP and FOMP, respectively, against $\eta_o/\eta_0$ which are smaller than the slope of 1.0 expected for a diffusion-limited reaction.39 This shows that the rate of formation of the Michaelis complex is partly limited by a second step, which we propose is the conformational change that traps FOMP in a protein cage [$k_1$ (Scheme 3)].27

The small value of ($k_{\text{cat}}$)$_E$/$(k_{\text{cat}})_H = 6$ for ScOMPDC-catalyzed decarboxylation of OMP and FOMP, compared to ratios observed when the entire S-F substituent effect on carbanion stability is expressed at the rate-determining transition state (Scheme 2), shows that decarboxylation of FOMP is not limited by rate constant $k_{\text{chem}}$ (Scheme 3) for decarboxylation of bound substrate. Slopes of 0.39 and 1.0, respectively, were determined for plots of ($k_{\text{cat}}$)$_{obsd}$/$(k_{\text{cat}})_E$ for decarboxylation of OMP and FOMP, respectively, versus $\eta_o/\eta_0$. These are consistent with the conclusion that $k_1$ and/or $k'_{-c\text{-d}}$ are partly rate-determining for ScOMPDC-catalyzed decarboxylation of OMP (Scheme 3) and fully rate-determining for decarboxylation of FOMP.

**Simple Conclusions about the Effect of Mutations on S-F Substituent Effects.** The effect of the S-F at FOMP on the kinetic parameters $k_{\text{cat}}/K_m$ and $k_{\text{cat}}$ for ScOMPDC-catalyzed decarboxylation of OMP, when chemical decarboxylation is clearly rate-determining, should be similar to the 390-fold effect of S-F on $k_{\text{cat}}/K_m$ for ScOMPDC-catalyzed decarboxylation of the phosphodianion-truncated substrate EO (Scheme 2). Table 1 shows that the values of ($k_{\text{cat}}$)$_E$/$(k_{\text{cat}})_H$ increase as ScOMPDC is crippled by mutations of gripper side chains to 760 for the Q215A/Y217A/R235A triple mutant, consistent with rate-determining decarboxylation for the two substrates. The largest observed ratio of ($k_{\text{cat}}$)$_E$/$(k_{\text{cat}})_H$ is 240 for the Q215A/R235A mutant enzyme-catalyzed reactions. The more severely crippled mutants bind OMP and FOMP too weakly to approach saturation.
Table 1. Effect of Mutations of Phosphoribosylglyceri Amino Acid Residues on the Kinetic Parameters for ScOMPDC-Catalyzed Decarboxylation of FOMP

| ScOMPDC       | k_{cat} (s^{-1})^{b} | (k_{cat})_{f}/(k_{cat})_{t}^{b} | K_{m} (M)^{b} | k_{cat}/K_{m} (M^{-1} s^{-1}) | (k_{cat}/K_{m})_{f}/(k_{cat}/K_{m})_{t}^{b} |
|---------------|----------------------|---------------------------------|---------------|-----------------------------|------------------------------------------|
| wild-type     | 95                   | 6                               | 8.0 × 10^{-6} | 1.2 × 10^{5}                | 1.1                                       |
| Q215A^{d}     | 190 ± 10             | 8                               | (9.6 ± 1) × 10^{-5} | 2.0 × 10^{4} | 8                          |
| Y217F^{d}     | 430 ± 30             | 21                              | (4.2 ± 0.6) × 10^{-4} | 1.1 × 10^{6} | 6                          |
| R235^{d}      | 92                   | 92                              | 5.8 × 10^{-4}  | 1.6 × 10^{5}                | 180                                       |
| Q215A/Y217F^{d} | 49 ± 5               | 10                              | (9.2 ± 1.7) × 10^{-4} | 5.3 × 10^{4} | 16                         |
| Q215A/R235A^{d} | 4.7 ± 0.3            | 240                             | (6.5 ± 0.8) × 10^{-4} | 7200             | 500                        |
| Y217F/R235A^{d} | triple mutant        | 28                              |                |                            |                                           |
| S154A^{d}     | 16.0 ± 5             | 200                             | (5.5 ± 0.6) × 10^{-5} | 2.9 × 10^{4} | 460^{d}                    |
| S154A/Q215A^{d} | 6.6 ± 0.3            | 160                             | (8.6 ± 1.3) × 10^{-5} | 7.7 × 10^{4} | 200^{d}                    |
| Y217A^{d}     | 8.2 ± 0.5            | 26                              | (6.2 ± 0.6) × 10^{-4} | 1.3 × 10^{4} | 50^{d}                     |

