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ABSTRACT: Transition-state structures of human and bovine of purine nucleoside phosphorylases differ, despite 87% homologous amino acid sequences. Human PNP (HsPNP) has a fully dissociated transition state, while that for bovine PNP (BtPNP) has early SN 1 character. Crystal structures and sequence alignment indicate that the active sites of these enzymes are the same within crystallographic analysis, but residues in the second-sphere from the active sites differ significantly. Residues in BtPNP have been mutated toward HsPNP, resulting in double (Asn123Lys; Arg210Gln) and triple mutant PNPs (Val39Thr; Asn123Lys; Arg210Gln). Steady-state kinetic studies indicated unchanged catalytic activity, while pre-steady-state studies indicate that the chemical step is slower in the triple mutant. The mutant enzymes have higher affinity for inhibitors that are mimics of a late dissociative transition state. Kinetic isotope effects (KIEs) and computational chemistry were used to identify the transition-state structure of the triple mutant. Intrinsic KIEs from [1-3H], [1-14C], [2-3H], [5-3H], and [9-15N] inosines were 1.221, 1.035, 1.073, 1.062 and 1.025, respectively. The primary intrinsic [1-14C] and [9-15N] KIEs indicate a highly dissociative Sn1 transition state with low bond order to the leaving group, a transition state different from the native enzyme. The [1-14C] KIE suggests significant nucleophilic participation at the transition state. The transition-state structure of triple mutant PNP is altered as a consequence of the amino acids in the second sphere from the catalytic site. These residues are implicated in linking the dynamic motion of the protein to formation of the transition state.

Purine nucleoside phosphorylase (PNP, E.C. 2.4.2.1) catalyzes the phosphorylation of the N-ribosidic bonds of 6-oxypurine nucleosides and deoxynucleosides. The ratelimiting step for mammalian PNPs in the phosphorylase reaction is release of the purine base (1). Transition-state analysis for PNPs from the human, bovine and Plasmodium falciparum enzymes reveals a unique transition-state structure for bovine PNP (2–4). While the intrinsic [1-14C] kinetic isotope effects are unity for human and Plasmodium falciparum PNPs, consistent with fully dissociated Sn1 transition states, that for bovine PNP is 1.022, indicating an early dissociative mechanism for bovine PNP (Figure 1) (2, 3).

The amino acid sequences of human PNP (HsPNP) and bovine PNP (BtPNP) share 87% identity overall and are completely conserved at the catalytic sites (Figure 2), while their transition states are different (2, 4). Structural comparison of the catalytic sites with both early and late transition state analogue inhibitors reveals no significant changes to suggest differences in these transition states (Figure 3; 5, 6). Structural and sequence comparisons reveal significant differences in amino acids at the subunit interface and in the second sphere from the surface of the proteins. Twenty-two of the thirty-three nonconserved residues are located on the surface of the proteins. Residues not located on the surface were examined for their distance from the active site (Table 1). Of these, residues 39, 123, and 210 of BtPNP are highly dissimilar and are 10–14 Å from the active site. A double mutant (Asn123Lys and Arg210Gln) and a triple mutant BtPNP (Val39Thr, Asn123Lys, and Arg210Gln) were created, with these residues being changed to those found in HsPNP. The purpose of these changes was to test the hypothesis that changes remote from the catalytic site could alter the transition state of the bovine enzyme to become more like HsPNP. Inhibition and kinetic isotope effect studies were conducted to characterize the chimeric proteins for comparison with the parent enzymes. The results establish that transition-state structure of BtPNP can be altered by remote mutations and implicate dynamic contribution to transition-state structure.

MATERIALS AND METHODS

Enzymes and Reagents. The reagents and enzymes used in the synthesis of isotopically labeled inosines have been described (4, 7–9). All chemicals were the highest quality available and were purchased from commercial sources as described previously (1, 7).

