Crystal Structures of Carbamate Kinase from *Giardia lamblia* Bound with Citric Acid and AMP-PNP

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**Abstract**

The parasite *Giardia lamblia* utilizes the L-arginine dihydrolase pathway to generate ATP from L-arginine. Carbamate kinase (CK) catalyzes the last step in this pathway, converting ADP and carbamoyl phosphate to ATP and ammonium carbamate. Because the L-arginine pathway is essential for *G. lamblia* survival and absent in high eukaryotes including humans, the enzyme is a potential target for drug development. We have determined two crystal structures of *G. lamblia* CK (glCK) with bound ligands. One structure, in complex with a nonhydrolyzable ATP analog, adenosine 5'-adenylyl-[β,γ]-imidodiphosphate (AMP-PNP), was determined at 2.6 Å resolution. The second structure, in complex with citric acid bound in the postulated carbamoyl phosphate binding site, was determined in two slightly different states at 2.1 and 2.4 Å resolution. These structures reveal conformational flexibility of an auxiliary domain (amino acid residues 123–170), which exhibits open or closed conformations or structural disorder, depending on the bound ligand. The structures also reveal a smaller conformational change in a region associated with the AMP-PNP adenine binding site. The protein residues involved in binding, together with a model of the transition state, suggest that catalysis follows an in-line, predominantly dissociative, phosphotransfer reaction mechanism, and that closure of the flexible auxiliary domain is required to protect the transition state from bulk solvent.

**Introduction**

Carbamate kinase (CK; EC 2.7.2.2, ATP:carbamate phosphotransferase) reversibly converts ADP and carbamoyl phosphate into ATP and ammonium carbamate (Fig. 1). The enzyme functions in the L-arginine dihydrolase pathway, operative primarily in some bacteria but also in the enteric protozoan parasite, *Giardia lamblia*. In contrast, CK is absent in high eukaryotes including Human. Since *Giardia* lacks the enzymes of the citric acid cycle and the oxidative phosphorylation pathway, the L-arginine dihydrolase and glycolytic pathways supply *G. lamblia* with ATP [1]. In addition, the L-arginine hydrolase pathway deprives host intestinal epithelial cells of arginine for nitric oxide biosynthesis, thereby dampening the innate immunity defenses [2,3]. Northern blotting revealed that the *G. lamblia* CK (glCK) gene was primarily expressed in *Giardia* trophozoites and that the mRNA level significantly decreased after encystation [4]. The glCK gene is also one of eight genes whose expressions are upregulated in neomycin selected *Giardia* cell lines [5], suggesting its importance under stress conditions. To validate glCK as a potential drug target, we used RNAi to knock down the glCK gene, which showed that the enzyme is essential for the survival of the organism under optimal laboratory growth conditions (*Giardia* possesses two nuclei and gene knockout is impractical) [6]. These data establish glCK as a potential target for drug development.

Previously, we have determined the crystal structure of glCK at 3.0 Å resolution [6], which revealed a large active site crevice within a core α/β domain. A region comprising 30 amino acid residues flanking the active site was largely disordered in that crystal structure suggesting a possible role for enzyme conformational transition during catalysis. Here, we report two enzyme-ligand crystal structures determined at higher resolutions than the earlier structure. We identify residues that are likely to play key roles during catalysis, and analyze a range of ligand-dependent conformational flexibility that protects the catalytic site from bulk solvent during the chemical reaction. One structure contains a bound non-hydrolyzable ATP analog, 5'-adenylyl-[β,γ]-imidodiphosphate (AMP-PNP), and the second structure (determined in two slightly different crystal forms) contains citric acid bound in the postulated carbamoyl phosphate site. These structures enable comparison with structures of ligand-bound CK from *Enterococcus faecalis* (efCK) [7] and the modeling of the transition state complex.

**Materials and Methods**

**Protein Preparation and Crystallization**

The glCK gene was cloned into the pDEST-HisMBP expression vector [8], and produced in *E. coli* strain BL21(DE3)Star as a TEV protease-cleavable, His-tagged, maltose binding protein (MBP) fusion product [6]. The cleaved and purified protein was concentrated to 30 mg/mL in solution containing 50 mM Tris-HCl (pH 8.0), 0.1 M NaCl, 5 mM MgCl₂, and 1 mM DTT. DTT
was required to prevent the protein from aggregation. The protein sample was stored in aliquots at −80°C.