*For reactions at pH 7.1 (30 mM MOPS), 25 °C, and I = 0.105 (NaCl).^{a} The quoted errors are the standard deviations obtained from the nonlinear least-squares fits of data from panels A–C of Figure 3 to the Michaelis–Menten equation. From ref 29. Figure 3A. Figure 3B. Figure 3C. Kinetic parameters for reactions of OMP from ref 37. Kinetic parameters for reactions of OMP from ref 19, unless indicated otherwise. Kinetic parameters for reactions of OMP from ref 34.

Scheme 3

Single mutations of gripper side chains result in increases in k_{cat}/K_{m} for kinetic parameters k_{cat} and k_{cat}/K_{m} because of the stronger effect of the mutations on k_{cat} and k_{cat}/K_{m} for enzyme-catalyzed decarboxylation of OMP. The following effects of Q215A and Y217F single mutations on k_{cat}/K_{m} for kinetic parameters k_{cat}/K_{m} and k_{cat} provide strong evidence that the mutations affect the rate constants for binding and release of FOMP.

1. The Q215A and Y217F mutations of ScOMPDC cause (k_{cat}/K_{m})_{P}/(k_{cat}/K_{m})_{H} to increase to 8 and 6, respectively. These are still much smaller than the ratio of ≈400 for rate-determining decarboxylation. Therefore, formation of the E_{c}/FOMP complex remains rate-determining for the mutant enzyme-catalyzed reactions. The mutations result in surprising 6- and 10-fold decreases in k_{cat}/K_{m} for Q215A and Y217F mutant enzyme-catalyzed decarboxylation of FOMP, respectively (Table 1), which can reflect only decreases in k_{cat} for rate-determining conversion of E_{c}/FOMP to E_{c}/FOMP (Scheme 3). This shows that a weakening of the protein–phosphoribosylglyceri interactions by these mutations results in a decrease in the rate of enzyme conformational change, which allows expression of these interactions (Figure 2).

2. The Q215A and Y217F mutations of ScOMPDC cause an increase in (k_{cat})_{P}/(k_{cat})_{H} which is due mainly to a surprising increase in k_{cat} for decarboxylation of FOMP by rate-determining breakdown of E_{c}/FOMP (Scheme 3). We conclude that k_{cat} ≈ K_{cat} = 95 s^{-1} (Scheme 3) for the wild-type enzyme-catalyzed reaction and that weakening of the protein–dianion interactions results in an increase in k_{cat} (Scheme 3) for the rate-determining enzyme conformational change.

The S154A mutation results in large 180- and 15000-fold decreases in kinetic parameters k_{cat} and (k_{cat})_{f}/(k_{cat})_{t}, respectively, for decarboxylation of OMP but much smaller 6- and 41-fold decreases in k_{cat} and (k_{cat})_{f}/(k_{cat})_{t}, respectively, for decarboxylation of FOMP (Table 1). Consequently, the S154A mutant shows a high reactivity toward catalysis of decarboxylation of FOMP: (k_{cat}/K_{m})_{P}/(k_{cat}/K_{m})_{H} = 460, and (k_{cat})_{f}/(k_{cat})_{t} = 200. These S-F effects are similar to the 400-fold effect of the S-F on k_{cat}/K_{m} for decarboxylation of the phosphoribosylglyceri–truncated substrate EO (Scheme 2). We conclude that the stabilizing interactions between the S-F and neighboring C-6 carbanion are strongly expressed at the rate-determining transition state for decarboxylation of FOMP catalyzed by S154A mutant ScOMPDC. We suggest that the ~2-fold larger S-F substituent effect on k_{cat}/K_{m} compared with that on k_{cat} reflects the small stabilization of the Michaelis complex to FOMP by an interaction between the protein and the S-F substituent.

