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Abstract: Phosphomevalonate kinase (PMK) catalyzes an essential step in the mevalonate pathway, which is the only pathway for synthesis of isoprenoids and steroids in humans. PMK catalyzes transfer of the γ-phosphate of ATP to mevalonate 5-phosphate (M5P) to form mevalonate 5-diphosphate. Bringing these phosphate groups in proximity to react is especially challenging, given the high negative charge density on the four phosphate groups in the active site. As such, conformational and dynamics changes needed to form the Michaelis complex are of mechanistic interest. Herein, we report the characterization of substrate induced changes (Mg-ADP, M5P, and the ternary complex) in PMK, using NMR-based dynamics and chemical shift perturbation measurements. Mg-ADP and M5P Kd’s were 6-60 μM in all complexes, consistent with there being little binding synergy. Binding of M5P causes the PMK structure to compress (τc= 13.5 nsec), while subsequent binding of Mg-ADP opens the structure up (τc= 17.6 nsec). The overall complex seems to stay very rigid on the psec-nsec timescale with an average NMR order parameter of S2≈0.88. Data are consistent with addition of M5P causing movement around a hinge region to permit domain closure, which would bring the M5P domain close to ATP to permit catalysis. Dynamics data identify potential hinge residues as H55 and R93, based on their low order parameters and their location in extended regions that connect the M5P and ATP domains in the PMK homology model. Likewise, D163 may be a hinge residue for the lid region that is homologous to the adenylate kinase lid, covering the
“Walker-A” catalytic loop. Binding of ATP or ADP appears to cause similar conformational changes; but, these observations do not indicate an obvious role for γ-phosphate binding interactions. Indeed, the role of γ-phosphate interactions may be more subtle than suggested by ATP/ADP comparisons, since the conservative O to NH substitution in the β-γ bridge of ATP causes a dramatic decrease in affinity and induces few chemical shift perturbations. In terms of positioning of catalytic residues, binding of M5P induces a rigidification of Gly21 (adjacent to the catalytically important Lys22), although exchange broadening in the ternary complex suggests some motion on a slower timescale does still occur. Finally, the first 9 residues of the N-terminus are highly disordered, suggesting they may be part of a cleavable signal or regulatory peptide sequence.

**Keywords:** Phosphomevalonate kinase, chemical shift perturbation, relaxation dynamics, mevalonate, NMR, Modelfree

**Introduction**

Protein motions are critical for various biological functions and occur over a wide range of timescales. There is a strong correlation between structural dynamics and molecular function, and studying these dynamic processes can give site-specific information on motions that span timescales from psec to msec\(^1\). One class of enzymes that undergoes large structural and associated motional changes are the kinases. These changes occur due to the binding of their two ligands, and the need for reactive groups to move into close proximity for catalysis. A model kinase in this regard is adenylate kinase, which consists of two domains that adjust to bring their phosphate donor and acceptor ligands into proximity for reaction to occur \(^2, 3\).

Phosphomevalonate kinase (PMK), the topic of this study, catalyzes a reaction analogous to that of adenylate kinase in that the phosphate acceptor is itself a phosphate group. Human PMK catalyzes the transfer of the γ-phosphoryl group of ATP to mevalonate 5-phosphate (M5P), resulting in formation of mevalonate 5-diphosphate. This is the sole pathway for biosynthesis of steroids and other isoprenoids in mammals\(^4-7\). Despite the importance of this pathway in human heart disease and the fact that it is the biosynthetic source of a diverse class of metabolites, the mechanism of PMK is not completely characterized\(^8\), although recent studies have identified important catalytic residues\(^9, 10\). To better understand the molecular mechanism...
of catalysis, structural characterizations of protein-ligand complexes are required. Of particular interest are conformational and dynamic changes that are needed to bring the two highly charged substrates in proximity to react. To facilitate this, a homology model for human PMK had been generated using the PHYRE homology model method (≈24% overall sequence identity), since there is no experimental structure yet available for this protein\(^9\). Our recent NMR chemical shift assignments and subsequent secondary structure analysis has largely validated this model\(^11\), based on chemical shift index values that match helical and sheet regions in the model. But, the presence of disordered regions and loops has made it impossible to pursue a complete structure determination. Currently, 77% of the backbone atoms have been assigned, excluding two unassigned gaps that may be disordered regions (residues 56-72 and 99-111). As such, studies herein refer to the homology model. This model consists of two domains. The larger ATP-binding domain is comprised of a 5 stranded parallel \(\beta\)-sheet interweaved with 3 \(\alpha\)-helices. The smaller M5P domain is comprised of loop regions and two \(\alpha\)-helices. In the model, the ATP and M5P domains are tethered by two hinge regions, analogous to adenylate kinase. These may be involved in opening and closing motions, to permit binding and release of substrates\(^3\). Also, recent side-directed mutagenesis studies have identified a “Walker A” ATP binding motif (K17, R18, K19, K22)\(^12\).\(^13\). Mutagenesis of some of these residues can decrease catalysis up to 10,000-fold, suggesting this as the active site location for phosphoryl transfer\(^9\),\(^10\). “Walker-A” loop residues most important for catalysis include K22 and R18. Outside the “Walker-A” catalytic loop, R110 is also important \(^9\),\(^10\).

