The Organization of Active Site Side Chains of Glycerol-3-phosphate Dehydrogenase Promotes Efficient Enzyme Catalysis and Rescue of Variant Enzymes

Judith R. Cristobal, Archie C. Reyes, and John P. Richard*

ABSTRACT: A comparison of the values of $k_{cat}/K_m$ for reduction of dihydroxyacetone phosphate (DHAP) by NADH catalyzed by wild type and K120A/R269A variant glycerol-3-phosphate dehydrogenase from human liver (hlGPDH) shows that the transition state for enzyme-catalyzed hydride transfer is stabilized by 12.0 kcal/mol by interactions with the cationic K120 and R269 side chains. The transition state for the K120A/R269A variant-catalyzed reduction of DHAP is stabilized by 1.0 and 3.8 kcal/mol for reactions in the presence of 1.0 M EtNH$_3^+$ and guanidinium cation (Gua$^+$), respectively, and by 7.5 kcal/mol for reactions in the presence of a mixture of each cation at 1.0 M, so that the transition state stabilization by the ternary $E$·EtNH$_3^+$·Gua$^+$ complex is 2.8 kcal/mol greater than the sum of stabilization by the respective binary complexes. This shows that there is cooperativity between the paired activators in transition state stabilization. The effective molarities (EMs) of $\sim$50 M determined for the K120A and R269A side chains are $<10^8$ M, the EM for entropically controlled reactions. The unusually efficient rescue of the activity of hlGPDH-catalyzed reactions by the HPi/Gua$^+$ pair and by the Gua$^+$/EtNH$_3^+$ activator pair is due to stabilizing interactions between the protein and the activator pieces that organize the K120 and R269 side chains at the active site. This “preorganization” of side chains promotes effective catalysis by hlGPDH and many other enzymes. The role of the highly conserved network of side chains, which include Q295, R269, N270, N205, T264, K204, D260, and K120, in catalysis is discussed.

There are many examples of the rescue of the activity of truncated variant enzymes and of truncated alternative substrates by small molecule analogues of the deleted enzyme or substrate piece. However, such rescue is not universally observed, and there has been relatively little consideration of the structural requirements for the observation of efficient small molecule rescue of the catalytic activity of truncated enzymes and substrates. These studies provide insight into the mechanism for small molecule activation of enzyme activity that is analogous to allosteric activation, while enabling practical uses of chemical rescue in the activation of enzymes.

Glycerol-3-phosphate dehydrogenase (GPDH) catalyzes the reduction of dihydroxyacetone phosphate (DHAP) by NADH to form L-glycerol 3-phosphate [G3P (Scheme 1A)]. We have examined the activation of wild type human liver GPDH-catalyzed reduction of glycolaldehyde (GA) by phosphite dianion (HPi) and determined an EM value of 290 M for the phosphodianion of DHAP at the Michaelis complex with the wild type enzyme.3,4 The R269 side chain of hlGPDH interacts with the phosphodianion of DHAP and provides a 9.1 kcal/mol stabilization of the hydride transfer transition state (Figure 1).16 We determined an EM value of 60 M for this side chain in a study of the rescue of the R269A variant by the guanidine cation (Gua$^+$).16 We next examined the activity of...
The R269A variant is responsible for a large fraction of the ∼15 kcal/mol stabilization of the hydride transfer transition state.21,22 The K120 and R269 side chains are part of a network of highly conserved amino acid side chains that extend from Q295 to K120 and that includes N270, N205, K204, T264, D260, and K120 (Figure 1).23,24 The importance of this network is highlighted by the large effects of N270A25 and D260G25 substitutions on the activity of wild type GPDH, but the network’s full role in catalysis of hydride transfer has not been determined.

We report here the results of characterization of the efficiency of the rescue of K120A/R269A variant-catalyzed reduction of DHAP by the combined action of Gua⁺ and ethylammonium (EtNH₃⁺) cations (Scheme 1C) and the reduction of GA by the combined action of the phosphite dianion, Gua⁺, and EtNH₃⁺. The first set of experiments shows the efficient rescue by these two cations and gave an EM² value of 2400 M⁻² for the product of the effective molarity of the K120 and R269 side chains at the wild type enzyme. We conclude that the K120A/R269A variant provides a good template for binding of the excised cationic enzyme pieces and for organization of the K120A and R269A side chains at the active site of wild type GPDH. By contrast, we did not detect the fifth-order K120A/R269A variant-catalyzed reduction of GA in the presence of HPᵦ, Gua⁺, and EtNH₃⁺ activators. This sets a limit on the capacity of wild type GPDH to usefully assemble small molecule activators at the enzyme active site.