The gCK was crystallized using the vapor-diffusion method in hanging drops. The protein solution was mixed with an equal volume of mother liquor containing 0.4 M ammonium citrate dibasic (pH 5.0), and 21% (w/v) polyethylene glycol (PEG) 3350. These gCK crystals contained citric acid bound in the active site (designated hereafter as gCK-citrateL). Crystals appeared within 1 to 3 days. For X-ray diffraction data acquisition, the crystals were transferred to mother liquor supplemented with 20% (w/v) glycerol and flash-cooled in liquid nitrogen.

A second gCK structure, also with bound citrate, was obtained after an unsuccessful attempt to soak a gCK-citrateL crystal with the drug Disulfiram, recently identified as a gCK inhibitor [9]. Flush cooling of the crystal led to shrinkage of the b unit cell dimension by >6%, and improvement in the diffraction resolution from 2.4 to 2.1 Å (Table 1). This structure is designated hereafter as gCK-citrateS. Finally, soaking a gCK-citrateL crystal with 50 mM AMP-PNP for 4 hours displaced the citric acid and yielded a structure with AMP-PNP bound in the active site (designated hereafter gCK-AMPPNP).

Data Collection and Refinement

X-ray diffraction data for both gCK-citrate crystals were acquired using an R-AXIS IV+ image-plate detector, mounted on a Rigaku rotating-anode MicroMax-007 X-ray generator (Rigaku MSC Inc). The diffraction data for the AMP-PNP soaked gCK crystal were collected at the GM/CA-CAT ID beam line at the Advanced Photon Source, Argonne National Laboratory, Illinois. The beam line was equipped with a MARmosaic 300 CCD detector (Marresearch GmbH).

Diffraction data were processed with the XDS program [10]. The previously determined gCK crystal structure was used as the model for the calculation of the initial difference Fourier electron density map [6]. Rigid body minimizations and refinements of the structures were carried out with the Phenix program [11]. Anisotropic scaling, bulk solvent correction, and isotropic temperature factors were used. TLS refinement did not improve the R values and therefore was not used. Water molecules were assigned using peaks in the Fo−Fc difference Fourier map with electron density >3σ as the acceptance criteria. Model building and structure modifications were performed with the Coot interactive graphics program [12].

Figures were generated with Raster3D linked to Molscript [13,14]. The coordinates and structure factors were deposited in the Protein Data Bank (entry codes 4JZ0 for gCK-citrateL, 4JZ8 for gCK-citrateS, and 4JZ7 for gCK-AMPPNP).

Results and Discussion

Conformational Flexibility of CK

CK assembles into homodimers, consistent with the oligomeric form detected in solution (Fig. 2). Two homodimers occupy the crystal asymmetric unit. Each subunit contains an eight-stranded β-sheet surrounded by three α-helices on one side and four α-helices on the other side of the β-sheet. This is a modified “Rossmann fold” organization with two additional β-strands at one edge of the β-sheet running antiparallel to the remaining six parallel β-strands [6]. The dimer interface is formed along the edge of the core Rossmann fold and also involves an additional α-helix that protrudes from this core. The new structures reveal an auxiliary domain, comprising a short β-strand followed by an α-helix and a β-hairpin (amino acids 123–170; Fig. 2B), which was mostly disordered in the previously reported gCK structure [6].

Depending on the bound ligand, the new crystal structures reveal this auxiliary domain in two different orientations with respect to the core domain. The large conformational changes, induced by soaking with ligands, are tolerated within the gCK crystals because the auxiliary domains are located close to crystal solvent channels. These conformational changes were accompanied by substantial changes to the crystal’s b cell dimension (Table 1). The b unit cell expanded by 5% upon soaking with AMP-PNP, and shrank by 8% upon the non-productive soaking with Disulfiram (Table 1). Thus, this pliable crystal form enables the sampling of multiple conformations while maintaining a reasonably good diffraction quality. The structural flexibility of the auxiliary domain is reflected in the higher crystallographic temperature factors (Bs) when compared to the entire structure. The average overall B values for gCK-citrateL and gCK-citrateS are 44 Å² and 31 Å², respectively (calculated for the four molecules in the asymmetric unit). By contrast, the average B values for the corresponding auxiliary domains are higher, at 63 Å² and 42 Å². For the gCK-AMPPNP structure, the overall B value is 24 Å², lower than that of two well-defined auxiliary domains at 63 Å². The remaining two auxiliary domains within the asymmetric unit are associated with weak electron density that cannot be interpreted.

