Evolutionary plasticity in the allosteric regulator binding site of pyruvate kinase isoform PykA from *Pseudomonas aeruginosa*.

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Running title: Structure and regulation of PykA

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Abstract

Unlike many other well-characterized bacteria, the opportunistic human pathogen Pseudomonas aeruginosa relies exclusively on the Entner-Doudoroff pathway (EDP) for glycolysis. Pyruvate kinase (PK) is the main "pacemaker" of the EDP, and its activity is also relevant for P. aeruginosa virulence. Two distinct isozymes of bacterial PK have been recognized, PykA and PykF. Here, using growth and expression analyses of relevant PK mutants, we show that PykA is the dominant isoform in P. aeruginosa. Enzyme kinetics assays revealed that PykA displays potent K-type allosteric activation by glucose 6-phosphate and by intermediates from the pentose phosphate pathway. Unexpectedly, the Xray structure of PykA at 2.4 Å resolution revealed that glucose 6-phosphate binds in a pocket that is distinct from the binding site reported for this metabolite

in the PK from Mycobacterium tuberculosis (the only other available bacterial PK structure containing bound glucose 6-phosphate). We propose a mechanism by which glucose 6phosphate binding at the allosteric site communicates with the PykA active site. Taken together, our findings indicate remarkable evolutionary plasticity in the mechanism(s) by which PK senses and responds to allosteric signals.

Introduction

Pyruvate kinase (EC.2.7.1.40) is a key enzyme in glycolysis, and a major site of pathway regulation. The enzyme catalyses the conversion of phospho*enol*pyruvate and ADP into pyruvate and ATP (1). Given its central position in metabolism, pyruvate kinase (PK) activity can have a major impact on the carbon economy of the cell (2, 3). More recently, it has been shown that PK activity can also influence pathogencity (4). Bacterial PK is encoded by two isozymes, *pykA* and *pykF* (5–7). Most organisms encode either *pykA* or *pykF*, although a few species also encode both isozymes. Among the organisms encoding both pykA and pykFthat have been characterised to date, PykF is usually the dominant isoform. The PykA and PykF isozymes have been especially well-studied in the Enterobacteriaceae, and are distinguished by being differentially regulated and cladistically-distinct (at a sequence level). For example. PykF isozymes are typically activated by fructose 1,6-bisphosphate (F1,6P) (8), whereas PykA is activated by adenosine 5' monophosphate (AMP) (9). Several crystal structures are available for PykF isozymes, although no structures have been solved for PykA. Recently, the structure of Mycobacterium tuberculosis PK bound to AMP and glucose 6-phosphate was solved (the first structure to be solved for a bacterial PK with bound allosteric regulators). The regulators were found to bind at distinct sites in the allosteric cleft and were proposed to communicate with the active site through a series of domain re-orientations denoted the "rockshape-lock" mechanism.

Pseudomonas aeruginosa is an opportunistic pathogen which commonly causes pulmonary (10), urinary tract (11), and soft tissue-associated infections in hospitalized patients. The pathogen is intrinsically resistant to most clinically-used antibiotics (12) and was recently designated by the World Health Organisation (WHO) as a "critical priority pathogen". Unlike the Enterobacteriaceae, which mainly employ the Embden-Meyerhof-Parnas (EMP) pathway for alucose catabolism. Ρ aeruginosa lacks phosphofructokinase and utilizes the Entner-Doudoroff pathway (EDP) instead (13, 14). As a consequence, PK is likely to be the major site of glycolytic flux regulation in this organism. P. aeruginosa encodes two uncharacterized PK isozymes, annotated as pykA (PA4329) and pykF (PA1498). Here, we show that in contrast with most organisms encoding both PK isozymes, in *P. aeruginosa*, PykA plays the dominant physiological role. We also present the first structure of a PykA isozyme. Surprisingly, this structure revealed that glucose 6-phosphate binds in a site distinct to that observed for the *M. tuberculosis* enzyme.