To characterize the effects of substrate addition on protein dynamics, we have used NMR relaxation experiments in conjunction with a Modelfree analysis\(^14\) to quantify any changes to the fast timescale motion (psec-nsec) of the protein backbone, especially for catalytically relevant regions. We also report on the conformational effects of substrate addition, by comparing chemical shift changes and backbone \(^15\)N relaxation data for various complexes of PMK: (a) Apo-PMK, (b) the binary complex with Mg-ADP, (c) the binary complex with M5P, and (d) the ternary complex with Mg-ADP and M5P.
Materials and Methods

Docking of M5P and ATP into Human PMK Homology Model

The homology model used in our studies was that which was previously prepared and reported by Herdendorf and Miziorko. The binding orientation of ligands in this PMK (human) homology model was determined using Autodock4. Gasteiger charges and hydrogens were added using AutoDock Tools (ADT). The docking grids were also prepared using ADT, with a grid size of 60 × 60 × 60 Å and a spacing of 0.375 Å. These grids were centered on the ligands (AMP and GMP) of the homology model template, bacteriophage T4 deoxynucleotide kinase. The homology model and template were superimposed on each other, in order to obtain coordinates for the grid box used to dock the ligands. The overlay was done using Sybyl 6.8 (Tripos Inc., St Louis, MO, USA). Default docking parameters were used, except that 50 genetic algorithm runs were used with 2,500,000 as the maximum number of evaluations. The docking of both ligands was done separately, due to inter-ligand repulsion from the negative charges on M5P and the ATP triphosphate, even after neutralization of two charges (as would be the case with bound Mg²⁺). This repulsive interaction was recognized based on earlier docking attempts to form the ternary complex, which produced erroneously distorted structures with the phosphate groups on the two ligands as far apart as possible in the docking box. This led us to do independent docking of ATP and M5P to produce the separate binary complexes, and these pdb files were then merged to create the ternary complex. This approach positioned the phosphate groups on ATP and M5P in close proximity, as required for the phosphate transfer reaction, and therefore provides strong validation for the docking poses that were generated (Fig. 1).
Figure 1 Ternary structure of docked ligands ADP and M5P in the active site, near the "Walker A" loop, of PMK. Catalytically important amino acids are labeled, including Lys's 17, 19, and 22, as well as Arg's 18 and 110.

Protein Expression and Purification

PMK is normally a 192 residue 22.0 kDa protein, although it was expressed with an N-terminal histidine tag and additional linker residues, giving a total molecular weight of 24.2 kDa. Uniformly $^{15}$N labeled PMK samples were prepared using $^{15}$NH$_4$Cl as the sole source of nitrogen in minimal media and protein was expressed and purified as described previously$^{9,10}$. Briefly, *E. coli* BL21-(DE3) Rosetta cells were transformed with a pET15b(+) expression construct which encoded human PMK with an N-terminal His$_6$ affinity tag. The transformed cells were plated onto LB (Luria Bertani) agar containing ampicillin (amp) and chloramphenicol (chl). Plates were incubated overnight at 37 °C, and a single colony was picked to inoculate 2 mL of media, and grown to $A_{600} \sim 0.3$. This culture was then used to inoculate 20 LB-amp-chl plates. The plates were incubated overnight at 37 °C, and resulting lawns were used to inoculate 500 mL of LB-amp-chl to give $A_{600} \sim 1.0$. The liquid culture was then incubated at 30°C for 1 h prior to induction with 1 mM IPTG. The culture was harvested 4 h post induction at $A_{600} \sim 2.0$. Bacterial pellets were resuspended in 100 mL of a 50 mM KP$_i$ (potassium phosphate) buffer containing 100 mM KCl, 5 mM imidazole, and 0.5 mM DTT at pH 7.8.
Lysis was accomplished by passage through a microfluidizer at \( \sim 17 \) kpsi. The lysate was clarified by centrifugation at \( \sim 100,000 \) g and the supernatant was loaded onto \( \sim 0.5\text{-}1.0 \) mL of Ni-Sepharose Fast Flow resin. The column was washed with lysis buffer until \( A_{280} < 0.005 \), and the protein was eluted with lysis buffer supplemented with 300 mM imidazole. The fractions containing PMK were pooled and the concentration was determined spectrophotometrically using an extinction coefficient of \( \varepsilon_{280} = 32,290 \text{ M}^{-1} \text{ cm}^{-1} \).

**NMR Sample Preparation**

All protein samples were buffer exchanged, using ultrafiltration with an Amicon (YM10) membrane, into 5 mM DTT, 20 mM KH\(_2\)PO\(_4\), 10\% D\(_2\)O, and 0.02\% NaN\(_3\), and concentrated to 400-600 \( \mu \)M. ADP was complexed with Mg\(^{2+}\) by adding a 1:1 ratio of MgCl\(_2\). Mg-ADP and M5P were both concentrated to 50 mM prior to titration, with pH's of 5.0 for ADP and 7.0 for M5P. All experiments were performed at pH 6.5.