Mutagenesis, Transformation and Expression. The cDNAs were obtained from DNA 2.0 Inc., with nucleotide sequences optimized for expression in Escherichia coli. A thrombin cleavable 6-His tag was encoded at the N-terminus of the
genes, which were subcloned into the pDNR vector. The plasmids were inserted into the pBAD-DEST49 vector by LR clonase II, and the resulting plasmids were transformed into XL-Blue competent cells. The sequences of the plasmids were verified by automated DNA sequencing. The recombinant plasmids were transformed into BL21A.I.(DE3) competent cells and plated onto LB/agar plates containing 100 μg/mL ampicillin. Single colonies were inoculated into 50 mL of LB containing 100 μg/mL ampicillin with shaking at 220 rpm, 37 °C. The overnight cultures were then transferred into 10 L of LB containing 100 μg/mL ampicillin and grown at 37 °C with shaking at 220 rpm to OD₆₀₀ of 0.7. L-Arabinose (0.1%) was added to induce the overexpression for 8–10 h at 28 °C. Cells were harvested at 4000 rpm for 30 min at 4 °C, and cell pellets were stored at −80 °C before use.

**Purification and Protein Concentration Assay.** Cells were resuspended in 50 mL of lysis buffer, containing 20 mM
Table 1: Distance from the Active Site of Residues That Are Not Conserved between BtPNP and HsPNP

<table>
<thead>
<tr>
<th>residue no.</th>
<th>PNP identity</th>
<th>distance from active site (Å)</th>
</tr>
</thead>
<tbody>
<tr>
<td>2</td>
<td>bovine Glu</td>
<td>20.4</td>
</tr>
<tr>
<td>12</td>
<td>Asp</td>
<td>26.1</td>
</tr>
<tr>
<td>23</td>
<td>Gln</td>
<td>23.5</td>
</tr>
<tr>
<td>29</td>
<td>Val</td>
<td>10.4</td>
</tr>
<tr>
<td>39</td>
<td>Val</td>
<td>13.1</td>
</tr>
<tr>
<td>40</td>
<td>Asp</td>
<td>16.8</td>
</tr>
<tr>
<td>93</td>
<td>Phe</td>
<td>11.9</td>
</tr>
<tr>
<td>123</td>
<td>Asn</td>
<td>14.3</td>
</tr>
<tr>
<td>152</td>
<td>Glu</td>
<td>26.6</td>
</tr>
<tr>
<td>210</td>
<td>Arg</td>
<td>10.8</td>
</tr>
<tr>
<td>277</td>
<td>Leu</td>
<td>19.3</td>
</tr>
</tbody>
</table>

*Residues targeted for construction of the chimeric proteins are highlighted. Distance from the active site was a measure of the distance to the nearest point of bound immmuclin-H.

KHPo₄, pH 7.5, 1 protease inhibitor cocktail tablet and ~10 mg of DNase I. Cells were disrupted by three passes through a French pressure cell. After centrifugation (15,000 rpm for 30 min at 4 °C), the supernatant was applied to a 70 mL Ni-NTA column. Wash buffer (150 mL) containing 10 mM imidazole, 100 mM NaCl and 50 mM KHPo₄, pH 7.5 was applied followed by a 1 to 10,000 mM imidazole gradient in the same buffer. The PNPs eluted at approximately 300 mM imidazole. The purified proteins were concentrated, in the same buffer. The PNPs eluted at approximately 300 mM imidazole. The purified proteins were concentrated, dialyzed against 20 mM Hepes, pH 7.4, 100 mM NaCl and 10% glycerol, and stored at ~80 °C, where they are stable for more than 1 year. The PNP concentrations were determined from the corresponding ATP molecules using the molar extinction coefficients of 9540 M⁻¹ cm⁻¹, calculated by the ExPaSy ProtParam Tool, us.expasy.org (10).