### EXPERIMENTAL SECTION

#### Materials

Water was obtained from a Milli-Q Academic purification system. Q-Sepharose and Sephacryl S-200 were purchased from GE Healthcare. Nicotinamide adenine dinucleotide, reduced form (NADH, disodium salt), glycolaldehyde dimer, 2-(N-morpholino)ethanesulfonic acid sodium salt (MES, ≥99.5%), triethanolamine hydrochloride (≥99.5%), guanidinium chloride, and sodium phosphate dibasic pentahydrate were purchased from Sigma-Aldrich. Ethylammonium chloride, d,l-dithiothreitol (DTT), sodium hydroxide (1.0 N), and hydrochloric acid (1.0 N) were purchased from Fisher Scientific. All other chemicals were reagent grade or better and were used without further purification. The solution pH was determined at 25 °C using an Orion model 720A pH meter equipped with a Radiometer pHC4006-9 combination electrode that was standardized at pH 4.00, 7.00, and 10.00 at 25 °C. Stock solutions of NADH were prepared by dissolving the disodium form of the coenzyme in water and then stored at 4 °C. The concentration of NADH in these solutions was determined from the absorbance at 340 nm using the extinction coefficient ε of 6220 M⁻¹ cm⁻¹. Stock solutions of DHAP were prepared by dissolving the lithium salt of DHAP in water, adjusting the pH to 7.5 with 1.0 NaOH, and storing the solution at −20 °C. The concentration of DHAP was determined as the concentration of NADH consumed during an hGPDH-catalyzed reduction. Published procedures were used to prepare stock solutions of the guanidine cation,3 ethylammonium cation,15 and phosphite dianion8 at pH 7.5. Triethanolamine (TEA) buffers were prepared by addition of 1 M NaOH or 1 M HCl and solid NaCl to give the desired acid/base ratio and final ionic strength. Stock solutions of glycolaldehyde dimer (200 mM monomer) were prepared by dissolving the dimer in water and waiting for 3 days at room temperature to allow for quantitative breakdown of the dimer to the monomer.6
The K120A/Q295A and K120A/R269A variants of hGPDH were expressed and purified by published procedures. Concentrated solutions of these variants were dialyzed exhaustively against 20 mM TEA buffer (pH 7.5) at 4 °C. When necessary, these solutions were diluted with 20 mM TEA buffer (pH 7.5) that contained 10 mM DTT and 0.1 mg/mL bovine serum albumin (BSA). The concentration of these enzyme variants was calculated from the absorbance at 280 nm using the extinction coefficient ε of 18450 M⁻¹ cm⁻¹ and a subunit molecular mass of 37500 Da that were determined using the ProtParam tool available on the ExPASy server.

**Enzyme Assays.** All enzyme assays were conducted at an ionic strength I of 0.12 (NaCl) in a volume of 1.0 mL at 25 °C. hGPDH was assayed by monitoring the oxidation of NADH (0.2 mM) by DHAP. Initial velocities of NADH oxidation over ≤10% reaction of DHAP were calculated from the change in absorbance at 340 nm using a molar extinction coefficient of 6220 M⁻¹ cm⁻¹ for NADH. Published procedures were used to assay the activity of K120A/Q295A and K120A/R269A variant-catalyzed reduction of DHAP by NADH at 25 °C, pH 7.5 (20 mM TEA), and I = 0.12 (NaCl) for the K120A/R269A variant-catalyzed reduction of GA under the same conditions.

**Activation of Variant hGPDH-Catalyzed Reactions.**

(A) Ethylammonium Cation. The assay mixtures at 25 °C and pH 7.5 (20 mM TEA) contained 0.1 mg/mL BSA and 0.2 mM NADH. For the K120A/Q295A variant, the assay mixture contained 1–3 mM DHAP, 0–80 mM activator, and 2.1 μM K120A/Q295A variant hGPDH. For the K120A/R269A variant, the reaction mixture contained 0.800 mM DHAP and 22 μM K120A/R269A variant hGPDH. The reactions were monitored for 20 min for the K120A/Q295A variant and for 720 min for the K120A/R269A variant.

(B) Guanidinium Cation. The assay mixtures at 25 °C and pH 7.5 (20 mM TEA) contained 0.1 mg/mL BSA, 0.2 mM NADH, 0.4–0.8 mM DHAP, 0–60 mM activator, and 20 μM K120A/R269A variant. The initial velocities for oxidation of NADH were calculated from the change in absorbance at 340 nm for a 20–40 min reaction time.

**Activation of the K120A/R269A Variant by Mixtures of Guanidinium and Ethylammonium Cations.** The assay mixtures for the K120A/R269A variant-catalyzed reduction of DHAP by NADH at I = 0.12 (NaCl) and 25 °C in the presence of mixtures of guanidinium and ethylammonium ions contained 20 mM TEA buffer (pH 7.5), 0.1 mg/mL BSA, 0.2 mM NADH, 0.4–0.8 mM DHAP, 0–60 mM guanidinium and ethylammonium cation activators (total concentration of the mixture), and 20 μM K120A/R269A variant hGPDH. The initial velocities for oxidation of NADH were calculated from the change in absorbance at 340 nm for a 20 min reaction time.