The CH_{3}OH side chain of Ser154 accepts a hydrogen bond from the pyrimidine NH group and acts as a hydrogen bond donor to the oxygen of the amide side chain of Gln215 (Figure 4).^{18,34} The second H-bond acts to clamp two enzyme loops over the pyrimidine ring and the phosphoribosylglyceri of OMP. The effect of these individual mutations on the kinetic parameters for decarboxylation of OMP depends upon the order of the amino acid substitutions.^{34} The 6 kcal/mol total side chain

![Phosphoribosylglyceri Gripper Loop](image)

![Pyrimidine Umbrella](image)

Figure 4. Partial X-ray crystal structure of ScOMPDC complexed with 6-hydroxypyrimidine 5′-monophosphate (PDB entry 1DQX) superimposed over the structure for unliganded ScOMPDC (PDB entry 1DQW). The movement of the phosphoribosyl gripper loop (Pro202–Val220) toward the hydrophobic pyrimidine umbrella (Glu152–Thr165) is shown for the unliganded (olive green) and liganded (orange) enzymes. The closure of these loops is cooperative and driven by the formation of a hydrogen bond between the side chains of Gln215 from the “gripper” loop and of Ser 154 from the “umbrella.” Reproduced from ref 29. Copyright 2013 American Chemical Society.
interactions can be partitioned into an ∼2 kcal/mol interaction between the amide side chain of Gln215 and the phosphodianion group of OMP, observed for the Q215A mutation of wild-type ScOMPDC, and an ∼4 kcal/mol interaction between the -CH₂OH side chain of Ser154 and the pyrimidine ring of OMP, observed for the S154A mutation of Q215A mutant ScOMPDC. The opposite order of these mutations leads to expression of nearly the entire 6 kcal/mol effect observed for the S154A/Q215A double mutation at the S154A single mutant, with only a small 0.3 kcal/mol additional effect of the second Q215A mutation.34 These results reflect cooperativity in the interactions of these side chains with the transition state for decarboxylation, due to the requirement for a hydrogen bond between the -CH₂OH group of Ser154 and the amide side chain of Gln215 to position the latter side chain to interact with the phosphodianion group of OMP. It was proposed that the S154A single mutation leads to loss of the transition-state stabilization contributed by the side chains of both Ser154 and Gln215. These trends are hardly discernible in the effects of the same mutations on the kinetic parameters for ScOMPDC-catalyzed decarboxylation of FOMP, because the large effects on the rate of chemical decarboxylation are weakly expressed as changes in the enzyme kinetic parameters.

**Eliminating Protein–Phosphodianion Interactions.**

The results from a recent study on the effects of mutations of gripper side chains on the kinetic parameters for ScOMPDC-catalyzed decarboxylation of EO and OMP provide strong evidence that the effects of these mutations on the enzyme kinetic parameters for decarboxylation of OMP are due essentially entirely to changes in the relative stability of the inactive open and active closed forms of ScOMPDC (Scheme 4).4,19,20 We will summarize the experimental evidence that supports the model shown in Scheme 4 and then use this model in the interpretation of the effects of these mutations on the kinetic parameters for ScOMPDC-catalyzed decarboxylation of FOMP.