**NMR Spectroscopy**

All NMR experiments were performed on a 600 MHz Varian NMR System at 599.515 MHz using a triple resonance cryoprobe with z-axis gradients at 25 °C. Titrations were performed using 100 \( \mu \)M increments for both ligands, until saturation was achieved based on \(^1\)H-\(^{15}\)N HSQC chemical shift changes (1 mM for M5P and 2 mM for MgADP). There were two sets of titrations performed, first starting with M5P then adding Mg-ADP. To assess binding order and synergy, the second titration started with Mg-ADP followed by M5P. \(^{15}\)N relaxation experiments were performed on free PMK, PMK saturated M5P (2 mM), PMK saturated Mg-ADP (20 mM; higher concentration was needed due to slow hydrolysis of ADP), and then the ternary complex (using the same saturation concentrations). \(^{15}\)N-\( T_1 \), \(^{15}\)N-\( T_2 \), and \( \{^1\)H\}-\(^{15}\)N NOE experiments were all performed using the BioPack pulse sequences from Varian, Inc (Palo Alto, CA). Delay times for the \( T_1 \) experiment were 10.8, 108.3, 216.8, 379.2, 541.7, 758.4, 1083.4, 1516.8, and 2166.8 ms and 4.31, 8.62, 12.9, 17.2, 21.6, 30.2, 38.8, 47.4, 56.1 ms for \( T_2 \). \( T_1 \) experiments employed the standard inversion-recovery pulse sequence\(^{17} \), while \( T_2 \) experiments employed the CPMG
sequence, as implemented previously\textsuperscript{18,19}. NOE's were obtained by measuring HSQC spectra with and without \textsuperscript{1}H saturation for a time of 3 s (same for both), and using an interscan delay of 1 sec. An interleaved approach was used for the \textit{T}_1 and \textit{T}_2 experiments, to average out any potential instability over time\textsuperscript{20,21}.

### NMR Spectroscopy Data Analysis

NMR data were processed using NMRPipe/NMRDraw\textsuperscript{22}, and analyzed using NMRView\textsuperscript{23}. For all experiments, \textit{\textsuperscript{1}H-\textsuperscript{15}N} spectra were processed using a 90° shifted sine function in the \textit{\textsuperscript{1}H} and \textit{\textsuperscript{15}N} dimensions. For \textit{\textsuperscript{15}N-\textit{T}_1} and \textit{\textsuperscript{15}N-\textit{T}_2} experiments, the spectra with the shortest relaxation delay were peak picked using NMRView. For \{\textit{\textsuperscript{1}H}\}-\textit{\textsuperscript{15}N} NOE measurements only one spectrum was peak picked, and for each subsequent spectrum the peak ellipses were manually adjusted to fit each peak.

The \textit{\textit{R}_1} and \textit{\textit{R}_2} relaxation rates were determined by fitting the \textit{\textit{T}_1} and \textit{\textit{T}_2} curves to Equation 1:

\[
I_t = I_0 \cdot e^{(-Rt)}
\]

where \(I_t\) is the intensity after time \(t\), \(I_0\) is the intensity at time \(t=0\), and \(R\) is either \(\textit{\textit{R}_1}\) or \(\textit{\textit{R}_2}\). Fitting was done using the Rate Analysis package included in NMRView. NOE values were obtained by taking the ratio of the intensity versus the control. This was done with two sets of experiments in order to obtain an error for the analysis. \textit{\textit{T}_1} and \textit{\textit{T}_2} values were measured for PMK at various concentrations and used to calculate \(\tau_c\) values, to demonstrate that PMK remains monomeric under the conditions of our NMR studies (previous analytical gel filtration studies also established that PMK is monomeric\textsuperscript{9}).

Dissociation constants were obtained by measuring the chemical shift changes in going from free to various bound states, monitoring peaks in fast exchange in both \textit{\textsuperscript{1}H} and \textit{\textsuperscript{15}N} dimensions. These changes were then combined using Equation 2:
The chemical shift change ($\Delta \text{shift}_{\text{obs}}$) along with the concentration of the ligand and protein were fitted to a quadratic equation, to determine the dissociation constant $K_d$:

$$\Delta \text{shift}_{\text{obs}} = \left( \frac{\Delta \text{shift}_{\text{max}}}{(2 [P_o])} \right) \left\{ ([L_o] + [P_o] + K_d) \\
- ([L_o] + [P_o] + K_d)^2 - 4 [L_o] [P_o] \right\}^{1/2}$$

using Sigmaplot8, where $L_o$ is the total ligand concentration at a particular titration point, $P_o$ is the total protein concentration, and $\Delta \text{shift}_{\text{max}}$ is the maximum chemical shift change observed for the particular peak in question. Fitting to the quadratic equation was required because $[P_o] > K_d$. Standard deviations resulted from the non-linear least squares fitting process (note: as $K_d$ gets very small relative to $[P_o]$, errors will become larger because $K_d$ become less well-defined in the fitting process).