**RESULTS**

Slow GPDH-catalyzed reactions may be monitored for at least 24 h, during which time there is no detectable (<10%) loss of the activity of the wild type or K120A/R269A variant enzyme. There was no detectable reduction of 1.8 mM GA in the presence of 0.2 mM NADH catalyzed by 30 μM K120A/R269A variant hGPDH (ΔA₃₄₀ < 0.004 for a 40–60 min reaction time). This sets a limit for kcat/Km of ≤0.003 M⁻¹ s⁻¹ for this enzymatic reaction. There was likewise no detectable reduction of 1.8 mM GA at 0.2 mM NADH catalyzed by 30 μM K120A/R269A variant hGPDH in the presence of single activator HP, (20 mM), Gua⁺ (20 mM), or EtNH₃⁺ (20 mM), and in the presence of a mixture of 20 mM HP, 20 mM Gua⁺, and 20 mM EtNH₃⁺. There is no detectable effect (<5%) of 60 mM Gua⁺ or 60 mM EtNH₃⁺ on v/[E] for reduction of DHAP by 0.2 mM NADH catalyzed by the K120A or R269A variant, respectively, so that small molecule rescue of these variant enzymes is specific for the cation analogue of the excised side chain.

Figure 2 shows plots of v/[E] against [DHAP], with slopes (kcat/Km)obs for K120A/Q295A hGPDH-catalyzed reduction of DHAP by 0.2 mM NADH (saturating) at different fixed EtNH₃⁺ concentrations. The inset of Figure 2 shows the plot of (kcat/Km)obs against [EtNH₃⁺], with slope kcat/Km of 880 ± 20 M⁻² s⁻¹ (Table 1), for rescue by EtNH₃⁺. Figure 3A shows the plot of v/[E] against [EtNH₃⁺], with slope (kcat/Km)obs for K120A/R269A hGPDH-catalyzed reduction of 0.8 mM DHAP by 0.2 mM NADH (saturating). Combining the slope of this correlation (kcat/Km)obs (2.8 × 10⁻⁵ M⁻¹ s⁻¹) with a [DHAP] of 8 × 10⁻⁴ M gives a kcat/Km value of 0.035 M⁻² s⁻¹ (Table 1) for rescue of the K120A/R269A variant by EtNH₃⁺. Figure 3B shows the related plots of v/[E] against [Gua⁺], with slopes (kcat/Km)obs for K120A/R269A hGPDH-catalyzed reduction of 0.8 or 0.4 mM DHAP by 0.2 mM NADH. The inset of Figure 3B shows the plot of (kcat/Km)obs against [DHAP], with slope kcat/Km of 3.9 ± 0.1 M⁻² s⁻¹ (Table 1) for rescue of the K120A/R269A variant by Gua⁺ (Table 1).
Table 1. Kinetic Parameters and Derived Gibbs Free Energy Terms for Reactions of the Substrate and Pieces Catalyzed by hPGDH at 25 °C, pH 7.5 (20 mM TEA), and I = 0.12 (NaCl)

| variant       | $k_{cat}/K_m$ (M$^{-1}$ s$^{-1}$) | activator  | $k_{cat}/K_m$ (M$^{-2}$ s$^{-1}$) | $\Delta\Delta G^0_{\text{EM}}$ (kcal/mol) | $\Delta G_{\text{EM}}^{0} / \Delta G_{\text{EM}}^{0/2}$ (kcal/mol) | $\Delta G_{EM}^{0}$ (kcal/mol) |
|---------------|---------------------------------|------------|----------------------------------|-------------------------------------------|-------------------------------------------------|-------------------------------|
| K120A         | $550 \pm 30$                   | EtNH$_3^+$ | $(8.5 \pm 0.4) \times 10^9$      | $5.3 \pm 0.1$                             | $3.0 \pm 0.1$ (0.57)                              | $2.3$ (50 M)                  |
| K120A/Q295A   | $2.0 \pm 0.1$                   | EtNH$_3^+$ | $880 \pm 20$                    | $6.2 \pm 0.1$                             | $3.6 \pm 0.1$ (0.58)                              | $2.5$ (70 M)                  |
| R269A         | $1.0 \pm 0.2$                   | Gua$^+$    | $(8.0 \pm 0.5) \times 10^8$      | $9.1 \pm 0.1$                             | $6.7 \pm 0.1$ (0.74)                              | $2.4$ (60 M)                  |
| K120A/R269A   | $(6.2 \pm 0.4) \times 10^{-3}$ | Gua$^+$    | $(3.5 \pm 0.4) \times 10^{-2}$   | $2.9 \pm 0.1$                             | $1.0 \pm 0.1$ (0.33)                              | $2.0$ (30 M)                  |
|               |                                 | EtNH$_3^+$ | $1.9 \pm 0.1$                    | $6.7 \pm 0.1$                             | $3.8 \pm 0.1$ (0.56)                              | $2.9$ (140 M)                 |

From ref 4. *From ref 28. Second-order rate constant for variant hPGDH-catalyzed reduction of DHAP. bThird- or fourth-order rate constant for rescue of the activity of variant enzyme-catalyzed reduction of DHAP by the given activator(s). cEffect of the amino acid substitution on the stability of the transition state for wild type hPGDH-catalyzed reduction of DHAP, unless stated otherwise. dEffect of 1.0 M activator on the stability of the transition state for the variant hPGDH-catalyzed reduction. e,fTransition stabilization obtained from the covalent connection between the enzyme pieces ($\Delta G_{EM}^{0} = \Delta G^{0} - \Delta G_{EM}^{0/2}$). gEffective molarity, of wild type or variant hPGDH, of the deleted amino acid side chain ($k_{cat}/K_m$). hEffect of the K120A substitution on Q295A hPGDH. iEffect of the K120A substitution on R269A hPGDH. jEffect of the R269A substitution on K120 hPGDH. kUsing eq 6. lUsing eq 7. mUsing eq 3. nUsing eq 5. oUsing eq 6.