(1) The Q215A, Y217A, and R235A single mutations result in ≥2.5-fold decreases in $k_{cat}/K_a$ for decarboxylation of the phosphodianion-truncated substrate EO,4 but in much larger decreases in the third-order rate constant for phosphodianion activation of ScOMPDC-catalyzed decarboxylation of EO,4 and in the second-order rate constant $k_{cat}/K_m$ for decarboxylation of OMP.19 The first result shows that there is little or no direct interaction between the loop side chains and the site of chemical decarboxylation of the orotate ring. It was concluded that the large effect of these mutations on the kinetic parameters for dianion activation of decarboxylation of EO and for decarboxylation of OMP is entirely due to the loss of stabilizing interactions between the deleted side chains and the enzyme-bound dianion.4,19

(2) The model in Scheme 4 was used to rationalize the effects of mutations of gripper side chains on the kinetic parameters for ScOMPDC-catalyzed decarboxylation of OMP. The complex between wild-type ScOMPDC and OMP is proposed to exist mainly in the closed form $[K_a \gg 1 \text{(Scheme 4A)}]$. Consequently, the first mutations of gripper side chains, which result in up to 4 kcal/mol destabilization of $E_0$, are entirely expressed as a decrease in the Michaelis constant to the limiting value of $(K_m)_\text{EO} = K_a = 1 \text{ mM}$ for dissociation of OMP from $E_0\cdot\text{OMP}$. Mutations that result in further decreases to $K_a$ values of <1 then lead to a decrease in $k_{cat}$ which falls below the invariant value of $k_{cat}$ for ScOMPDC-catalyzed decarboxylation of OMP (Scheme 3).28 This is because the loop dianion interactions now only develop on proceeding from the dominant Michaelis complex ($E_0\cdot\text{OMP}$) to the transition state for decarboxylation at $E_0\cdot\text{OMP}$ [ES$^\ddagger$ (Scheme 4B)]. We estimate a $K_a$ value of ≈1000 (Table 2) for the reaction catalyzed by wild-type ScOMPDC from the difference in $K_a = 1 \mu M$ for decarboxylation of OMP and estimated $K_a \approx 1 \mu M$ for dissociation of OMP from $E_0\cdot\text{OMP}$.19

**Microscopic Rate Constants. Wild-Type ScOMPDC.**

Substrate binding ($k_{cat}$) and enzyme conformational change $k_{cat}$ are each partly rate-determining for wild-type ScOMPDC-catalyzed decarboxylation of FOMP at [FOMP] $\ll K_m$ so that $k_{cat} \approx k_f$ (Scheme 5).27 Combining the values of $(k_{cat}/K_m) \approx k_f = 1.2 \times 10^7 \text{ M}^{-1} \text{s}^{-1}$ for encounter-limited decarboxylation of FOMP determined for wild-type OMPDC and $K_a = (k_{cat}/K_m) \approx 10^{−3} \text{ M} \text{ (see above)}$ for breakdown of the $E_0\cdot\text{FOMP}$

**Table 2. Microscopic Rate Constants for Decarboxylation of FOMP (Scheme 5) by Wild-Type and Mutant Forms of ScOMPDC at pH 7.1 (30 mM MOPS), 25 °C, and I = 0.105 (NaCl), Calculated As Described in the Text.**

| ScOMPDC | $k_{cat}$ (s$^{-1}$) | $k_{cat}$ (s$^{-1}$) | $[(k_{cat}/K_m)_{FOMP}]$ | $K_d^d$ | $k_f$ (s$^{-1}$) | $k_{cat}$ (s$^{-1}$) | $K_a^d$ | $A^e$ (kcal/mol) | $k_f$ (s$^{-1}$) |
|---------|-------------------|-------------------|---------------------------|--------|----------------|-------------------|--------|----------------|----------------|
| wild-type | 95 | 5000 | 1000 | 10$^4$ | 10 | 13.9 | 350 | |
| R235A | 92 | 5000 | 15 | 0.071 | 120 | 1800 | 13.7 | 500 |
| Q215A/R235A | 4.7 | 5000 | 750 | 0.0013 | 17 | 13000 | 13.7 | 560 |
| Q215A/Y217F | 49 | 5000 | 3.1 | 0.48 | 50 | 100 | 14.9 | 70 |
| Y217A | 15 | 5000 | 3.2 | 0.45 | 8.2 | 18 | 15.9 | 12 |