**Modelfree Analysis**

The parameters of internal motion were determined from the NMR relaxation data according to the model-free formalism established by Lipari and Szabo\textsuperscript{24-26} using Modelfree4 software (version 4.20, A. G. Palmer, Columbia University). The residues were optimized with an isotropic diffusion model using an initial estimate derived from the R2R1_diffusion program (A. G. Palmer, Columbia University). Backbone dynamics calculations were performed with 300 Monte Carlo simulations per run using an internuclear distance $r_{\text{NH}}$ of 1.02 Å and a chemical shift anisotropy (CSA) for the $^{15}$N nucleus of $-172$ ppm. Five models were used to fit our experimental data and were iteratively tested in order of increasing complexity ($M1 = S^2$; $M2 = S^2, \tau_e$; $M3 = S^2, R_{ex}$; $M4 = S^2, R_{ex}, \tau_e$; $M5 = S^2_{ef}, S^2_s, \tau_e$) until an acceptable fit was achieved. $S^2$ is the generalized order parameter, $\tau_e$ is the internal correlation time, $R_{ex}$ is the exchange contribution term, and $S^2_{ef}$ and $S^2_s$ are for sub-nanosecond and nanosecond motions respectively. These models were tested until they could reproduce the experimental
relaxation data within 90% confidence limits using appropriate $\chi^2$ and F-tests$^{14}$.

**Results**

**Docking of M5P and ATP to Human PMK**

The docking of M5P and ATP (Fig. 1) seems reasonable based on the proximity of the reactive phosphate groups to each other, and also their location within 4.0 Å of the “Walker A” catalytic loop, which includes Lys17, Arg18, Lys19, and Lys22 as well as other basic residues, especially Arg110. It is striking that the independent docking of M5P into PMK positioned its phosphate group within 2.7 Å of the ATP γ-phosphate and 2.8 Å from Arg110 and Arg138.

**Chemical Shift Perturbation Studies to Make the Binary and Ternary Complexes**

For chemical shift perturbation studies, a dead-end complex was created to ensure that there would be no catalytic turnover during the NMR measurements. AMP-PNP (Fig. 2) is often used to create such dead-end complexes in kinases, but it binds only weakly to PMK and causes few structural changes in PMK compared to ATP (Fig. 3a and supplementary materials) or ADP (Fig. 3c and supplementary materials). As a result, ADP was used to make the dead end complex (Fig. 3c) in our studies. To justify the use of ADP as a suitable replacement for ATP, an ATP titration experiment was carried out and is shown in Fig. 3a. Comparing these spectra, it is clear that they show the same chemical shift changes for the same residues, which infers that the two protein-ligand complexes undergo very similar structural changes. From the HSQC spectra for the Mg-ADP titration (Fig. 3c), a pronounced conformational change is observed upon substrate addition, due to multiple residues that are affected when the complex is formed. Fig. 4a shows the chemical shift perturbations due to Mg-ADP binding mapped onto the PMK model, and is consistent with Mg-ADP causing the protein to undergo a gross conformational change; that is, the whole protein appears to adjust structurally in order to accommodate the addition of Mg-ADP. Notable changes are observed in the loops near the substrate phosphate groups, as well as the region
where the two ATP-domain helices contact each other. The Mg-ADP addition was then followed by further titration with M5P to form the ternary complex (Fig. 4b). To determine if there is a preferred sequence for binding, due to synergy, M5P was then titrated first to Apo-PMK (Fig. 4c).

![Structures of adenine nucleotides ATP, AMP-PNP, and ADP with corresponding K_d values.](image)

**Figure 2** Structures of adenine nucleotides ATP, AMP-PNP, and ADP with corresponding K_d values.

![Chemical shift perturbations due to binding of nucleotides a) Mg-ATP titrated to saturation onto Apo-PMK, b) Mg-AMP-PNP titrated to saturation onto Apo-PMK, and c) Mg-ADP titrated to saturation onto Apo-PMK. All experiments begin at the](image)

**Figure 3** Chemical shift perturbations due to binding of nucleotides a) Mg-ATP titrated to saturation onto Apo-PMK, b) Mg-AMP-PNP titrated to saturation onto Apo-PMK, and c) Mg-ADP titrated to saturation onto Apo-PMK. All experiments begin at the
red cross-peaks, with additions in 100 μM increments to saturation, which is indicated by the transition to the blue cross-peaks. The corresponding chemical shifts are mapped on to the human PMK homology model, where the red indicates a large chemical shift change (>0.09 ppm) and pink indicates a medium chemical shift change (0.05-0.09 ppm), while blue indicates a small or no chemical shift change (or no data).

**Figure 4** Chemical shift perturbations of: a) Mg-ADP added to saturation to Apo-PMK b) M5P added to saturation to Mg-ADP/PMK c) M5P added to saturation to Apo-PMK and d) Mg-ADP added to saturation to PMK/M5P. All experiments begin with the red cross-peaks and titrations occur at 100 μM increments to saturation, which is indicated by transition to the blue cross-peaks. The corresponding chemical shift changes are mapped on to the human PMK homology model, where the red indicates a large chemical shift change (>0.09 ppm) and pink indicates a medium chemical shift change (0.05-0.09 ppm) while blue indicates small or no chemical shift change (or no data). Note: Gly21 is visible in panel (a) at a lower threshold.