**Steady-State Kinetic and Pre-Steady-State Kinetic Studies.** Initial rate studies were performed for the mutant and wild type enzymes using a xanthine oxidase coupled assay (13, 14). Initial rates were measured at pH 7.5, 50 mM Na₂HPO₄ buffer, as a function of inosine concentration (0.5–10 Kₘ) at 25 °C. The formation of uric acid was monitored at 293 nm using the published molar extinction coefficient of 12,900 M⁻¹ cm⁻¹ (15). Pre-steady-state kinetic studies were carried out using an Applied Photophysics stopped flow spectrophotometer (model π⁺–180 spectrometer) equipped with a thermostated water bath. Equal volumes of ~4 μM enzyme in 50 mM KH₂PO₄, pH 7.4 and ~2 mM guanosine were rapidly mixed, and the fluorescence increase caused by enzyme-bound guanine was monitored above 290 nm while exciting at 280 nm. The temperature was varied from 4 °C to 20 °C. Two hundred data points were collected within 0.1 to 0.5 s at each temperature in triplicates. The single turnover rates at different temperatures were calculated by fitting the stopped-flow traces to the single-exponential decay equation (eq 1), where kₖ清远 represents the first-order rate constant of the chemical step (the conversion of PNP:guanosine:PO₄ to PNP:guanine:R-1-P, Figure 3), Fᵢ is the fluorescence emission intensity at 290 nm at time t, A is the amplitude of the total change and Fᵢ is the fluorescence intensity at infinite time. The temperature dependence of the single turnover rate was determined by fitting the data with the Arrhenius and Eyring equations (eq 2 and eq 3, respectively), where kᵦ and h are the Boltzmann and Planck constants, respectively, A is the preexponential factor, E₀ is activation energy, R is the gas constant (8.31 J mol⁻¹ K⁻¹) and T is the temperature in kelvins. The enthalpy of activation (ΔH°) is calculated from the slope of the plots, whereas the entropy (ΔS°) is calculated from y-intercepts of the plots.

\[
Fᵢ = A(e^{-kt}) + Fᵢ
\]

\[
\ln(k) = \ln(A) - \frac{E₀}{RT}
\]

\[
\ln(k/T) = \ln(kᵦ/h) + \Delta S°/R - \Delta H°/RT
\]

**Inhibition Studies.** Slow onset inhibition was measured by the addition of enzyme to reaction mixtures containing 1 mM inosine, 50 mM Na₂HPO₄, pH 7.5 with varied inhibitor concentrations (typically 1 nM to 1 μM) at 25 °C (Figure 4). Inhibitor concentrations were determined spectrophotometrically using the molar extinction coefficients of 9540 M⁻¹ cm⁻¹ at 261 nm and 8920 M⁻¹ cm⁻¹ at 269 nm for 9-deazainosine (ImmH based inhibitors) and 9-deazaguanosine (ImmG based inhibitors), respectively (16). For analysis of slow onset tight-binding inhibition, the lowest inhibitor concentration is usually >10 times the total enzyme concentration (17). In conditions where inhibitor concentration does not exceed ten times the enzyme concentration, the effective inhibitor concentration was obtained by the expression

\[
I' = I - \left(1 - \frac{V₀}{V₀} Et\right)
\]

where I' is the effective inhibitor concentration, V₀ and V₀ are the initial rate in the presence and absence of inhibitor, and Eᵣ is the total enzyme concentration (17, 18). The Kᵦ and Kᵦ* (initial and equilibrium dissociation constants) values were determined by fitting the initial reaction rate or the
equilibrium reaction rate and inhibitor concentrations to the following expression for competitive inhibition:

\[ \frac{V_0'}{V_0} = \frac{K_m + [S]}{K_m \left( \frac{1}{K_i} + [I] \right) + [S]} \]  

(5)

where \( V_0' \) is the initial reaction rate or the equilibrium reaction rate in the presence of inhibitor, and \( V_0 \) is the initial rate in the absence of inhibitor, [I] is the inhibitor concentration, and [S] is the substrate concentration, which is held at saturating level (1 mM). The dissociation constants of ImmH, ImmG, DADMe-ImmH and DADMe-ImmG were obtained for native and mutant enzymes.

**Kinetic Isotope Effects.** The KIEs were measured for the arsenolysis of inosine as previously reported (2, 4, 7, 19, 20). Briefly, reaction mixtures contained 100 mM sodium arsenate, 50 mM Tris-HCl (pH 7.5), 250 \( \mu \)M inosine (including label) in 1 mL. 100 \( \mu \)L of the reaction mixture was resolved on charcoal-cellulose columns, and the radioactive ribose was eluted as described above. Scintillation fluid (20 mL) was added directly to the eluate, and the radioactivity was counted. The forward commitment (\( C_f \)) was added directly to the eluate, and the radioactivity was counted. The reactions were quenched after 2 s by adding 50 \( \mu \)L of 1 M HCl. 200 \( \mu \)L of the reaction mixture was resolved on charcoal-cellulose columns, and the radioactive ribose was eluted as described above. Scintillation fluid (20 mL) was added directly to the eluate, and the radioactivity was counted. The forward commitment (\( C_f \)) was calculated from the fraction of bound inosine escaping to product divided by the fraction of bound inosine escaping to catalysis at saturating arsenate (as in 19).