$^a$Table 1.4 Calculated from $k_{cat} = 15 \text{ s}^{-1}$ for decarboxylation of OMP and the ratios $k_{cat}/K_m = 1.8$ for decarboxylation of OMP, where $k_{cat}$ is the true decarboxylation rate constant, and $(k_{cat}/K_m)_FOMP \approx 200$ (Table 1) when chemistry is rate-determining for ScOMPDC-catalyzed decarboxylation of OMP and FOMP (Table 1). $^b$The ratio of observed rate constants $k_{cat}$ for decarboxylation of OMP catalyzed by wild-type $[(k_{cat})_{wild}]$ and mutant $[(k_{cat})_{mutant}]$ forms of ScOMPDC. $^c$Estimated equilibrium constants for conversion of $E_0\cdot\text{OMP}$ to $E_0\cdot\text{OMP}$ (Scheme 5), calculated for mutants of ScOMPDC using eq 1. $^d$Rate constant for conversion of $E_0\cdot\text{OMP}$ to $E_0\cdot\text{OMP}$, calculated using eqs 2 and 3. $^e$Rate constant for conversion of $E_0\cdot\text{OMP}$ to $E_0\cdot\text{OMP}$. $^f$Marcus intrinsic reaction barrier, calculated using eq 5.
complex gives $k_i \approx k_{-i} = 10^5$ s$^{-1}$ for the enzyme conformational change (Table 2). The value of $k_{chem} = 5000$ s$^{-1}$ for decarboxylation at $E_C$-FOMP by wild-type ScOMPDC (Table 2) was calculated from $k_{cat} = 15$ s$^{-1}$ for decarboxylation of OMP, and the ratios $k_{chem}/k_{cat} = 1.8$ for decarboxylation of OMP, where $k_{chem}$ is the true decarboxylation rate constant, and $(k_{cat})_{R}/(k_{cat})_{H} \approx 200$ when chemistry (Table 1) is rate-determining for OMPDC-catalyzed reactions of both substrates.

**Mutants of ScOMPDC.** The Q215A and Y217F mutations each cause $k_{cat}$ for turnover of FOMP to increase above the value of $k_{cat} = 95$ s$^{-1}$ for wild-type ScOMPDC, because of an increase in the rate of release of product to water (Table 1). By contrast, the more severe R235A, Q215A/Y217F, and Q215A/R235A mutations cause $k_{cat}$ to decrease to a value as small as 4.7 s$^{-1}$ for the Q215A/R235A mutant (Table 1). The value of $k_{chem}$ for the reaction of $E_C$-FOMP is not affected by mutations of the gripper residues, because there are only weak interactions between these side chains and the orotate ring at the transition state for ScOMPDC-catalyzed decarboxylation of enzyme-bound substrate. Rather, the decreases in $k_{cat}$ as the protein dianion interactions are weakened and product release becomes much faster than turnover, show that the values of rate constants $k_{cat}$, $k_{-cat}$ and $k_{chem}$ (Scheme 5, which omits the step for product release) begin to control the observed value of $k_{cat}$.

Equation 1 relates the equilibrium constant $K_c$ for loop closure at mutant ScOMPDCs to the effect of the individual mutation on $(k_{cat})_{mut}$ for wild-type ScOMPDC-catalyzed decarboxylation of OMP. This equation was derived by assuming that these mutations do not affect microscopic rate constant $k_{chem}$ for decarboxylation of $E_C$-OMP, and that the entire decrease in $(k_{cat})_{mut}$ compared to $(k_{cat})_{wt}$ is due to the decrease, from the value of $(k_{cat})_{mut} = (k_{cat})_{wt}$. The estimated values of $K_c$ reported in Table 2 for mutant ScOMPDC-catalyzed decarboxylation of FOMP were calculated using eq (1) (Scheme 3), and assuming identical values of $K_c$ for loop closure over OMP and FOMP.