**Binding Affinity and Synergy**

To quantitatively assess binding affinity and binding synergy, if any, cross-peak perturbations (Figs. 3 and 4) were fitted to obtain $K_d$ values for each binding event. Monitoring chemical shift changes and fitting to the quadratic equation (Eq. 3) permitted
determination of dissociation constants \( (K_d) \) for the various nucleotides (Fig. 2) and for all four complexes (Fig. 5). The \( K_d \) for the Mg-ADP titration to Apo-PMK is 19 \( \pm \) 9 \( \mu \)M and <20 \( \mu \)M for subsequent M5P binding to form the ternary complex. For the titration of M5P to Apo-PMK the \( K_d \) is 6 \( \pm \) 3 \( \mu \)M and 56 \( \pm \) 16 \( \mu \)M for the subsequent Mg-ADP addition to form the ternary complex. These values are consistent with the previously reported \( K_m \) values of 47 \( \pm \) 5 \( \mu \)M for ADP and 34 \( \pm \) 3 \( \mu \)M for M5P \(^9\), keeping in mind that \( K_m \) values can often deviate from \( K_d \) values due to kinetic effects.

![Diagram](https://example.com/diagram.png)

**Figure 5** Thermodynamic box for the formation of the dead-end ternary complex of human PMK with corresponding \( K_d \) and \( \tau_c \) values for each complex. Cartoon representations of each complex are shown, with the circles representing M5P and the squares representing ADP.

**Relaxation Dynamics**

To obtain dynamic information on the psec-nsec time scale, longitudinal (\( R_1 \)) and transverse (\( R_2 \)) relaxation rates as well as \( ^{1}H \)–\( ^{15}N \) NOE values were obtained, and are summarized in Fig. 6 for Apo-PMK, Mg-ADP bound PMK, M5P bound PMK, and the ternary (Mg-ADP/M5P) complex. All of the complexes have relatively rigid structures (high \( S^2 \)), indicative of a well-ordered protein backbone on the psec-nsec timescale. Trends in \( R_1 \) and \( R_2 \) values in Table 1 can best be explained based on changes in the overall correlation time of the protein in the different complexes. It should again be noted that
PMK remains in a monomeric state in our NMR studies (Supplementary material), so changes in correlation time are not associated with aggregation effects. The changes in PMK correlation time were verified and quantified in the Modelfree analysis, where it was found that relaxation data could not be properly fitted for the different PMK complexes using the same correlation time. The correlation times obtained from the Modelfree analysis of each complex are summarized in Fig. 5. These correlation time changes are consistent with there being ligand-induced change in overall shape of PMK in the different complexes, as had been previously observed in analogous NMR studies of adenylate kinase.

Figure 6 NMR dynamics data including $R_1$, $R_2$, NOE, and $S^2$ values, where blue is the Apo-PMK complex, red is the M5P saturated complex, yellow is the Mg-ADP saturated complex, and green is the ternary complex.
Table 1 Average Values of Relaxation Parameters for PMK Complexes

<table>
<thead>
<tr>
<th></th>
<th>Apo-PMK</th>
<th>PMK-M5P</th>
<th>PMK-ADP</th>
<th>PMK-ADP-M5P</th>
</tr>
</thead>
<tbody>
<tr>
<td>$R_1$ (s$^{-1}$)</td>
<td>0.945 ± 0.034</td>
<td>0.994 ± 0.046</td>
<td>0.856 ± 0.051</td>
<td>0.933 ± 0.036</td>
</tr>
<tr>
<td>$R_2$ (s$^{-1}$)</td>
<td>20.7 ± 1.9</td>
<td>17.3 ± 1.9</td>
<td>24.0 ± 2.0</td>
<td>20.0 ± 1.7</td>
</tr>
<tr>
<td>NOE</td>
<td>0.820 ± 0.080</td>
<td>0.766 ± 0.028</td>
<td>0.834 ± 0.065</td>
<td>0.818 ± 0.112</td>
</tr>
<tr>
<td>$S^2$</td>
<td>0.881 ± 0.035</td>
<td>0.859 ± 0.046</td>
<td>0.891 ± 0.036</td>
<td>0.873 ± 0.036</td>
</tr>
</tbody>
</table>

The studies reported herein are directed to defining the structural and dynamics changes in human PMK, induced by ligand binding events. Studies were based on human PMK that was titrated with MgADP and M5P, and interpreted in light of a previously reported homology model. Results can be interpreted in the context of extensive studies of related proteins, such as adenylate kinase which binds ATP and the substrate to be phosphorylated in two separate domains, and which also contains a “Walker A” catalytic loop capped by a flexible “lid” region.

The docked complex – a structural hypothesis

In our docked complex of human PMK with M5P and ATP (Fig. 1), the key catalytic residues identified by Herdendorf and Miziorko$^{9,10}$ (Arg18, Lys22, and Arg110) are all reasonably well positioned for catalysis. The only possible exception is Lys22, which is in the active site but is pointed away from the phosphate; while this may reflect an inaccuracy of the homology model, it is noteworthy that a simple side chain rearrangement could easily position its ε-amino group near the ATP γ-phosphate. Arg18, Lys22, and Arg110, along with the other basic residues in Fig. 1, may be important for forming the ternary complex by neutralizing the negative charges on the two substrates, so that their phosphate groups can move into proximity for phosphoryl transfer. Additional charge stabilization (from Lys22?) may also be needed to stabilize the transition state if it goes by an associative-type mechanism, which would place added charge density on the γ-phosphate. An important feature of the model in Fig. 1 is that the γ-
phosphate of ATP is ideally positioned for nucleophilic attack by the M5P phosphate (separated by only 2.7 Å). This provides further validation that this structural model of the ternary complex is reasonably accurate.