A citrate ion occupies each subunit of the gCK homodimer in both gCK-citrateS and gCK-citrateL complexes (Figs. 2A & 3A). The main difference between the two structures is in the conformational state of one of the auxiliary domains. In the gCK-citrateS structure, all four auxiliary domains in the asymmetric unit are placed in proximity to the bound citrate, a state referred to as the closed conformation. In contrast, only three auxiliary domains in the crystal asymmetric unit of the gCK-citrateL structure adopt the closed conformation and one domain is placed more remotely from the citrate ligand, a state referred to as the open conformation.

In the gCK-AMPPNP structure, the citrate ions were displaced by the AMP-PNP because of partial overlap at the location of the γ-phosphate moiety (Fig. 3A&B). Of the two homodimers in the asymmetric unit, the auxiliary domains of only one homodimer adopt the open conformation while those in the second homodimer are structurally disordered.

The different conformational states of the auxiliary domain are accommodated within the same crystal form by changes in the spatial relationship between the two dimers in the asymmetric unit. Compared with the gCK-citrateL structure, one dimer in the gCK-AMPPNP crystal asymmetric unit is rotated by 3.0° and translated by 1.6 Å relative to the partner dimer. The corresponding values between gCK-citrateL and gCK-citrateS are smaller, 0.3° and 0.6 Å, consistent with a changed disposition of only a single auxiliary domain in the crystal asymmetric unit.

Flexibility of the auxiliary domain has been observed in two previous crystal structures of Enterococcus faecalis CK (gCK), one with bound ADP and Mg²⁺ and the second with bound SO₄²⁻ [7]. Each of these two structures was determined in a different crystal form. In particular, the gCK-SO₄²⁻ structure exhibited the closed conformation and was associated with extensive crystal contacts. In contrast, the three gCK complexes are accommodated within
the same crystal framework. Because crystal contacts do not drive the conformational changes, the gCK structures substantiate the proposed role of conformational flexibility during substrate binding and reaction pathway.

In addition to the gCK and efCK structures, three more CK or CK-like crystal structures have been deposited in the PDB. None of these exhibit the closed conformation. The structure of the enzyme from Pyrococcus furiosus contains ADP and exhibits the auxiliary domain in an open conformation [15]. The Mycoplasma penetrans CK structure contains two sulfate ions in the active site, one in the carbamoyl phosphate binding site and the other in the nucleotide’s 3-phosphate site. The auxiliary domain in this crystal structure is disordered [16]. Finally, a structure of CK from Aeropyrum pernix K1 has been determined by the Riken Structural Genomics team but has not been published (PDB entry code 2E9Y). This structure does not contain any bound ligand and the auxiliary domain exhibits the open conformation.

The relative disposition of the auxiliary domain in the open and closed conformation of gCK was analyzed using the program DynDom [17]. The conformational transition between the open and closed conformation within the gCK-citrateL protomers was modeled by a rigid body rotation of 44° (with a small translational component of 1.2 Å) around an effective hinge axis that runs perpendicular to a plane at the base of the domain (defined by residues 126–131 and 166–168). Similarly, the transition from the closed conformation in the gCK-citrate structures to the open conformation in the gCK-AMPPNP structure can be described by 36° and 31° rotations and 1.3 Å and 0.2 Å translations around effective hinge axes for the two structurally defined auxiliary domains.

A similar conformational transition was reported for the auxiliary domain in gCK with a 33° rotation about the effective hinge axis [7]. Comparison of the gCK and efCK auxiliary domains was facilitated by alignment of the secondary structure units of the core domains (161 of the 316 residues align with root-mean-squares deviation (RMSD) of 0.6–0.7 Å between 31{}-carbon atoms). For the gCK-AMPPNP and gCK-ADP structures, the orientations of the respective auxiliary domains differ on average.