$$1/(K_c)_{mut} = (k_{cat})_{wt}/(k_{cat})_{mut} - 1$$  

(1)

Table 2 reports estimated rate constants (Scheme 5) for mutants of ScOMPDC, which were calculated from $k_{chem} = 5000$ s$^{-1}$, the values of $(k_{cat})_{mut}$ for decarboxylation of FOMP (Table 1) and of $K_c$ for the enzyme conformational change (see above), and using eqs 2 and 3 derived for Scheme 5. We note again this treatment assumes that mutations of gripper side chains do not affect the value of $k_{chem}$ (Scheme 5) for decarboxylation of OMP and FOMP. The observation that this treatment provides a rationalization for the following otherwise confusing effects of the mutations of gripper side chains on the kinetic parameters for decarboxylation of OMP and FOMP provides strong justification for the simplifying assumptions used in the calculation of the rate constants from Table 2.

(2) The large value of $(k_{cat})_{F}/(k_{cat})_{H} = 240$ for decarboxylation catalyzed by Q215A/R235A mutant ScOMPDC shows that the step for $k_{chem}$ for decarboxylation of $E_C$-FOMP limits the value of $k_{cat}$ for turnover (Scheme 3). The $k_{cat}$ value of 4.7 s$^{-1}$ for decarboxylation of FOMP catalyzed by this mutant is much smaller than the $k_{chem}$ of $\approx 5000$ s$^{-1}$ and the $k_{cat}$ of $= 17$ s$^{-1}$ (Table 2), because the barrier to decarboxylation $k_{cat}$ includes the $\sim 4$ kcal/mol barrier for conversion of $E_O$-FOMP to $E_C$-FOMP $[K_c = 1/750, k_{cat} = 13000$ s$^{-1}$) (Table 2)]. In other words, loop closure over enzyme-bound FOMP is partly reversible $(k_{-cat} > k_{chem})$ so that decarboxylation at $E_C$-FOMP $(k_{chem})$ is the rate-determining step.

(3) The Y217A mutation results in similar large decreases in $k_{cat}$ and $k_{cat}/K_m$ for wild-type ScOMPDC-catalyzed decarboxylation of both OMP and FOMP, so that there is only a small effect of the S-F substituent on the kinetic parameters for this mutant enzyme-catalyzed reaction: $(k_{cat})_{F}/(k_{cat})_{H} = 2$, and $(k_{cat})_{F}/(k_{cat})_{H} = 5$ (Table 1). We conclude that the Y217A mutation results in a similar increase in the barriers to decarboxylation of OMP, where chemistry is rate-determining, and for decarboxylation of FOMP, where the enzyme conformational change is rate-determining. This is reflected by the large decrease in rate constant $k_{cat}$ for the enzyme conformational change (Scheme 5) to form the loop-closed ScOMPDC (Table 2).

$$k_{cat} = k_{cat}/K_m = (1/(K_c)_{mut}) \times (k_{cat})_{wt}/((k_{-cat})_{mut})$$

(4) Equations 2 and 4 are derived for mutants (Table 2) whose reaction is described by Scheme 5. These equations predict that kinetic parameters $k_{cat}$ and $k_{cat}/K_m$ for decarboxylation of FOMP are related by $1/K_c$ for formation of $E_O$-FOMP. This provides a rationalization for the observation that the kinetic data from Table 1 can be used to calculate a nearly constant ratio of $k_{cat}/(k_{cat}/K_m) \approx K_c = 6-9 \times 10^{-4}$ M for decarboxylation of FOMP catalyzed by the different mutants from Table 2, which is similar to $K_c \approx 10^{-3}$ M estimated in earlier work for the release of substrate from $E_O$-OMP to form $E_O$. 

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Opening at the crippled enzyme, relative to the barrier for formation of EC decarboxylation show that the step for chemical decarboxylation of OMP.