Figure 3. (A) Effect of increasing [EtNH$_3^+$] on $v/[E]$ (s$^{-1}$) for K120A/R269A variant hPGDH-catalyzed reduction of 0.8 mM DHAP by saturating 0.2 mM NADH for reactions at pH 7.5 (20 mM TEA buffer), 25 °C, [NADH] = 0.2 mM, and I = 0.12 (NaCl). (B) Effect of increasing [Gua$^+$] on $v/[E]$ (s$^{-1}$) for K120A/R269A variant hPGDH-catalyzed reduction of DHAP by 0.2 mM NADH for reactions at pH 7.5 (20 mM TEA buffer), 25 °C, saturating [NADH] = 0.2 mM, and I = 0.12 (NaCl): (●) 0.8 mM DHAP and (●) 0.4 mM DHAP. The inset shows the plot of $(k_{cat}/K_m)_{\text{obs}}$ against [DHAP], where $(k_{cat}/K_m)_{\text{obs}}$ values are the slopes of the correlations from the main panel. The empty symbols show the agreement of data obtained for two different preparations of the K120A/R269A variant.

Figure 4A ([DHAP] = 0.4 mM) and Figure 4B ([DHAP] = 0.8 mM) show plots of $v/[E]$ against [Gua$^+$], with slopes $(k_{cat}/K_m)_{\text{obs}}$ for K120A/R269A hPGDH-catalyzed reduction of 0.2 mM NADH at different fixed concentrations of EtNH$_3^+$. Figure 5 shows plots of $(k_{cat}/K_m)_{\text{obs}}$ from panels A and B of Figure 4 against [EtNH$_3^+$], with slopes $(k_{cat}/K_m)_{\text{obs}}$ for K120A/R269A hPGDH-catalyzed reduction of 0.4 or 0.8 mM DHAP by 0.2 mM NADH. The inset of Figure 5 shows the plot of $(k_{cat}/K_m)_{\text{obs}}$ against [DHAP], with a slope $(k_{cat}/K_m)_{\text{obs}}$ of 1900 M$^{-1}$ s$^{-1}$ (Scheme 2 and Table 1).

**DISCUSSION**

The failure to detect reduction of GA catalyzed by 20 μM R269A hPGDH results in a $(k_{cat}/K_m)_{\text{obs}}$ of $\leq 0.003$ M$^{-1}$ s$^{-1}$ for the variant hPGDH-catalyzed reaction. The failure to observe activation of the reduction of GA catalyzed by 30 μM K120A/R269A hPGDH by 20 mM HP$_2$ Gua$^+$, or EtNH$_3^+$ activators or by a mixture of 20 mM HP$_2$ and 20 mM Gua$^+$, and 20 mM EtNH$_3^+$, shows that $(k_{cat}/K_m)_{\text{obs}} \leq 0.003$ M$^{-1}$ s$^{-1}$ for the K120A/R269A variant-catalyzed reaction in the presence of these activators. Equation 1 gives the relationship between this limit for $(k_{cat}/K_m)_{\text{obs}}$ and the fifth-order rate constant $k_Q$ for the reaction catalyzed by the quaternary complex of the K120A/R269A variant, GA, HP$_2$, Gua$^+$, and EtNH$_3^+$. These data set an upper limit for $k_Q$ of 375 M$^{-4}$ s$^{-1}$ (eq 2). This shows that two small molecules may act together to give detectable activation of GPDH under our experimental conditions [HP$_2$, and Gua$^+$ for R269A variant hPGDH-catalyzed reduction of GA$^+$ and Gua$^+$ and EtNH$_3^+$ for K120A/R269A variant hPGDH-catalyzed reduction of DHAP (this work, Table 1)] but that it is not
Gua+, and EtNH₃ for the reaction of 0.4 mM DHAP at different fixed EtNH₃⁺ concentrations. (B) Increase in v/[E] with increasing [Gua⁺], for the reaction of 0.4 mM DHAP at different fixed EtNH₃⁺ concentrations. (A) Increase in v/[E] with increasing [Gua⁺], for the reaction of 0.4 mM DHAP at different fixed EtNH₃⁺ concentrations.

**Figure 4.** Effect of increasing [Gua⁺] and [EtNH₃⁺] on v/[E] for K120A/R269A variant hGPDH-catalyzed reduction of DHAP by NADH at pH 7.5 (20 mM TEA buffer), 25°C, saturating [NADH] = 0.2 mM, and I = 0.12 (NaCl). (A) Increase in v/[E], with increasing [Gua⁺], for the reaction of 0.4 mM DHAP at different fixed EtNH₃⁺ concentrations. (B) Increase in v/[E], with increasing [Gua⁺], for the reaction of 0.8 mM DHAP at different fixed EtNH₃⁺ concentrations: (●) 5 mM EtNH₃⁺, (▲) 10 mM EtNH₃⁺, (■) 20 mM EtNH₃⁺, and (○) 30 mM EtNH₃⁺. The empty symbols show the agreement of data obtained for two different preparations of the K120A/R269A variant hGPDH.