<u>Results</u>

PykA is the dominant pyruvate kinase isozyme in *P. aeruginosa*

P. aeruginosa encodes two PK isozymes; PykA (PA4329, 58% amino acid identity with *E. coli* PykA) and PykF (PA1498, 37% amino acid identity with E. coli PykF). An alignment of selected relevant PykA and PykF sequences is shown in Figure S1. Phylogenetic classification of *P. aeruginosa* PykA and PykF (Figure S2) indicates that the isozymes belong to two distinct sub-classes of PK. To establish which isozyme is dominant in *P. aeruginosa*, we measured the PK activity in cell extracts of mutants defective in pykA, pykF, or pykA and pykF together (*pykA pykF*). Following growth on medium minimal containing glucose, glycerol, or acetate as a sole carbon source, pyruvate kinase activity was >90% lower in the *pykA* mutant and in the *pykApykF* mutant than it was in the wild-type control (Figure 1). However, pyruvate kinase activity in the pykFmutant remained high on these substrates. These data indicate that PykA contributes most to PK activity in P. aeruginosa. Consistent with this, western analyses using antibodies raised against purified Ρ. aeruginosa PykA or PykF confirmed that the latter is essentially undetectable during growth on the carbon sources tested unless over-expressed in trans from a plasmid (Figure 1). Note that the antibodies were preabsorbed against a pykA mutant (for the anti-PykA antibodies) or a *pykF* mutant (for the PykF antibodies). The absence of PykF expression in a *pykA* mutant led to weak cross-reactivity of the anti-PykA antibodies against shared PykA/PykF epitopes in PykF, although this was only apparent when PykF was over-expressed. We conclude that pvkA is expressed at higher levels in these conditions than *pykF*.

To explore the phenotypic consequences of *pykA* and *pykF* mutation, we assessed the growth characteristics of each mutant on different carbon sources (Figure 1). The *pykA* mutant and the *pykA pykF* mutant grew slowly compared with the wild-type and the *pykF* mutant in minimal media containing glucose or glycerol as a sole carbon source. That there was any observable growth of these mutants at all on glucose is

presumably due to the pyruvate produced at 2-keto-3-deoxy-6-phosphogluconate the (KDPG) aldolase-catalyzed step of the EDP. Similarly, the slow, but non-negligible growth in glycerol (which feeds in to the EDP after the KDPG aldolase-catalyzed step) can be rationalized by invoking operation of the recently-defined EDEMP cycle (15). The EDEMP cycle allows triose phosphates such as those derived from glycerol to be recycled through the KDPG aldolase step, thereby yielding pyruvate. Growth on glucose or glycerol was restored in the *pykA* mutant and the *pykA pykF* mutant by expression of either pykA or pykF in trans, suggesting that both isozymes have the potential to complement the *pykA* mutant growth defect. These data also indicate that the low PK activity in the *pykA* mutant is likely due to low expression of PykF, rather than low intrinsic activity of the isozyme. Consistent with this, pyruvate activity increased kinase and PykF became detectable expression during growth on glucose when pykF was expressed in trans in the pykF mutant (Figure 1). [PykF is predicted to be part of a bicistronic operon alongside a putative glycerate kinase, PA1499. The ppykF+ complementation plasmid encoded both PA1499 and pykF, see Materials and Methods.] In contrast, during growth on acetate, the pykA, pykF, and pykA pykF mutants all grew at essentially the same rate as the wild-type. This suggests that PykA plays an important role in catabolising substrates that feed into the EDP such as glucose and glycerol, but contributes less to growth on acetate (which does not require flux through the PK catalysed reaction).

Kinetic properties and regulation of PykA

Purified PykA was an approximately 200 kDa tetramer in solution based on analytical ultracentrifugation data (Figure S5, S6). At saturating phospho*enol*pyruvate (PEP) concentration, the enzyme displayed Michaelis-Menten kinetics with respect to [ADP], with a K_M value of 0.07mM (Figure 2, Table S1). In contrast, at saturating ADP concentration, the PEP-dependency was sigmoidal, with an $S_{0.5}$ value of 0.67mM(Figure 2, Table S1) and a Hill coefficient of 2.14, indicative of positive cooperativity.

All PK enzymes require divalent cations for activity (mostly Mg²⁺), and some have also been reported to require K⁺ in order to achieve maximum catalysis (16). The presence of these ions is important to facilitate binding of the substrate, transfer of the phosphoryl group from PEP to ADP, and possibly also acquisition of an active enzyme conformation (17, 18). We found that the activity of *P. aeruginosa* PvkA was independent of K⁺ concentration (Figure 2). Indeed, addition of monovalent cations (K⁺, NH₄⁺ or Na⁺) even somewhat decreased the overall catalytic activity of the enzyme (Figure S7, Table S1). The K⁺-mediated activation of pyruvate kinase activity is known to be dependent on the nature of the residue immediately preceding strand B_{β1} in the structure (19); in K⁺-dependent enzymes, this residue is a glutamate, whereas in K⁺independent enzymes, it is often a lysine. In *P. aeruginosa* PykA, the corresponding residue is a lysine (Figure S1).