− estimated for the Q215A/R235A [100 s−1] (Scheme 5) as the reaction barrier in the absence of a change for the Y217A mutant is slow because of the large unfavorable thermodynamic activation barrier to conversion of EO enzymes, drawn using the kinetic parameters from Table 2. (A) Decarboxylation of FOMP catalyzed by wild-type OMPDC, which shows (a) the similar barriers to partitioning of EO between dissociation of OMP and the enzyme conformational change to form EC (Scheme 5), limits the values of these kinetic parameters. This reflects the small barrier to kcat for fast loop opening at the crippled enzyme, relative to the barrier for kcat for decarboxylation, which is not affected by these mutations. Figure 5B shows that the overall barrier to kcat/Km is the sum of the barriers to (i) formation of the Michaelis complex ([S] ≪ Km), (ii) Kc for the enzyme conformational change, and (iii) the activation barrier to kchem.

By comparison, the small 2- and S-fold effects of S-F on kcat and kcat/Km for Y217A mutant ScOMPDC-catalyzed decarboxylation of OMP (Table 2) show that the slow enzyme conformational change is rate-determining for the mutant enzyme-catalyzed decarboxylation reaction (Figure 5C). Now, the expected increase in kcat for loop opening for the mutant enzyme, due to the effect of the mutation on Kc (Table 2), has been approximately balanced by a second effect of this mutation that results in a decrease in k−c.

Intrinsic Reaction Barriers. Similar rate constants (kcat) are estimated for the Q215A/R235A [100 s−1 (Table 2)] and Y217A (18 s−1) mutant enzyme-catalyzed reactions; however, the conformational change for the Q215A/R235A mutant is relatively slow because of the large unfavorable thermodynamic driving force (kC = 1.3 × 103), while this conformational change for the Y217A mutant is slow because of the large intrinsic kinetic reaction barrier. These intrinsic kinetic barriers Λ (kilocalories per mole) may be defined by the Marcus equation (eq 5) as the reaction barrier in the absence of a thermodynamic driving force (ΔG° = 0).40−45 Intrinsic barriers Λ for the loop closure, with different ScOMPDC mutants, were estimated from rate constants kcat and corresponding equilibrium constants Kc reported in Table 2, using eq 5, derived for a reaction at 298 K. The Marcus intrinsic rate constants for hypothetical thermoneutral loop closure reactions at the different mutant enzymes (Table 2) were then estimated from Λ using eq 6, also derived at 298 K.

\[
\log k_c = \frac{1}{1.36} \left[ 17.44 - \Lambda \left( 1 - \frac{1.36 \log K_c}{4 \Lambda} \right)^2 \right]
\]

\[
\log k_{oc} = 12.8 - \Lambda \frac{1}{1.36}
\]

Table 2 shows that the R235A/Q215A double mutation results in an ≈106-fold decrease in Kc for loop closure but causes little change in intrinsic barrier Λ for the enzyme conformational change. By contrast, the Q215A/Y217F double mutation results in a 1.2 kcal/mol increase in Λ, which we attribute mainly to the effect of the conservative Y217F substitution, while the less conservative Y217A mutation results in a larger 2.2 kcal/mol increase in Λ. We propose that the phenol group at position 217 functions both to stabilize the closed form of ScOMPDC, through formation of a hydrogen bond to the substrate phosphodianion,18,19,34 and to reduce the intrinsic kinetic barrier to this enzyme conformational change. We suggest that the effect of these mutations of Y217 on Λ is related to their effect on the rate of formation of the highly organized intraloop clamping interaction between the -CH2OH side chain of Ser154 and the amide side chain of Gln215, which sits close to the phenol side chain of Y217 (Figure 4).18,34

CONCLUDING REMARK

The simplest interpretation of these results is that the large conformational change of OMPDC is necessary to lock OMP and FOMP into catalytically active caged Michaelis complexes. We have no evidence that this conformational change is coupled in any way to the decarboxylation step of the enzyme.