### Table 1. X-ray data collection and structure refinement statistics.

| Crystal       | gCK-citrateL | gCK-citrateS | gCK-AMPPNP |
|---------------|--------------|--------------|------------|
| Data collection |              |              |            |
| Cell dimension (Å) | a = 69.9, b = 92.7, c = 101.8, β = 106.2 | a = 69.9, b = 86.4, c = 102.0, β = 106.5 | a = 70.6, b = 97.1, c = 102.1, β = 106.7 |
| Wavelength (Å) | 1.5418       | 1.5418       | 1.0332     |
| Resolution (Å) | 2.4          | 2.1          | 2.6        |
| No. of observed reflections | 162,457      | 202,519      | 130,375    |
| Completeness (%) | 99.4 (99.9)  | 99.6 (97.5)  | 99.2 (99.2) |
| No. of unique reflections | 48,667       | 67,363       | 40,467     |
| R<sub>merge</sub> | 0.097 (0.355) | 0.092 (0.322) | 0.093 (0.272) |
| <I<sub>cryst</sub>/ <I<sub>free</sub> | 7.2 (3.3) | 7.9 (2.9) | 8.1 (3.6) |
| Redundancy | 3.3 (3.4) | 3.0 (2.8) | 3.2 (3.1) |
| Refinement |              |              |            |
| No. of reflections used | 48,666       | 67,363       | 40,450     |
| No. of protein atoms | 9,448        | 9,313        | 8,960      |
| No. of ligand atoms | 52 (citrate) | 52 (citrate) | 124 (AMPPNP) |
| No. of water atoms | 275          | 700          | 475        |
| R<sub>cryst</sub> | 0.218 (0.302) | 0.220 (0.285) | 0.216 (0.275) |
| R<sub>free</sub> | 0.278 (0.367) | 0.288 (0.380) | 0.298 (0.344) |
| RMSd from ideal geometry |              |              |            |
| Bond length (Å) | 0.012 | 0.013 | 0.010 |
| Bond angle (°) | 1.4 | 1.4 | 1.3 |
| ΔB bonded (Å²) | 1.2 | 1.4 | 1.7 |
| Wilson B (Å²) | 31 | 24 | 18 |
| Average B factor (Å²) | Protein | 44 | 31 | 24 |
| | Ligand | 47 | 35 | 28 |
| | Water | 35 | 33 | 21 |
| Ramachandran plot (%) | | 86.0, 13.7, 0.3, 0.0 | 85.9, 13.9, 0.2, 0.0 |

*The values in parentheses are for the highest resolution shell, 2.40–2.49 Å for gCK-citrateL, 2.10–2.15 Å for gCK-citrateS, and 2.60–2.73 Å for gCK-AMPPNP.

R<sub>merge</sub> = Σ<sub>j=1</sub>Σ<sub>i=1</sub> | I<sub>j</sub> − | F<sub>j</sub> | / Σ<sub>j=1</sub> | F<sub>j</sub> |

R<sub>cryst</sub> = Σ<sub>j=1</sub>Σ<sub>i=1</sub> | F<sub>j</sub> − | F<sub>j</sub> | / Σ<sub>j=1</sub>Σ<sub>i=1</sub> | F<sub>j</sub> |

R<sub>free</sub> is computed using 2,009 randomly selected reflections omitted from the refinement for gCK-citrateL, 6,707 for gCK-citrateS, and 1,616 for gCK-AMPPNP.

Ramachandran plot categories are most favored, allowed, generously allowed, and disallowed.

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by a 16° rotation. For the closed conformations, the orientations of the respective auxiliary domains differ on average by an 11° rotation in the gl\textsubscript{CK}-citrate\textsubscript{S} and gl\textsubscript{CK}-phosphate\textsubscript{S} structures. The differences in the open conformations may simply reflect the multiple orientations that the auxiliary domain can adopt in the absence of substrate or substrate analog. However, the difference in the orientation of the auxiliary domain in the closed conformational state is probably due to the differences in the open conformations. The auxiliary domains of the second homodimer are disordered, which is not shown. (D) Amino acid sequence conservation of gl\textsubscript{CK} based on multiple alignment of the top 100 sequences identified using the BLAST protein sequence homology search [28]. Multiple sequence alignment was performed with ClustalW [29]. Invariant residues are colored in red. The N-terminus of two CKs in the redundant sequence database are truncated, thus only 98 sequences were used to define the first three invariant residues. Secondary structure units are boxed. β-strands and α-helices are show in red and blue colors, respectively, and in addition, the auxiliary domain is boxed in magenta color. doi:10.1371/journal.pone.0064004.g002