The intrinsic kinetic KIEs (KIE\(_{\text{int}}\)) are calculated from the experimental KIEs (KIE\(_{\text{exp}}\)) and forward commitment factor (\( C_f \)) using Northrop’s equation (eq 9) (22).

\[ \text{KIE}_{\text{exp}} = \frac{\text{KIE}_{\text{int}} + C_f}{1 + C_f} \]  

(9)

**Computational Modeling of PNP Transition State.** The transition state for the phosphorolysis of inosine was calculated in vacuo using hybrid density functional methods implemented in Gaussian 98 (23). The optimization process for the transition state stabilized by PNP was started by varying bond distances between leaving hypoxanthine or nucleophilic phosphate and the C1 of inosine. The structures of inosine (substrate), \( \alpha \)-d-ribose 1-phosphate (product) and transition state were optimized using the B3LYP functional and the 6-31G (d, p) basis set. Bond frequencies for the substrate and the transition state were calculated using the same level of theory. All vibrational modes were used for calculating kinetic isotope effects using the ISOEFF98 program (24). Frequencies of the substrate, the transition state and a reaction-coordinate imaginary frequency of 50i cm\(^{-1}\) or greater were used as the inputs. The KIEs calculated by this procedure were compared with the intrinsic KIEs obtained experimentally to give the transition-state structure most closely representing the intrinsic KIEs.

**RESULTS AND DISCUSSION**

A goal of this study is to understand how bovine and human PNP s stabilize distinct transition states despite crystallographically conserved catalytic sites. Dynamic contributions from the protein architecture remote from the catalytic sites are implicated and were explored by mutating residues that (1) differ between human and bovine enzymes and (2) are neither surface nor catalytic site residues. Site-directed mutagenesis changed Val39Thr, Asn123Lys, and Arg210Gln in BtPNP to make the structure more similar to HsPNP. Steady-state and pre-steady-state kinetic studies were used to characterize the mutant enzymes. Kinetic isotope effects were measured to determine the effect of the substitutions on the transition-state structure. Altered transition-state structure was additionally validated by altered dissociation constants with transition-state analogues.

**Steady-State and Pre-Steady-State Kinetics.** Steady-state kinetic parameters showed only small changes in \( k_{\text{cat}} \) and \( K_m \) for the double and triple mutant enzymes, compared to the native HsPNP and BtPNP (Table 2). Since the rate-limiting step of PNP s is release of purine base, changes in the transition-state structure may not be reflected in steady-state kinetics.

Pre-steady-state kinetic studies with guanosine provide a convenient spectral probe since enzyme-bound guanine

<table>
<thead>
<tr>
<th>enzyme</th>
<th>( K_m ) (( \mu )M)</th>
<th>( k_{\text{cat}} ) (s(^{-1}))</th>
<th>( k_{\text{cat}}/K_m ) (M(^{-1}) s(^{-1}))</th>
</tr>
</thead>
<tbody>
<tr>
<td>native Hs</td>
<td>40 ± 6</td>
<td>56 ± 6</td>
<td>1.4 \times 10^6</td>
</tr>
<tr>
<td>native Bt</td>
<td>32 ± 4</td>
<td>44 ± 2</td>
<td>1.4 \times 10^6</td>
</tr>
<tr>
<td>triple mutant Bt</td>
<td>38 ± 7</td>
<td>38 ± 3</td>
<td>1.0 \times 10^6</td>
</tr>
<tr>
<td>double mutant Bt</td>
<td>39 ± 6</td>
<td>25 ± 4</td>
<td>0.6 \times 10^6</td>
</tr>
</tbody>
</table>

*Values are for the phosphorolysis of inosine.*
shows strong fluorescence (the conversion of PNPKguanosine: PO4 to PNPKguanine:R-1-P). The single turnover rate (kchem) of guanosine was obtained at different temperatures for the triple mutant and parent enzymes (Table 3). The triple mutant has a slower kchem of 66 s^{-1} (25 °C) than the rates of 154 and 316 s^{-1} for HsPNP and BtPNP, respectively. This change in rate and a smaller entropic term for catalysis (TAS^E = 4.8 kJ mol^{-1}) suggests that the active sites of the mutant enzyme have transition-state features altered from both BtPNP and HsPNP (25). The activation enthalpy (ΔHE) of triple mutant PNP is 67 kJ mol^{-1}, unchanged from BtPNP, but smaller than the value of 78 kJ mol^{-1} for HsPNP. Mutations distant from the active site alter both the transition-state barrier (ΔG^‡) and its thermodynamic properties (Table 4).