**Binding synergy ... or lack thereof**

Based on chemical shift perturbations, it appears that binding of either M5P or ADP to apo PMK can induce a large conformational change. To quantitatively assess binding synergy, if any, cross-peak perturbations were fitted to obtain $K_d$ values for each binding event (Fig. 5). These $K_d$ values are consistent with their being no preferred binding sequence or synergy for the ligands. This is in contrast to the report of an ordered sequential mechanism (M5P binding first and ADP released last) for the pig liver PMK enzyme\textsuperscript{27}, but is consistent with recently reported steady state kinetics on human PMK, where competitive inhibition was observed using mevalonate 5-diphosphate as a product inhibitor versus M5P\textsuperscript{10}. If anything, our data suggest there is a modest (~3-fold) anti-synergy, such that affinity for the second ligand is weakened due to presence of the first, which could be due to the charge/charge repulsion by the phosphate groups on both ligands. But, this anti-synergy could be unique to our dead-end complex (MgADP/M5P), and does not necessarily reflect the situation for the catalytically competent Michaelis complex (MgATP/M5P)\textsuperscript{28}.

**Ligand-induced structural changes**

Based on the chemical shift perturbations that occurred during titration to form binary and ternary complexes, it can be seen that the largest changes occur in forming the binary complex, whether it is ADP or M5P that binds first (Fig. 4). The chemical shift changes include residues outside the binding site, such as hinge regions between domains, so suggest a conformational change has occurred. In terms of adenine nucleotide binding, it is interesting that while the $\gamma$-phosphate of ATP seems to not be important for binding or inducing structural changes, based on comparisons of ATP and ADP (Figs. 2 and 3), the situation is actually more complicated - because simple O=\textgreater NH substitution of the bridging heteroatom between the $\beta$ and $\gamma$ phosphates has a dramatic effect on binding. In
fact, independent kinetic measurements performed at 34 μM ATP (< 1/3rd $K_m$ for ATP) indicates no inhibition of PMK by AMPNP at levels ranging from 269-1074 μM (Miziorko and Herdendorf, unpublished results). This suggests there may be important interactions with the β-γ bridging heteroatom, which might be important for stabilizing this leaving group during phosphate transfer. It is also noteworthy that ADP (and ATP) binding causes chemical shift perturbations in the Walker A loop (ex. Arg18) as well as the lid that covers the Walker A loop, and a hinge residue for the lid (D163). Opening of this lid motion should only produce a modest change in size of the protein, and so would not be expected to produce a change in correlation time. In contrast, domain movement between ATP and M5P binding domains would be expected to produce a much larger change in size and correlation time, as had been observed in adenylate kinase, and appears to also occur in human PMK (vide infra).

**Ligand-induced changes to psec-nsec timescale dynamics ... or lack thereof**

Based on Modelfree analysis of NMR dynamics parameters (Fig. 6), generalized order parameters were obtained. These clearly indicate that PMK adopts a fairly rigid structure ($S^2$ values >0.8) for all the binary and ternary complexes. Furthermore, the average $S^2$ values remain largely unchanged upon formation of the binary complex with either ADP or M5P, indicating that the local dynamics for all the complexes are virtually the same (Table 1) on the psec-nsec timescale. The large $S^2$ values, which are consistent with a rigid, highly ordered backbone, are not unexpected given the well-structured core of the protein, which includes a 5 stranded parallel β-sheet. One might have expected some ordering in the M5P domain upon ligand binding, given the high content of extended and loop regions, but this does not appear to be occurring.

**Inter-domain and loop motion: ligand binding domains, lid and “Walker A” catalytic loop**

While the protein core may be largely rigid, it is possible that there is some motion of domains relative to each other, and of other defined regions. Indeed, chemical shift perturbation studies suggested
there was a large substrate-induced conformational change (Fig. 4). In fact, there were a small number of residues with small or negative NOE's that gave rise to small $S^2$ values (H55, R93, and D163), and these are located in loop and potential hinge regions in the homology model (*vide infra*). Interestingly, the “Walker A” ATP binding loop shows the same $S^2$ values as the rest of the protein (i.e. it is rigid), indicating that if there is any catalytically relevant change in dynamics, it might only occur during turnover with the actual Michaelis complex\textsuperscript{28}, rather than with the “dead end” complex used in our studies. One important exception is Gly21, which is adjacent to Lys22, the catalytically most important residue in PMK\textsuperscript{9}. Gly21 undergoes an increase in $S^2$ in going from the apo (blue) to the M5P (red) complex, indicating it becomes less mobile in the binary complex (Fig. 6), perhaps as Lys22 moves into position for catalysis (note: Lys22 was not assigned, so changes to its dynamic state are not known). Interestingly, Gly21 exchange broadens in the ternary complex, suggesting it undergoes msec timescale motion. But, in the conversion of apo PMK to the PMK-M5P binary complex, one can see a slow exchange process for Arg18, and a fast exchange process for Gly21 (Fig. 4c, and expansions in supplementary materials). Assuming the same motion occurs for these two “Walker A” loop residues, this timescale flanking permits an estimation of the rate constant for the “Walker A” catalytic loop exchanging between the two states associated with M5P binding as \( k_{ex} = 100-600 \text{ sec}^{-1} \) (see supplementary materials). This is likely to be due to a lid opening/closing motion, which would be coupled to “Walker A” loop motion, based on analogy to adenylate kinase where a flexible lid caps the catalytic “Walker A” loop\textsuperscript{31-36}.