Figure 2. Ribbon depiction of dimeric gl\textsubscript{CK} structures. The flexible auxiliary domains and the core α/β domains are highlighted in different colors. Bound ligands are shown as stick models. (A) One homodimer of the gl\textsubscript{CK}-citrate\textsubscript{L} structure contains one auxiliary domain in an open conformation and the second in a closed conformation. (B) The second homodimer of the gl\textsubscript{CK}-citrate\textsubscript{L} structure and both homodimers of the gl\textsubscript{CK}-citrate\textsubscript{S} structure exhibit the auxiliary domains only in the closed conformation. (C) The homodimer in the gl\textsubscript{CK}-AMP-PNP crystal asymmetric unit that exhibits the open conformation. The auxiliary domains of the second homodimer are disordered, which is not shown. (D) Amino acid sequence conservation of gl\textsubscript{CK} based on multiple alignment of the top 100 sequences identified using the BLAST protein sequence homology search [28]. Multiple sequence alignment was performed with ClustalW [29]. Invariant residues are colored in red. The N-terminus of two CKs in the non-redundant sequence database are truncated, thus only 98 sequences were used to define the first three invariant residues. Secondary structure units are boxed. β-strands and α-helices are show in red and blue colors, respectively, and in addition, the auxiliary domain is boxed in magenta color. doi:10.1371/journal.pone.0064004.g002

Nucleotide Binding Site

AMP-PNP binds in a pocket flanked by amino acid residues 11–13, 253–250, and 267–278, (Fig. 3B). It exhibits the same interactions whether CK’s auxiliary domain adopts the open conformation or is disordered, as the nucleotide does not interact with the auxiliary domain. An AMP-PNP molecule binds to each of the four subunits within the crystal asymmetric unit. Consistent with the amino acid sequence homology analysis, the key protein-nucleotide interactions in the gl\textsubscript{CK}-AMP-PNP structure are conserved with those seen in the gl\textsubscript{CK}-ADP structure [7] with a few notable exceptions. The following interactions are common to both structures. The loop comprising amino acid residues 245–250 undergoes a conformational transition upon AMP-PNP binding such that the side chain of Tyr245 (Tyr238 in gl\textsubscript{CK}) stacks against the adenine ring and its hydroxyl group forms a hydrogen bond with the ribose O2’ hydroxyl group (see Fig. 3C for atom numbering and protein-ligand interactions). Although this tyrosine residue is sometimes replaced in other carbamoyl kinases by a phenylalanine or tryptophan, these replacements still permit aromatic-aromatic interactions.

Met275 (Met268 in gl\textsubscript{CK}) side chain stacks against the opposite face of the adenine moiety. Interestingly, other nucleotide binding proteins also exhibit methionine side chains that are placed to interact with ATP in a similar manner, for example carbamoyl phosphate synthetase [18] and acetylglutamate kinase [19].

Additionally, Val238 (Val231 in gl\textsubscript{CK}) forms a van der Waals contact with the ribose C3’. Thr236 O9 atom (Thr229 in gl\textsubscript{CK}) forms a hydrogen bond with one of the oxygen atoms of the β-phosphate. The loop encompassing residues 267–275 adjusts such that the backbone carbonyl oxygen of His269 (His262 in gl\textsubscript{CK}) forms a hydrogen bond with N6 of the adenine ring. Ser274 (Ser267 in gl\textsubscript{CK}) shifts away from the ribose site to form a van der Waals interaction with the ribose O4’.

Lys278 (Lys271 in gl\textsubscript{CK}) forms a salt bridge with the β-phosphate. Finally, the nitrogen atom of the AMP-PNP γ-aminophosphate interacts with the backbone amide nitrogen atom of Gly13 (Gly11 in gl\textsubscript{CK}).