Inhibition Studies. Altered binding to transition state analogue inhibitors (ImmH, DADMe-ImmH, ImmG, and DADMe-ImmG) would be anticipated if transition-state properties have been altered (5). BtPNP has an early dissociative transition state and higher affinity for ImmH, while HsPNP has a late dissociative transition state and shows higher affinity for DADMe-ImmH. The transition-state structures established for HsPNP use inosine as the substrate, which ImmH and DADMe-ImmH mimic.

Both double and triple mutant enzymes exhibited higher affinity for DADMe-ImmH than for ImmH, suggesting that the transition states of the mutant enzymes are more dissociative than that of their parent enzyme, and more closely resemble HsPNP (Table 4). Triple mutant BtPNP gave K_i values for DADMe-ImmH and ImmH similar to those of HsPNP, while the double BtPNP has lower affinity for both inhibitors. Thus, mutation of residues Asn123Lys and Arg210Gln alters interactions with transition-state analogues of BtPNP, and mutation of three residues alters BtPNP to be more like HsPNP. Both double and triple mutant BtPNPs show small decreases in affinity for ImmG and DADMe-ImmG, and both enzymes prefer DADMe-ImmG to ImmG. According to X-ray structures of BtPNP with immucillin-H and immucillin-G bound at the catalytic sites, the 9-deazaguanine group alters the position and H-bond contacts between inhibitors and Glu201 (see below). These contacts are proposed to be perturbed by the introduction of the mutations, thus causing decreased K_i values for inhibitors containing 9-deazaguanine.

Experimental Kinetic Isotope Effects. KIEs for the triple mutant BtPNP were measured on the arnosylation of inosine to solve the transition-state structure under the conditions used earlier for BtPNP and HsPNP. The product, ribose 1-arsenate, is unstable and hydrolyzes to ribose and arsenate, which renders the reaction physiologically irreversible (26). The apparent KIEs for [1^-14C], [1'-14C], [2^-14C], [5^-14C], [9^-15N] and [1'-14C, 9^-15N] with the triple mutant enzyme were measured by the competitive radiolabeled method (Table 5). [5^-14C]inosine was used as a remote control for measuring [1^-14C], [2^-14C] and [5^-14C] isotope effects, and [5^-13C]inosine was used as a remote label to measure [1'-14C], [9^-15N] and [1'-14C, 9^-15N] KIEs. The remote [1^-14C] KIE is assumed to be unity since 5^-13C is three bonds away from the reaction center and 14C is not sensitive to geometric variation in the transition state. [5^-14C]H, however, gave a 5% KIE for the triple mutant enzyme, similar to that reported for HsPNP (4). The [5^-13C]H arises from CS'-H5' bond geometric changes between unbound substrate and the enzyme-bound transition state and includes the dihedral angle H(O5')-O5'-CS'-H5' (7). The experimental [1'-14C], [9^-15N] and [1'-14C, 9^-15N] KIEs were then corrected for the [5^-13C]H KIE to give 1.028, 1.020 and 1.053, respectively.

Commitment Factor. Forward commitment to catalysis is defined as the ratio of rate constants for product formation to substrate released from the Michaelis complex (21, 22). The forward commitment of the triple mutant BtPNP is 0.243 ± 0.026, slightly greater than for BtPNP and HsPNP (0.19 and 0.15, respectively). The intrinsic KIEs were calculated by correcting the observed KIEs for the forward commitment (Table 5). The intrinsic KIEs for triple mutant BtPNP are similar to those for HsPNP, with the exception of the [1'-14C] KIE (Figure 5). Since intrinsic KIEs are related to bond geometry at the transition state, triple mutant BtPNP is characterized by a different transition state from the native enzyme.