It should be noted that while there doesn't appear to be significant motion of “Walker A” catalytic loop residues on the psec-nsec timescale (besides Gly21), there does appear to be motion on slower (ex. msec) timescales. For example, in the ADP binary and ternary complexes, most crosspeaks for residues in the loop were no longer visible, due to exchange broadening (Fig. 4). For this reason, $S^2$ for ADP complexes could not be measured. Accordingly, it is simply not known in these complexes what motion might be occurring on the psec-nsec timescale.
Overall, the psec-nsec timescale dynamics data indicate that M5P binding drives the immobilization of catalytic residues (with Gly21 as our “Walker A” reporter), but there still appears to be msec timescale motion, especially when ADP is bound. This motion might be important for catalysis, but could also be unique to the particular dead-end inhibitory complex used in our study. There is certainly literature precedent for kinase active site loop dynamics being very sensitive to the nature of the inhibitory complex. For example, residual dipolar coupling and $^{15}$N relaxation measurements of various ABL kinase complexes indicated that inhibitors differed significantly in their ability to immobilize the catalytic/activation loop.$^{30}$

**Ligand-induced changes to protein size: inter-domain conformational change?**

The chemical shift perturbation studies indicated that binding of either ADP or M5P may cause a large conformational change (Fig. 4), but the above analysis of $S^2$ values indicated this did not correspond to any significant changes in dynamics within the two ligand binding domains, other than some changes to the “Walker A” catalytic loop, and the lid that covers it. What about inter-domain motion? The Modelfree analysis provided some insights in this regard. As mentioned in the Methods section, relaxation data fitting was optimized for each complex to find the rotational correlation time $\tau_c$ for that complex, since it was not possible to fit all complexes using a single $\tau_c$. Changes to fitted $\tau_c$ values for all the complexes (Fig. 5) reveal that the ADP binary complex has the largest $\tau_c$ of 17.6 nsec while the M5P binary complex has the smallest, with a $\tau_c$ of 13.5 nsec. The Apo-PMK and the ternary complex fall in between these two extremes, with similar $\tau_c$ values of $\sim$15.7 nsec (Fig. 5). This suggests that both ADP and M5P binding induce conformational changes, but in opposite directions. That is, M5P causes a compression of the structure (so PMK tumbles faster), while ADP causes an opening up of the structure (so PMK tumbles slower). Such changes would be consistent with a movement of the two ligand binding domains relative to each other, as occurs in the nucleoside monophosphate kinases$^{29,31-35}$.

Why ADP binding causes the structure to open up is not clear. Since the product of the PMK reaction is ADP, it could be that the open
conformation reflects the protein trying to release ADP. However, the chemical shift perturbation study with ATP (Fig. 3a) shows the same chemical shift perturbations as with ADP (see supplementary materials), arguing against there being a dramatically different conformation in the ATP vs. ADP binary complexes. The changes in $\tau_c$ for the various complexes indicates that protein closure occurs upon binding to M5P, with or without ADP present. This tightening/closure of the structure induced by M5P binding might be associated with domain movement around a His55/Arg93 hinge (Figs. 4 and 6). The chemical shift perturbation studies of PMK bound to M5P, with subsequent addition of Mg-ADP, demonstrate that the protein may open back up to bind ADP/ATP for catalysis.

**Hinge regions for lid and domain movement**

The previous discussion presented data in support of ligand-induced structural changes, which may be related to inter-domain (ATP/M5P) motion as well as to motion of the “Walker A” catalytic loop and the lid that covers it. Regarding movement of the lid, it is noteworthy that data on adenylate kinase indicates lid movement occurs on the same timescale as catalysis. This is consistent with the lid controlling access to the active site, as well as its packing up against the catalytic “Walker A” loop. In adenylate kinase, a key catalytic residue in the ATP binding site is Lys21, which twists into an unfavorable double gauche rotamer in order to interact with the ATP phosphate, just as Lys22 would in the human PMK homology model (Fig. 1) if its sidechain were adjusted to permit interaction with the ATP phosphate. This lysine may cycle into a catalytically useful orientation in the transition state to stabilize the increasing negative charge on the $\gamma$-phosphate, which would be created in the associative mechanism that is thought to be operative for nucleoside monophosphate kinases. Such a mechanism would permit selective stabilization of the transition state, and would require motion of the “Walker A” loop as the transition state is approached. So, lid motion is potentially relevant to catalysis. How does lid motion occur in PMK?