Present in gl\textsubscript{CK}-AMP-PNP but absent in the gl\textsubscript{CK}-ADP structure is a hydrogen bond between the O3’ hydroxyl of the ribose and the carboxyl group of Asp237 (Gly230 is the equivalent residue in gl\textsubscript{CK}). In addition, an interaction that requires the presence of a γ-phosphate is missing in the gl\textsubscript{CK}-ADP structure - a salt bridge to Lys216 amino group (Lys209 in gl\textsubscript{CK}). The Lys209 in gl\textsubscript{CK} adopts an alternative conformation to interact with Asp210 carboxylate, a residue that also interacts with Lys271. This second interaction occurs also in gl\textsubscript{CK} between Asp217 and Lys278. Thus, the dispositions of Asp217 (Asp210 in gl\textsubscript{CK}) and the two lysine residues with respect to the nucleotide are pivotal for nucleotide binding and catalysis, consistent with the previous gl\textsubscript{CK}
electron density is associated with Mg^{2+}. The figure was generated using the program LIGPLOT [30].

Scheme and atom numbering of the ligands and their interactions with omitting the ligand from the model, are contoured at 2.5σ value of ADP by 5-fold.

The γ-phosphate in the gCK-AMPPNP structure is located in an equivalent position to one of the four water molecules that coordinates to Mg^{2+} in the gCK-ADP structure, suggesting that the γ-phosphate would also coordinate to Mg^{2+}. However, no electron density is associated with Mg^{2+} in the gCK-AMPPNP structure reported here. We attributed the lack of bound divalent ion to the relatively high citric acid concentration in the crystalization solution (0.4 M), a chelating compound that might have reduced the effective concentration of the metal ion. Attempts to obtain crystals with higher magnesium concentration were unsuccessful. We therefore propose that although catalytic activity requires the presence of a divalent cation [20], the metal ion is not required for nucleotide binding in the active site. Our interpretation is consistent with the gCK-ADP structure where Mg^{2+} coordinates solely to the ADP and water molecules. We note that the absence of a divalent ion from protein crystal structures containing AMP-PNP is not unusual. A survey of the Protein Data Bank showed that of >100 protein structures with bound AMPPNP, ~25% lack an accompanying Mg^{2+}.

Citrate Binding Site

Superposition of the gCK-citrateS and gCK-AMPPNP structures reveals that the citric acid binds adjacent to the AMP-PNP such that the C4–C5 acetyl group of citric acid overlaps with the AMP-PNP’s γ-phosphate group [see Fig. 3C for atom numbering and protein-ligand interactions]. In addition, superposition of the gCK-AMPPNP structures places the SO_{4}^{2−} close to the C6-carboxyl moiety of the citric acid. In both gCK-citrate and gCK-AMPPNP structures the auxiliary domains are in the closed conformations, and the closure partially protects the ligands from bulk solvent. As proposed earlier, the carbamoyl phosphate binds in this site [6,7]. Note that the citrate binds in the glycerol site identified in the earlier low-resolution crystal structure of gCK, however in the low-resolution structure, all auxiliary domains were disordered [6].

As with the nitrogen atom of the AMP-PNP β,γ-imidophosphate, the C5-carboxylate of citric acid interacts with the backbone nitrogen of Gly13 (Fig. 3A). An analogous interaction occurs in the gCK-AMPPNP structure (Gly11 in gCK), where SO_{4}^{2−} binding is also accompanied by a conformational change of residues 11–12 relative to the gCK-ADP structure, resulting in the backbone amides of both residues Gly11 and Asn12 interacting with the sulfate ion. Whereas the sulfate ion binds 4.3 Å away from the ADP’s β-phosphate, the citrate’s C4–C5 acetyl group is placed within a covalent bond distance from the β-phosphate of the AMP-PNP, and there is no conformational difference between the backbone conformation of Gly13 and Asn14 in the gCK-citrate and gCK-AMPPNP structures.