**Figure 5.** Effect of increasing [EtNH₃⁺] on (kₘcat/KₘGua)obs for K120A/R269A variant hGPDH-catalyzed reduction of DHAP by NADH at pH 7.5 (20 mM TEA buffer), 25°C, saturating [NADH] = 0.2 mM, and I = 0.12 (NaCl). The slope of these correlations is (kₘcat/KₘGua)[km]obs for variant hGPDH-catalyzed reduction activated by the combined action of Gua⁺ and EtNH₃⁺: (■) 0.4 mM DHAP and (○) 0.8 mM DHAP. The inset shows the plot of (kₘcat/KₘGua,KₘEtNH₃)[km]obs against [DHAP], with a slope of (kₘcat/KₘGua,KₘEtNH₃)[km] of 1900 M⁻¹s⁻¹ (Scheme 2 and Table 1).

**Scheme 2. Rescue of the K120A/R269A Variant hGPDH by EtNH₃ and Gua⁺**

E-NADH + DHAP & (kₘcat/KₘGua,KₘEtNH₃)[km] & E-NAD⁻ - Gly³-P⁺

possible to detect activation by the combined action of HP, Gua⁺, and EtNH₃⁺.

\[
\frac{(k_{\text{cat}}/K_{\text{Gua}})_{\text{obs}}}{[\text{Gua}^+][\text{HP}][\text{EtNH}_3^+]} = k_Q \leq 0.003 \text{ M}^{-1}
\]

\[
\leq 8 \times 10^{-9} \text{ M}^3
\]

\[
\leq 375 \text{ M}^{-1} \text{s}^{-1}
\]

(2)

**Rescue of the K120A/Q295A Variant-Catalyzed Reactions of DHAP.** The K120A substitution results in similar 5.3 and 6.2 kcal/mol increases in the activation barrier for wild type- and Q295A variant-catalyzed reduction of DHAP, respectively (Table 1). The efficiency of rescue of the K120A/Q295A variant by EtNH₃⁺ was characterized in this work (Figure 2) and compared with the efficiency of rescue of the K120A variant (Table 1). The transition states for the K120A and K120A/Q295A hGPDH-catalyzed reductions of DHAP show similar stabilizations of 3.0 and 3.6 kcal/mol, respectively, for reactions in the presence of 1.0 M EtNH₃⁺, while the EMs of 50 and 70 M determined for the K120 side chain of the wild type and the Q295A variant of hGPDH, respectively, are similar (Table 1). These data provide additional support for the conclusion that the Q295A substitution only slightly impairs the transition state stabilization by the K120 side chain.

**K120A/R269A Variant-Catalyzed Reactions of DHAP.** The position of side chains at the ternary hGPDH-NAD-DHAP complex is shown in Figure 1, while panels A–C of Figure 6 show the side chains as representations of the surface of the binary hGPDH-NAD complex (Figure 6A), the binary hGPDH-NAD complex with DHAP inserted at the position of the ternary complex (Figure 6B), and the ternary hGPDH-NAD-DHAP complex (Figure 6C). The K120 and R269 side chains lie on opposite sides of the active site cavity, with the K120 side chain positioned to interact with the carbonyl group and the R269 side chain positioned to interact with the phosphodianion of the substrate. The side chains lie at the two ends of a network of highly conserved side chains (Figure 1), which extends from Q295 to K120 and includes R270, N205, K204, D260, and K120. A comparison of the structures shown in panels A–C of Figure 6 shows that the ligand-driven enzyme conformational change results in the folding of a flexible protein loop (292-LNGQKL-297) over the DHAP and NAD cofactor that is facilitated by formation of a hydrogen bond between the Q295 and R269 side chains. Loop closure traps the substrate in a solvent-occluded cage, where the K120 and R269 side chains are optimally placed for catalysis, and where electrostatic catalysis is presumably enhanced by the decrease in the effective dielectric constant of the closed compared to the open form of the complex (Figure 6A–C).

The consecutive K120 and R269 substitutions result in a total 12 kcal/mol increase in the activation barrier ΔG° for hydride transfer (Scheme 3). The efficient rescue of the K120A/R269A variant of hGPDH by Gua⁺ and EtNH₃⁺ (Figures 4 and 5) shows that the wild type and variant enzymes proceed through similar transition states, which are strongly stabilized by interactions with the K120 and R269 side chains (wild type hGPDH) or with bound Gua⁺ and EtNH₃⁺. The larger sum of the effects of individual K120A and R269A substitutions on ΔG° (5.3 + 9.1 = 14.4 kcal/mol) shows that the interaction energies of the single side chains of wild type hGPDH are higher than the total 12 kcal/mol interaction determined by deleting the two side chains.

The 2.4 kcal/mol difference between the total side chain interaction estimated when the remaining side chain is preserved and the interaction determined by deleting both
side chains (Scheme 3) represents the stronger side chain interactions of the tight, organized, conformation of wild type hlGPDH, compared to interactions of the K120A or R269A variant. We propose that this difference is due to effects of the first substitution, which reduce the transition state stabilization by the second side chain, such as an increase in the side chain conformational flexibility. This proposal is consistent with the notion that wild type hlGPDH derives a catalytic advantage from the high degree of organization of the catalytic side chains at the wild type active site and that the effect on $\Delta \Delta G^\ddagger$ of substitutions that erode this organization is greater than the effect of the lost interaction between the transition state and excised side chain.