Pyruvate kinase is a major site for the regulation of glycolytic flux in many species, and most of the known regulators in other species are associated with the EMP. However, *P. aeruginosa* relies solely on the EDP for glycolysis. To investigate the regulation of *P. aeruginosa* PykA further, we screened a wide range of metabolic intermediates - guided by a knowledge of the pathways feeding in and out of the EDP - for their ability to activate or inhibit PykA in vitro. Putative regulatory molecules were screened at low PEP concentration (0.3 mM) to identify activating ligands, and high PEP concentration (3 mM) to identify inhibitory ligands (Figures 2 and S8, respectively). More detailed kinetic analyses $(S_{0.5}, k_{cat})$ were carried out on the "hits" identified this way (Figure 2 and S9, Table S1). No inhibitors were identified. However, PykA was strongly activated by glucose 6phosphate (G6P) from the EDP and by intermediates from the reductive PPP (fructose 6-phosphate (F6P), glyceraldehyde 3-phosphate (G3P), ribose 5-phosphate (R5P), ribulose 5-phosphate (RL5P) and xylulose 5-phosphate (X5P)). The main effect of these regulators was to decrease suggestive of K-type allosteric $S_{0.5}$ regulation, and to change the substrate dependency kinetics on PEP from a sigmoidal to a hyperbolic (i.e., Michaelis-Menten) profile. Although these regulators had a large impact on the PEP-dependency of the reaction, they had little or no effect on its ADP-dependency. Interestingly, AMP and fructose 1,6-bisphosphate (two wellestablished regulators of PK in many other species) had almost no effect on PykA activity.

Structure of PykA

Currently, there is no structure available for any PykA isozyme. We were therefore fortunate to obtain crystals of PykA containing a substrate analogue (malonate) and Mg²⁺ bound in the active site, and a bound allosteric regulator (G6P). The crystals diffracted x-rays to 2.4 Å resolution. The asymmetric unit of PykA contained 12 polypeptides assembled into three complete homotetramers. This quaternary structure is consistent with analytical ultracentrifugation analyses (AUC) data indicating that PykA is tetramer in solution (Figure S6). а Refinement statistics are shown in Table 2.

Each protomer of PykA had the typical tripartite domain organization associated with PKs, with the A and C domains of one protomer interacting with the A and C domains (respectively) from the adjacent protomers (Figure 3A,B). The A domain is comprised of 8 α/β structures assembled into a TIM barrel motif. Helices A α 6 and A α 8 are further subdivided into shorter segments forming Aa6' and Aa8'. The Aa6'helix contains part of a highly conserved segment (M₂₄₅VARGDLGVE₂₅₄) which plays a key role in catalysis (Figure 3C). The active site is located in the cleft between domains A and B (Figure 3B). Relative to the A domain, the spatial disposition of the β -rich B domain varied somewhat between the different protomers, indicating that the domain is configurationally flexible as a unit. The $\alpha\beta$ rich C domain contains the bound allosteric regulator, G6P.

The PykA active site contains a bound substrate analogue, malonate (present in the crystallization buffer) and Mg²⁺ (Figure 3C). The malonate is bound by hydrogen bonds from both the polypeptide backbone (residues Gly₂₄₉ (2.8 Å) and Asp₂₅₀ (3.1 Å)) and the sidechains of Lys₂₂₁ (2.9 Å), Thr₂₈₂

(2.6 Å) and Arg_{34} (3.3 Å). In addition, the active site Mg²⁺ also contributes to malonate binding through electrostatic interactions. The Mg²⁺ is octahedrally coordinated through interactions with the sidechains of Glu₂₂₃ and Asp₂₅₀, and also with two water molecules. These active site residues are absolutely conserved in all bacterial PKs (Figure S1), as is their spatial disposition, even when comparing malonate-bound PykA with pyruvate-bound PK from rabbit muscle (PDB 1F3W) (Figure 3D). In contrast, superposition of the PykA active site with that of E. coli apo-PykF (PDB 1PKY) reveals that almost all the residues just discussed undergo significant spatial re-orientation in the presence of the substrate, indicative of induced fit (Figure 3E).

An alignment of PykA sequences reveals that all PykA orthologues contain 3 residues not present in any of the PykF orthologues, corresponding to Ala₂₂₉, Asp₂₃₀ and Asp₂₃₁ in *P. aeruginosa* PykA (Figure S1). The presence of these three additional residues lengthens the loop connecting A α 5-A β 5 resulting in the displacement of helix A α 5 away from the core of the enzyme (Figure S10). The functional significance of this is not yet clear, although it is worth noting that the same loop of structure contributes two key residues to the active site; Glu₂₂₃ and Lys₂₂₁ (Figure S10).