Other CK-citrate interactions that may mimic CK-carbamoyl phosphate interactions include the interaction of the C6 carboxyl group with the backbone amides of Asn55 and Gly56, located at the N-terminus of an α-helix (Fig. 3A&C). A similar “oxyanion hole” is also conserved in the gCK, where it serves as a second anchor for the sulfate anion. The C6 carboxyl group also interacts with Lys131’s amino group, located on the auxiliary domain. Again, a similar interaction between the sulfate and Lys128 is present in the gCK-ADP structure [7]. Finally, the C1 carboxyl group is bridged to Arg163 on the auxiliary domain via a water molecule. In the gCK structure, a second bound sulfate ion forms a salt bridge with the equivalent arginine residue (Arg158 in gCK).

Phosphotransfer Transition State Model

Superposition of the AMP-PNP and citrate bound structures reveals an overlap between the positions of the two ligands. The γ-phosphate of AMP-PNP and the C5 carboxylate group of the citrate are placed such that the β-phosphorous atom of AMP-PNP and one of the carboxylate oxygen atoms of the citrate are separated by 1.7 Å. These two structures provide further support to the previously proposed catalytic mechanism, which involves an in-line nucleophilic substitution at phosphorous [6,7]. Moreover, the active site lacks an appropriately positioned histidine or aspartic acid residues that may serve as a temporary phosphoryl group acceptor in a two-step phosphotransfer reaction involving a phospho-enzyme intermediate (see reviews of phosphotransfer reaction mechanisms [21,22,23]).

A model of the transition state formed during the phosphoryl group transfer is shown in Fig. 4A. The protein model in this transition state is a composite of two crystal structures; the closed conformation of the auxiliary domain as seen in the gCK-citrate structure and the C-terminal domain’s loop containing residues 245–250 as seen the gCK-AMPPNP structure. The Mg^{2+} was positioned by analogy to the gCK-ADP structure. The location of the AMP-PNP defines the ADP binding site. The ADP-protein...
interactions correspond to those described above for the AMP-PNP, which include interactions with both the protein backbone and the side chains. The β-phosphate is anchored not only by the interactions with the protein (Gly13 backbone amide, Lys278 amino group, and Thr236 hydroxyl group) but also by coordination to Mg²⁺, as observed in the gCK-ADP structure.

The location of the carbamoyl moiety overlaps with the citrate and is delineated by favorable electrostatic interactions and also by the avoidance of van der Waals clashes. Specifically, the carbamoyl's nitrogen atom binds adjacent to the side chain of Val213 (Fig. 4A), an invariant residue in all CK sequences (Fig. 2D). Together with the carbamoyl orientation relative to ADP, the restricted position of the carbamoyl's nitrogen atom defines the location of the carbamoyl phosphate site. Whereas the proximity of the nitrogen atom to Val213 still allows a hydrogen bond interaction of the carbamoyl's nitrogen atom with the side chain of the invariant Asn55, it prevents the nitrogen atom from forming a strong electrostatic interaction with the backbone carbonyl oxygen of Ile214 (the distance is 3.6 Å). In contrast, the carboxylate group of the carbamoyl moiety forms four favorable interactions. The free oxygen interacts with the amino group of the invariant Lys131, a residue located on the auxiliary domain, and with the backbone amide of Lys216. The phosphoryl group donor oxygen atom is located in an oxyanion hole formed by the backbone amide groups of Asn55 and Gly56 observed in all available CK structures.

The transition state's planar PO₃ group is located half-way between the donor and acceptor oxygen atoms, such that the O(donor)-P-O(acceptor) atoms form the apical axis and the PO₃ oxygen atoms lie in the equatorial plane of a trigonal bi-pyramid centered on the phosphorous. The PO₃ group interacts with the amino group of the invariant Lys216, as does the AMP-PNP γ-phosphoryl group in the experimental gCK-AMPPNP structure. However, unlike the AMP-PNP's γ-phosphate, the PO₃ in the transition state model is located 3.7 Å from Mg²⁺, i.e. too remote for direct coordination to Mg²⁺. The reaction direction towards ATP production may be governed in part by the preference to establish a γ-phosphate-Mg²⁺ coordination.