Rescue of K120A/R269A hlGPDH by EtNH$_3^+$ and Gua$^+$. The kinetic data for rescue of the K120A/R269A variant of hlGPDH by Gua$^+$ and EtNH$_3^+$ give a $(\Delta \Delta G^\ddagger)_{E+EtNH_3+Gua}$ of 7.5 kcal/mol (eq 3, Table 1) for stabilization of the transition state by interaction with the bound cations that is only 4.5 kcal/mol smaller than the transition state stabilization by the K120 and R269 side chains of wild type hlGPDH ($\Delta \Delta G^\ddagger = 12.0$ kcal/mol, eq 4). The difference corresponds to a $12.0 - 7.5 = (\Delta G^\ddagger)_{E+EtNH_3+Gua} = 4.5$ kcal/mol advantage to the reaction catalyzed by wild type hlGPDH compared to catalysis by the K120A/R269A variant in the presence of 1.0 M Gua$^+$ and EtNH$_3^+$ (eq 5). By comparison, the binding of 1.0 M EtNH$_3^+$ or 1.0 M Gua$^+$ to the K120A/R269A variant provides a $(\Delta G^\ddagger)_{EtNH_3} = 1.0$ (eq 6) or $(\Delta G^\ddagger)_{Gua} = 3.8$ kcal/mol (eq 7) stabilization, respectively, of the transition state for the reaction catalyzed by this variant, so that the sum of transition state stabilization by the individual cations in binary complexes (1.0 + 3.8 = 4.8 kcal/mol) is 2.7 kcal/mol smaller than the 7.5 kcal/mol stabilization observed for the cations in the ternary complex.

Scheme 3. Cycle That Shows the Effect of Consecutive K120A and R269A Substitutions on $\Delta \Delta G^\ddagger$ for Wild Type hlGPDH-Catalyzed Reduction of DHAP by NADH

\[
\begin{align*}
\text{WT} & : \Delta \Delta G^\ddagger_{\text{DHAP}} = 9.1 \text{ kcal/mol} \\
\text{K120A} & : \Delta \Delta G^\ddagger_{\text{DHAP}} = 6.7 \text{ kcal/mol} \\
\text{R269A} & : \Delta \Delta G^\ddagger_{\text{DHAP}} = 2.9 \text{ kcal/mol}
\end{align*}
\]

Scheme 4. Comparison between the Total Transition State Stabilization for Wild Type hlGPDH by Interactions with the K120 and R269 Side Chains and the Transition State Stabilization for the K120A/R269A Variant by Interactions with Exogenous Cations

\[
\begin{align*}
(\Delta \Delta G^\ddagger)_{E+EtNH_3+Gua} &= -RT \ln \left( \frac{k_{\text{cat}}/K_{\text{cat}}K_{\text{Gua}}K_{\text{EtNH_3}}}{(k_{\text{cat}}/K_{\text{cat}})_{\text{K120A/R269A}}} \right) \\
&= -7.5 \text{ kcal/mol} & (3)
\end{align*}
\]

\[
\begin{align*}
(\Delta \Delta G^\ddagger)_{E+EtNH_3+Gua} &= -RT \ln \left( \frac{k_{\text{cat}}/K_{\text{cat}}}{(k_{\text{cat}}/K_{\text{cat}})_{\text{K120A/R269A}}} \right) \\
&= -12.0 \text{ kcal/mol} & (4)
\end{align*}
\]

\[
\begin{align*}
(\Delta G^\ddagger)_{E+EtNH_3+Gua} &= RT \ln \left( \frac{(k_{\text{cat}}/K_{\text{cat}})_{\text{Gua}}}{(k_{\text{cat}}/K_{\text{cat}})_{\text{EtNH_3}}} \right) \\
&= 4.5 \text{ kcal/mol} & (5)
\end{align*}
\]
Figure 7. (A) Representations of the X-ray crystal structures of the following complexes with variant hiGPDH. (A) Complex of K120A/R269A hiGPDH with NAD, DHAP, Gua+, and EtNH3+. (B) Complex of R269A hiGPDH with NAD, GA, HPi, and Gua+. The complexes were generated in silico, starting with Figure 1 for the X-ray crystal structure of the nonproductive complex of wild type hiGPDH, DHAP, and NAD+ (PDB entry 6E90) with deletion of the relevant covalent linkage(s) while maintaining a fixed position for the remaining atoms of the hypothetical Michaelis complexes.

\[
\Delta G_{cat}^{T} = -RT \ln \left( \frac{k_{cat}/K_{m}}{k_{cat}^{0}/K_{m}^{0}} \right) = -1.0 \text{ kcal/mol} \quad (6)
\]

\[
\Delta G_{cat}^{T} = -RT \ln \left( \frac{k_{cat}/K_{m}}{k_{cat}^{0}/K_{m}^{0}} \right) = -3.8 \text{ kcal/mol} \quad (7)
\]