Figure 5. Free energy profiles for ScOMPDC-catalyzed decarboxylation of FOMP (Scheme 5), drawn for reactions at [S] ≪ Km using the kinetic parameters from Table 2. (A) Decarboxylation of FOMP catalyzed by wild-type OMPDC, which shows (a) the similar barriers to partitioning of EO and FOMP caged complex,19 and (c) the ≈3.5 kcal/mol difference [RT ln(390/1.1) (Scheme 2)] between the barriers for formation of EO and decarboxylation of FOMP. (B) Decarboxylation of FOMP catalyzed by the Q215A/R235A mutant of ScOMPDC. This mutation results in a 102-fold decrease in Kc for loop closure compared with that of wild-type ScOMPDC, but in little change in intrinsic barrier Λ for loop closure (Table 2). The decarboxylation step is rate-determining because kcat > kchem. (C) Decarboxylation of FOMP catalyzed by the Y217A mutant of ScOMPDC. This mutation results in a 2000-fold decrease in Kc for loop closure and a large increase in intrinsic barrier Λ for slow loop closure (Table 2), so that loop closure is rate-determining for this decarboxylation reaction.

Free Energy Reaction Profiles. Figure 5 shows free energy profiles for ScOMPDC-catalyzed decarboxylation of FOMP catalyzed by wild-type OMPDC (Figure 5A), the Q215A/R235A (Figure 5B), and Y217A (Figure 5C) mutant enzymes, drawn using the kinetic parameters from Table 2 for reactions at [S] ≪ Km. These profiles rationalize the two very different effects of the S-F substituent on mutant-enzyme catalyzed decarboxylation of OMP.

(1) The large 240- and 500-fold S-F substituent effects on kcat and kcat/Km for Q215A/R235A ScOMPDC-catalyzed decarboxylation show that the step for chemical decarboxylation, kchem (Scheme 5), limits the values of these kinetic parameters. This reflects the small barrier to k−c for fast loop opening at the crippled enzyme, relative to the barrier for k−c for decarboxylation, which is not affected by these mutations. Figure 5B shows that the overall barrier to kcat/Km is the sum of the barriers to (i) formation of the Michaelis complex ([S] ≪ Km), (ii) Kc for the enzyme conformational change, and (iii) the activation barrier to kchem.

(2) By comparison, the small 2- and S-fold effects of S-F on kcat and kcat/Km for Y217A mutant ScOMPDC-catalyzed decarboxylation of OMP (Table 2) show that a slow enzyme conformational change to form EC is rate-determining for this decarboxylation reaction. Now, the expected increase in kcat for loop opening for the mutant enzyme, due to the effect of the mutation on Kc (Table 2), has been approximately balanced by a second effect of this mutation that results in a decrease in k−c.
bound substrate. The steady-state kinetic parameters from Table 1 were obtained by the direct determination of decarboxylation reaction velocities. In several cases, the barrier to a nonchemical step, which we propose is an enzyme conformational change, was shown to control the observed reaction barrier. In these cases, the steady-state kinetic parameters were used to estimate the rate and equilibrium constants for this conformational change (Table 2). It would be interesting to examine the changes in the nuclear magnetic resonance or fluorescence spectral properties of ScOMPDC during turnover of FOMP, to obtain an independent estimate of the kinetic and thermodynamic barriers to this enzyme conformational change.

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**ABBREVIATIONS**
ScOMPDC, orotidine 5′-monophosphate decarboxylase from S. cerevisiae; OMP, orotidine 5′-monophosphate; UMP, uridine 5′-monophosphate; FOMP, 5-fluoroorotidine 5′-monophosphate; FUMP, 5-fluorouridine 5′-monophosphate; BMP, 6-hydroxyuridine 5′-monosophosphate; EO, 1-β-D-erythrofuranosyl)orotic acid; EU, 1-β-D-erythrofuranosyl)uracil; FEO, 1-β-D-erythrofuranosyl-5-fluoroorotic acid; FEU, 1-β-D-erythrofuranosyl-5-fluorouracil; MOPS, 3-(N-morpholino)propanesulfonic acid; PDB, Protein Data Bank.

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