Intrinsic [9^-15N], [1'-14C], and [1'-14C, 9^-15N] KIEs. The primary [9^-15N] KIE reports on the C1'-N9 bond order, protonation of the purine ring and aromatic bond rehybridization of hypoxanthine at the transition state. A large [9^-15N] KIE of 1.025 was obtained for the triple mutant BtPNP, close to the KIE for HsPNP (1.029). It suggests that dissociation of the N-glycosidic bond is complete and N7 is protonated at the transition state (4, 9, 19). The primary [1'-14C] is also informative for determining the nucleophilic participation in

### Table 3: Pre-Steady-State Kinetic Parameters

<table>
<thead>
<tr>
<th>parameters</th>
<th>HsPNP</th>
<th>BtPNP</th>
<th>triple mutant BtPNP</th>
</tr>
</thead>
<tbody>
<tr>
<td>TASA (kJ mol^{-1})</td>
<td>17 ± 1</td>
<td>9 ± 1</td>
<td>4.8 ± 0.3</td>
</tr>
<tr>
<td>ΔH (kJ mol^{-1})</td>
<td>78 ± 3</td>
<td>67 ± 3</td>
<td>67 ± 3</td>
</tr>
</tbody>
</table>

### Table 4: Inhibition Constants (pM) of the Native and Mutant PNPs

<table>
<thead>
<tr>
<th>analogues</th>
<th>native Hs</th>
<th>native Bt</th>
<th>triple mutant Bt</th>
<th>double mutant Bt</th>
</tr>
</thead>
<tbody>
<tr>
<td>DADMe-ImmH</td>
<td>100</td>
<td>23</td>
<td>10 ± 1</td>
<td>56 ± 8</td>
</tr>
<tr>
<td>ImmH</td>
<td>88 ± 3</td>
<td>4.7 ± 0.6</td>
<td>60 ± 11</td>
<td>462 ± 13</td>
</tr>
<tr>
<td>DADMe-ImmG</td>
<td>35 ± 1</td>
<td>35 ± 1</td>
<td>35 ± 1</td>
<td>35 ± 1</td>
</tr>
<tr>
<td>ImmG</td>
<td>9.3 ± 0.3</td>
<td>26 ± 2</td>
<td>26 ± 2</td>
<td>26 ± 2</td>
</tr>
</tbody>
</table>

### Table 5: Experimental and Intrinsic Kinetic Isotope Effects of the Triple Mutant BtPNP

<table>
<thead>
<tr>
<th>position</th>
<th>exptl</th>
<th>intrinsic</th>
<th>calc'd</th>
</tr>
</thead>
<tbody>
<tr>
<td>9^-15N</td>
<td>1.019 ± 0.005</td>
<td>1.025 ± 0.005</td>
<td>1.027</td>
</tr>
<tr>
<td>1^-14C</td>
<td>1.027 ± 0.006</td>
<td>1.035 ± 0.006</td>
<td>1.033</td>
</tr>
<tr>
<td>1^-14C, 9^-15N</td>
<td>1.048 ± 0.005</td>
<td>1.063 ± 0.005</td>
<td>1.061</td>
</tr>
<tr>
<td>1^-H</td>
<td>1.168 ± 0.006</td>
<td>1.221 ± 0.006</td>
<td>1.298</td>
</tr>
<tr>
<td>2^-H</td>
<td>1.055 ± 0.002</td>
<td>1.073 ± 0.003</td>
<td>1.057</td>
</tr>
<tr>
<td>5^-C/H2</td>
<td>1.047 ± 0.002</td>
<td>1.062 ± 0.002</td>
<td>1.101 (proR and S)</td>
</tr>
</tbody>
</table>

* Experimental KIEs are corrected to 0% substrate depletion.  
* Intrinsic KIEs are corrected for the forward commitment.
and matches a transition-state structure with C1’−OP phosphate distance equal to 2.40 Å when the dihedral to the leaving group is most favorable. Since the dihedral angle at the transition state is unknown, the [2'-H] KIE establishes significant ribooxacarbenium character at the transition state and that the dihedral to the leaving group is partially or fully aligned and is not near a perpendicular to the vacant orbital to the leaving group.