The proposed transition state model places the phosphoryl donor and acceptor oxygen atoms 5.8 Å apart, with the phosphorous at an equal distance (2.9 Å) to the donor and acceptor. The distance agrees closely with the quoted donor-acceptor oxygen atoms' distance of 6.0 Å in the model reported by Rubio's and colleagues [7]. A 2.9 Å long apical P-O distance corresponds to a predominantly dissociative mechanism. Using Pauling’s rule [24], the bond order is 0.02. The dissociative phosphotransfer mechanism has been implicated in previous crystal structures of creatine kinase, glycocoyamine kinase, and arginine kinase, where each enzyme formed a complex with a transition state analog [25,26,27]. In these structures, the transition state analogs comprised ADP, nitrate (a metaphosphate analog) and the respective substrates. The distances between the nitrate’s nitrogen atom to either donor and acceptor oxygen atoms are ~3 Å in all these cases, consistent with the distances in the carbamoyl kinase transition state model.

In contrast to the three phosphagen kinase structures listed above, where the metaphosphate analog, nitrate, is fully buried and protected from bulk solvent, the closed conformations in both gCK and eCK structures do not fully sequester the transition state, allowing bulk solvent to access the metaphosphate. Indeed, only one of the metaphosphate’s oxygen atoms interacts with a protein residue (Fig. 4A). An exposed transition state would be a target for hydrolysis. It is tempting to speculate that the auxiliary domain may close further so that the transition state is fully protected to promote the phosphotransfer reaction and prevent a non-productive hydrolysis. Two invariant residues, Asp157 and Arg163, are located on the surface of the auxiliary domain exposed to solvent and they could potentially facilitate further closure (Fig. 4B). In a true transition state, Arg163 could interact with the transferred phosphoryl group and Asp157 could perhaps coordinate to the Mg²⁺.

The superposition of the gCK and eCK structures suggests that the transition state would be further protected from bulk solvent if the gCK residues Gly13-Asn14 adjust to generate an oxyanion hole analogous to the oxyanion hole of Gly12-Asn13 in the gCK-SO₄ structure (invariant residues in all CK sequences). This generates electrostatic interactions between the two protein backbone amides and the remaining free metaphosphate oxygen atom, and also forms hydrogen bonds between Asn14 side chain

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Figure 4. A model of the transition state. (A) Stereoscopic representation showing the key protein and Mg²⁺ interactions with the transition state components. The apical axis between the donor and acceptor oxygen atoms is shown as dashed lines in magenta and the distances from the transferred phosphorous to the donor and acceptor oxygen atoms are 2.9 Å. Atoms are colored as follows: Carbon – gray, Oxygen – red, Nitrogen – blue, Magnesium – magenta. The carbamate is labeled CM (B) The relationship between the transition state and Arg163 and Asp157 on the auxiliary domain (colored in yellow), as defined by the closed conformation of gCK. The side chains of Arg163 and Asp157 project towards the transition state. However, the distances (listed in Å) are too far, allowing bulk solvent access into the active site. Protection of the transition state and prevention of phosphate hydrolysis requires a further closure of the auxiliary domain. doi:10.1371/journal.pone.0064004.g004
and both, the metaphosphate and the β-phosphate of ADP. Together, the further closure of the auxiliary domain and the transition of Gly13-Asn14 to form an oxyanion would fully sequester the transferred phosphoryl. Whether such conformational transitions could also induce contraction of the transition state towards a more associative phosphotransfer reaction mechanism remains to be determined.

Conclusion

The crystal structures of gCK bound with citric acid and with AMP-PNP reveal a dynamic molecule that exhibits substantial conformational flexibility within the same crystal form. The lack of bound Mg²⁺ in the gCK-AMP-PNP structure indicates that the divergent cation is not necessary for nucleotide binding, although it is required for catalysis. The transition state model, proposed based on the two crystal structures, is consistent with an in-line phosphotransfer reaction mechanism that is predominantly dissociative. Protection of the transition state from bulk solvent suggests a further closure of the auxiliary domain. Structures of additional complex with a transition state analog, kinetic isotope effect, analysis of stereochemical outcome, and site directed mutagenesis studies of residues postulated to facilitate catalysis and active site closure would help elucidate the catalytic mechanism. With respect to novel anti-giardiasis therapeutic approaches, the new structures provide a solid structural basis for drug development and show that new inhibitors may be designed with multiple conformational states of the enzyme in mind.

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Author Contributions

Conceived and designed the experiments: OH. Performed the experiments: KL. Analyzed the data: OH. Contributed reagents/materials/analysis tools: LK. Wrote the paper: OH. KL. AG.

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