These results are illustrated by Scheme 4, which partitions the total 12.0 kcal/mol effect of the K120A and R269A substitutions on transition state stability into the 7.5 kcal/mol interaction recovered in the EtNH3+·Gua+ complex and the 4.5 kcal/mol advantage to the reaction of the intact enzyme. The 7.5 kcal/mol recovered interaction is then partitioned into the 1.0 and 3.8 kcal/mol advantage for transition state stabilization of the individual binary complexes and the 2.7 kcal/mol advantage for cation activation of the ternary E·EtNH3+·Gua+ complex. By comparison, the sum of the effect of single K120 and R269 substitutions in the K120A/R269A variant of hiGPDH (6.7 + 3.0 = 9.7 kcal/mol) on the stability of the transition state for enzyme-catalyzed reduction of DHAP is 2.3 kcal/mol smaller than the overall 12 kcal/mol stabilization by consecutive K120 and R269 substitutions (Scheme 3). We conclude that substitution of a single cationic side chain, or binding of a small molecule side chain analogue, in the K120A/R269A variant enhances the interaction of the second side chain or cation. We propose that this is due to utilization of the binding energy of the first bound cation in the organization of the active site, which enhances the transition state stabilization by the second bound cation. It is interesting that these cooperative interactions are expressed between side chains that are separated by \( \sim 9 \) Å.

The 4.5 kcal/mol “connection energy” (Scheme 4) is an estimate for the advantage to connecting the Gua+ and EtNH3+ cations to the protein at the K120A/R269A variant of hiGPDH. This defines the effective concentration or effective molarity (EM)\(^{13}\) of the side chains of wild type hiGPDH compared to the value of 1.0 M of the free side chain in water (Table 1). EMs of 50 and 60 M were determined for the K120 and R269 side chains from the efficiency of rescue of the K120A and R269A variants, respectively.\(^{21}\) The product of the EMs from studies of single variants (50 M)(60 M) = 3000 M\(^2\) is similar to the value of 2400 M\(^2\) determined from rescue studies of the K120A/R269A variant (Table 1). This is consistent with similar stabilizing interactions between the catalyst and rescue agents of the K120A and R269A single variants and the K120A/R269A double variant.

The EMs from Table 1 are clustered between 30 and 140 M and correspond to an \(~2.5\) kcal/mol advantage in \( \Delta G_{cat}^{T} \) for reactions catalyzed by wild type hiGPDH compared to the reactions catalyzed by the complex between the variant enzyme and the missing piece. These values of EMs and \( \Delta G_{cat}^{T} \) are smaller than the values of \(~10^6\) M and \(~8\) kcal/mol predicted for cases in which the advantage for unimolecular compared with bimolecular reaction is wholly entropic.\(^{14,32}\) The low EMs from Table 1 reflect the effective stabilization of the complexes to the pieces by interactions with the protein catalyst (Figure 1),\(^{13}\) where the K120 chain is locked into place by an ion pair to the D260 side chain,\(^{21}\) and the R269 side chain is held by interactions with the phosphodianion of DHAP, the pyrophosphate anion of NAD, and the Q295 side chain (Figure 1). These same interactions stabilize the EtNH3+ and Gua+ pieces bound to the K120A/R269A variant of hiGPDH (Figure 7A).

Efficiency of Hydride Transfer Catalyzed by hiGPDH. The large transition state stabilization for hiGPDH-catalyzed reduction of DHAP is achieved largely through strong, focused, stabilizing interactions with the K120 and R269 side chains.\(^{21}\) The recovery of these interactions through the robust rescue of the K120A/R269A variant by Gua+ and EtNH3+ enzyme pieces reflects the strong stabilization of complexes with Gua+ and EtNH3+ (Figure 7A) by interaction with the neighboring amino acid side chains discussed above. Similarly, there is efficient activation of R269A variant hiGPDH-catalyzed reduction of glycolaldehyde by the combined action of the exogenous phosphate dianion and guanidine cation.\(^{1}\) This is a
consequence of the stabilization of the enzyme-bound HP; Ga− ion pair (Figure 7B) by hydrogen bonding and ionic interactions, respectively, of Ga− with the Q295 side chain and the cofactor pyrophosphate oxygen and by hydrogen bonds between HP; and the A269 backbone amide and the N270 amide side chain.

The side chain interactions, which promote efficient rescue of the K120A and R269A variants, serve to organize (or to preorganize) the K120 and R269 side chain cations of wild type hGPDH (Figure 1). We propose that this preorganization enables the large, focused, 12 kcal/mol transition state stabilization from interactions with the K120 and R269 side chains by minimizing the energetic price for side chain immobilization that occurs on proceeding from the Michaelis complex to the hydride transfer transition state. Additional examples of side chain preorganization at enzyme active sites, such as the catalytic triad found in serine proteases, have been discussed by Warshel and coworkers.