As noted earlier, a number of potential hinge residues had unusually small $S^2$ values (H55, R93, and D163), which indicates they are mobile on the psec-nsec timescale (note: often, fast timescale

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motion occurs along with msec timescale motion, which would be more relevant for a hinge region). Specifically, residues H55 and R93 are positioned (Fig. 4) such that they could operate as hinge residues that allow movement between the core ATP domain and the M5P domain, possibly for enabling an opening and closing motion upon substrate binding. Likewise, Asp163 is located in a potential hinge region for the “lid” loop region that is analogous to the “lid” that is present in the nucleoside monophosphate kinases. This lid caps the catalytic “Walker A” loop in the PMK model. The potential hinge residue, Asp163, also underwent dramatic chemical shift changes upon ligand binding (Fig. 4), as would be expected if there were motion around this hinge region induced by ligand binding. Motion around this hinge region would permit movement of the “Walker A” catalytic loop, which could be coupled to lid motion, as in adenylate kinase. The full extent to which catalytic loop and lid motions are coupled, and what role this motion might have in catalysis in human PMK, is not known, and will be the topic of future studies. But, extensive NMR studies on nucleoside monophosphate kinases, and especially adenylate kinase as a prototype, provide a rich literature that describes coupled motions between these loops, and their role in catalysis. These studies have also identified ligand-induced changes in protein correlation times (due to inter-domain movement; opening/closing), as we have now observed in human PMK. It will be interesting to see if human PMK follows the example of the adenylate kinase prototype, or whether it will provide unique and new insights into the role of such loop and domain motions in kinase-mediated phosphate transfer reactions. The studies reported herein provide a foundation for such future studies.

An N-Terminal Region of Unknown Function

The most noticeable feature of the order parameter profile (Fig. 6) is in the N-terminal region, which has no homology to adenylate kinase or to the T4 deoxynucleotidyl kinase template, upon which the homology model was based. But, there is a dramatic and sharp change in $S^2$ from a disordered state ($S^2 \approx 0.4$) to an ordered state ($S^2 > 0.8$), occurring immediately after Arg9, where the homology model starts. The fact that these nine residues are completely disordered in apo PMK, and all binary and ternary complexes, combined with the fact that organisms such as C. elegans lack this N-terminal segment.
entirely, suggests it is not an integral part of the PMK structure. Indeed, inspection of the homology model reveals no potential structural role for these nine N-terminal residues. This led us to speculate that these residues may be part of a signal or regulatory peptide, which might be cleaved off by an as-yet unidentified protease (note: this region shows no homology to membrane-binding peptides, and there are no data suggesting PMK is membrane bound). To begin exploring this hypothesis, and to identify which protease might cleave this N-terminal peptide, an exhaustive search of signal sequence databases was performed, searching only against the peptide sequence in question (i.e. Met1 to just past Arg9). While none of these searches yielded a strongly scoring motif hit, one did give a modest scoring prediction of cleavage immediately after Arg9 This site was identified using the ProP 1.0 server, using a neural network model trained using literature sequence data37. While the score for the cleavage site was relatively low (0.112; scores in the 0.5-1.0 range are considered high probability), it is the best scoring motif hit obtained thus far, and should be interpreted in the context of NMR data that indicate cleavage, if it occurs, would be expected at exactly this site (after Arg9). The enzyme predicted to cleave here is a furin-like protease, which belongs to a family of proteases broadly classified as “proprotein convertases”38. It should also be noted that other functions for this N-terminal region are also possible, such as binding to modular interacting domains of regulatory proteins (note: this sequence has no homology to existing SH3 motifs). This region also scored reasonably well for being a nuclear export signal peptide, using the NetNES 1.1 server. Clearly, the biological function of this N-terminal region remains to be elucidated, and is the topic of ongoing studies.

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Footnotes: Supporting Information Available. Spectra used to determine exchange rate limits (Suppl. Fig. 1), results of concentration-dependent T1 and T2 measurements to obtain \( \tau_c \) (Suppl. Fig. 2) and overlay of HSQC spectra of ATP and ADP binary complexes (Suppl. Fig. 3).
References


Supplementary Material

Click here to view.
http://www.ncbi.nlm.nih.gov/pmc/articles/PMC2649974/bin/NIHMS75666-supplement-Supp_Data.pdf
(152K, pdf)
**Figure S1.** Expansion of crosspeaks used to measure the kex range, along with chemical shifts used for the calculation.
Figure S2: PMK concentration effect on $\tau_c$, to ensure there was no protein aggregation effect on $\tau_c$. We did this by running 1D T1 and T2 relaxation experiments with the same nine time points used for the other relaxation experiments. The area under the peaks of the 1D spectra (1 representative is shown for each) was integrated to 1 and the intensity was measured at each time point using the VNMRJ program. The intensities of all nine experiments were plotted against time and fitted to an exponential curve to find $R_1$ and $R_2$. These $R_1$ and $R_2$ values were then used to calculate the $\tau_c$ value at each protein concentration: 500 $\mu$M, 250 $\mu$M, and 125 $\mu$M. The $\tau_c$'s are respectively $12.0 \pm 0.2$ ns, $13.5 \pm 0.4$ ns, and $12.5 \pm 0.5$ ns.
Figure S3. Overlay of $^1$H-$^{15}$N HSQC spectra for the PMK binary complexes with MgADP (black) or MgATP (red); spectra are from the highest concentration titration points (2 mM) from the titrations shown in Figures 3a and 3c.