The remote [5'-3H] KIE is normally expected to be unity in N-ribosyltransferases, since the 5'-3H is four bonds away from the reaction center. However, a large [5'-3H] KIE has been observed for human, bovine, and Plasmodium PNPs (2, 4, 7, 19). Significant distortions in the vibrational modes of the 5'-hydrogens are required to generate the [5'-3H] KIE and originate primarily from 5'-hydroxyl dihedral angle anchoring at the transition state (7). Computational modeling indicates that isotope effects as large as 10% can be caused by altering the O5'-C5'-C4'-O4' dihedral angle from −55° for free inosine to 60° at the transition state. The triple mutant BtBNP is similar to the human protein with a large [5'-3H] KIE of 1.062. Thus, the triple mutant BtBNP has similar geometrical distortion at C5' as does the human PNP.

**Transition-State Structure.** The transition-state structure of the triple mutant BtBNP was optimized at the B3LYP/6-31G (d, p) level of theory by systematically varying N-ribosidic and OP phosphate-ribosidic bond distances as well as the 2'-C-exolendo geometry. The calculated transition-state structure of the triple mutant BtBNP has the N-ribosidic bond completely broken (C1’−N9 distance ≥ 3.0 Å) in the transition state. There is nucleophilic phosphate participation at C1’ with C1’−OP phosphate = 2.26 ± 0.02 Å. N7 of the leaving group hypoxanthine is fully protonated, and the ribosyl pucker ring adopts a 2'-C-exo conformation (Figure 6 in ref 19). This transition state is within experimental error of that reported for the K22E:H104R mutant of human PNP (19).

Reactions with an intermediate have two transition states, and in two-transition-state systems (in the case of PNPs, N-ribosidic bond breaking, carbocation intermediate and phosphate attack) both transition states can contribute to intrinsic KIEs when their energetic barriers are similar (27). The value of the [1'-14C] KIE is greater with triple mutant BtBNP than for HsBNP or native BtBNP and together with the relatively large [9-15N] KIE indicates that the phosphate attack contributes to the transition state. This transition state lies near product on the reaction coordinate since phosphate attack has begun and N-ribosidic bond breaking is complete.

With the exception of the [1'-14C] KIE, the triple mutant BtBNP exhibited KIEs more similar to those of the human than to native bovine PNP. Transition-state analysis reveals that the triple mutant BtBNP has a transition state later in the reaction coordinate than native BtBNP and even later in the reaction coordinate than HsBNP, demonstrating significant nucleophilic participation at C1’.

Although the literature is replete with examples of kinetic properties being altered by remote mutations, this work is a departure from other transition-state analyses in demonstrating that three residues more than 1 Å from the active site can alter the BtBNP transition state to be more like HsBNP with little change of the steady-state kinetic parameters. Remote mutations may alter H-bond patterns that could couple conformational or dynamic changes into the catalytic site and thereby change the transition-state structure. This
work is consistent with the hypothesis that mutation of BtPNP toward HsPNP has altered the transition-state structure by changing the dynamics of the protein, or by mechanisms linked to transition-state conformational changes or by altered pKa values of groups linked to the catalytic site. The results demonstrate a surprising malleability of enzymatic transition-state structure while maintaining the catalytic rate enhancement.

CONCLUSION

Residues 39, 123 and 210 of BtPNP are more than 11 Å away from the active site and differ substantially between human and bovine PNPs. Mutant enzymes were obtained by mutating two or three of these residues in BtPNP to their counterparts from HsPNP. Steady-state kinetic studies showed only small differences between the mutant and parental enzymes, although pre-steady-state kinetic studies yielded a slower $k_{\text{chem}}$ for the triple mutant BtPNP. Changes in the triple mutant BtPNP transition state were suggested by altered affinity to DADMe-InmmH, which mimics the transition-state structure of HsPNP. Transition-state analysis from KIEs and computational chemistry established that the triple mutant BtPNP has a transition-state structure with the N-ribosidic bond completely broken and significant nucleophilic attack at C1\'. The energy barrier of nucleophilic attack is higher than it is in human PNP, which defines this step as the primary transition-state barrier and slows the chemical step. The transition-state structure for triple mutant BtPNP is much later in the reaction coordinate than for native BtPNP and is even later that the fully (or equilibrating) $S_\text{N}1$ transition state of HsPNP. Mutations of BtPNP toward HsPNP altered the transition-state structure to be more like the human enzyme. Even remote amino acids contribute to the transition-state structure of PNPs, likely through dynamic motions that form the transition state.

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SUPPORTING INFORMATION AVAILABLE

Table of transition state structure data. This material is available free of charge via the Internet at http://pubs.acs.org.

REFERENCES