There are surprising similarities between the mechanisms for hydride transfer from NADH to DHAP catalyzed by hGPDH and for isomerization of DHAP, with proton transfer, catalyzed by triosephosphate isomerase (TIM). Each enzyme shows strong phosphite dianion (HP;), activation of catalysis of the reaction of the common truncated substrate glyceraldehyde. The K12 side chain at TIM sits near the substrate dianion, and the K12G variant shows efficient rescue by alkyl ammonium cations. However, there is no rescue of K12G-catalyzed deprotonation of GA by the combined action of HP; and RH;+. The stronger stabilization of the transition state for R269A variant-catalyzed hydride transfer to GA by the HP; Ga− ion pair compared with the transition state for K12G variant-catalyzed deprotonation of GA by the HP; EtNH3+ ion pair is consistent with a higher degree of preorganization of the ion pair at the active site of hGPDH. This reflects, at least in part, the presence of an intervening water molecule between the substrate phosphodianion and the TIM K12 side chain.

**A Conserved Network of Amino Acid Side Chains.**

The highly conserved side chains from Q295, R269, N270, N205, T264, K204, D260, and K120 form a continuous chain of hydrogen bonds that stretch from the bound cofactor to the carbonyl group of DHAP (Figures 1 and 7). This side chain conservation suggests that the network operates as a unit, with K120, K204, and R269 providing direct transition state stabilization; N205, D260, and Q295 functioning directly to immobilize the catalytic side chains; and N270, N205, and T264 playing secondary roles in maintaining the network’s structural integrity. We suggest that this tight network of side chain interactions promotes effective catalysis and that the preorganization of this network by intra-side chain interactions with K120 and R269 provides a mechanism for the expression of cooperative interactions between the two cations, which are separated by 9 Å (Figure 1).

The D260 side chain shows no direct stabilizing interaction with the hydride transfer transition state, and there should be only a weak interaction with the N270 side chain. The large 6.5 and 5.6 kcal/mol effects of D260G and N270A substitutions, respectively, on the stability of the transition state for wild type hGPDH-catalyzed reduction of DHAP by NADH (neither of which is subject to small molecule rescue) are therefore consistent with a substantial reorganization of the extended side chain network at the D260G and N270A variants, which results in barriers to formation of the preorganized, catalytically active closed conformation of variant hGPDH. Finally, the observation that the N270A substitution results in an ~40-fold increase in kcat/Km for enzyme-catalyzed reduction of GA by NADH is consistent with a reorganization of the active site at this variant, which favors binding of GA in a reactive conformation. This unusual observation shows that there is still much to be learned about the exact role of this extended network of interactions in the organization of active site side chains that provides for optimal catalysis of the reactions of whole and truncated substrates.

**CONCLUSIONS**

The R269 side chain of GPDH functions to trap the substrate DHAP in a tight cage, which provides strong stabilization of the transition state for hydride transfer, while the K120 side chain acts to stabilize negative charge at the C-2 oxygen, which develops in this transition state. The K120A and R269A substitutions at GPDH result in a total 7 × 108-fold decrease in kcat/Km for catalysis of NADH by hGPDH, which corresponds to an enormous 12 kcal/mol destabilization of the transition state for hydride transfer: the K120A/R269A variant provides an excellent template for the expression of exogenous Ga− and EtNH3+ activators, which has a 7.5 kcal/mol transition state stabilization at standard states of 1.0 M Ga− and EtNH3+. The transition state stabilization from single R269 or K120 side chains, or from single Ga− or EtNH3+ activators, is enhanced by the presence of the second side chain or activator. These results provide compelling support for the conclusion that the network of conserved side chains at GPDH (Figure 1) functions in the preorganization of the K120 and R269 side chains into positions that provide optimal stabilization of the transition state for hydride transfer. This preorganization promotes cooperativity in the expression of the strongly stabilizing interactions of the K120 and R269 side chains across a separation distance of 9 Å. We propose that the side chains at enzyme active sites often function organically as a unit to provide for optimal stabilizing interactions between the transition state and a few key catalytic side chains.

**ASSOCIATED CONTENT**

**Accession Codes**

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**AUTHOR INFORMATION**

**Corresponding Author**

John P. Richard — Department of Chemistry, University at Buffalo, State University of New York, Buffalo, New York 14260-3000, United States; orcid.org/0000-0002-0440-2387; Email: jrichard@buffalo.edu

**Authors**

Judith R. Cristobal — Department of Chemistry, University at Buffalo, State University of New York, Buffalo, New York 14260-3000, United States

Archie C. Reyes — Department of Chemistry, University at Buffalo, State University of New York, Buffalo, New York 14260-3000, United States; orcid.org/0000-0001-9955-393X

Complete contact information is available at: https://pubs.acs.org/10.1021/acs.biochem.0c00175
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□ ABBREVIATIONS
ScOMPDC, orotidine 5’-monophosphate decarboxylase from yeast; TIM, triosephosphate isomerase; GPDH, glycerol-3-phosphate dehydrogenase; hGPDH, glycerol-3-phosphate dehydrogenase from human liver; DHAP, dihydroxyacetone phosphate; etG, ethylene glycol; GA, glycolaldehyde; NADH, nicotinamide adenine dinucleotide, reduced form; NAD, nicotinamide adenine dinucleotide, oxidized form; MES, 2-(N-morpholino)ethanesulfonic acid; TEA, triethanolamine; DTT, D,L-dithiothreitol; BSA, bovine serum albumin.

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