Kinetic analysis indicated that G6P is an activator of PykA, increasing the apparent catalytic efficiency ($k_{cat}/S_{0.5}$) of the enzyme around 3-fold (Table S1 and Figure 2,S9). Inspection of the PykA structure revealed a molecule of G6P buried in each C-domain, far from the active site, indicating that G6P is an allosteric regulator (Figure 3B). The structure of only one other bacterial PK with bound regulator has been solved, and that is the PykF family enzyme from *M. tuberculosis* (Mtb). A comparison of the allosteric binding site(s) observed in the Mtb enzyme reveals AMP bound in the same location as the G6P in *P. aeruginosa* PykA. The Mtb enzyme does bind G6P, but the ligand is located in an entirely different position in the allosteric cleft (PDB 5WSB) (Figure 4A).

The G6P binding pocket in PykA is comprised of two loops; a "phosphate loop" and a "ring loop". The phosphate loop ($C\beta$ 1-

Cα2, residues 383-388) and partly the Cα2 helix (Thr₃₈₉) binds the phosphate group of the ligand, whereas the ring loop (Cβ4-Cβ5, residues 460-471) anchors the hexose ring moiety. Superposition of the G6P binding site in PykA with the PykF apo-enzyme (1PKY) from *E. coli* reveals that in the presence of G6P, the ring loop of PykA is pulled tightly over the allosteric pocket (Figure 4B), whereas the phosphate loop shares a similar disposition in both structures. The movement of the ring loop in the presence of G6P causes a shift and partial unwinding of the Cα4 helix in PykA.

bound G6P The interacts with the surrounding residues through a network of hydrogen bonds involving both the phosphate group and the sugar ring (Figure hvdroxvls 4C). In addition. occupation of the allosteric site also appears to promote secondary interactions between the phosphate loop and the ring loop of PykA; Ser₃₈₆ and Thr₃₈₉ on the phosphate loop form hydrogen bonds with Gln₄₆₇ and Thr₄₇₀ on the ring loop, respectively. Interestingly, Lys₄₆₀ and Tyr₄₆₄ on the ring loop also interact, tightening up closure of the binding pocket (Figure 4D). This may be unique to the PK sub-class represented by P. aeruginosa PykA, because its ring loop is less well conserved than the ring loop in other enzymes PykA (Figure S1). Furthermore, half of the residues that mediate the interaction between the phosphate and the ring loops are unique to *P. aeruginosa* PykA (Lys₄₆₀, Tyr₄₆₄, Gln₄₆₇).

Comparison of the G6P-bound PykA structure with the E. coli apo-PykF structure revealed differences in (1 PKY)the disposition of residues at the protomer interfaces. The PvkA tetramer contains four inter-protomer interfaces, two of which are between adjacent A-domains (A-A interface) and two between adjacent C-domains (C-C interface) (Figure 3A). The A-A interface (approximately 1450 $Å^2$) is formed through pairing of Aa6, Aa6', Aa7, Aa8, loop Aa6-A α 6', loop A α 7-A β 7 and loop A α 8-C α 1 from one protomer with the same secondary (albeit, structural units with inverted symmetry) on the second protomer (Figure S11). By contrast, the apo-structure of PykF does not include Aa6' or loop Aa8-Ca1 as

parts of the interface and instead, there is an $A\alpha$ 7- $A\alpha$ 7 interaction which is absent in PykA (Figure 5A). The A-A interface in PykA is very close to the active site, so structural changes at the interface affect the disposition of active site residues including Arg₂₄₈, Gly₂₄₉, Glu₂₈₃, Asp₂₅₀ and Thr₂₈₂ (Figure S12). Indeed, the mainchain amide nitrogen of Gly₂₄₉ is directly involved in binding the substrate analogue, whereas the amide carbonyl moiety of this residue forms a hydrogen bond with the side chain of Arg₂₉₆ on helix Aa7 on the adjacent protomer (Figure 5C). The A-A interface is stabilized by just two salt bridges (Figure S13), which contrasts with the situation seen in E. coli apo-PykF, where most of the interactions at the A-A involve salt bridges.

Comparison of G6P-bound PvkA with apo-PykF (1PKY) also reveals major differences at the C-C interface (Figure 5B). The C-C interface in PykF is comprised of C α 4, C β 5, and the ring loop between C β 4 and C β 5. By contrast, the C-C interface in PykA is comprised of C β 5, the ring loop, C α 1 and the long, flexible loop between Aa8-Ca1. Essentially, the bound G6P in PykA displaces the ring loop and C α 4, resulting in an outward movement of $C\alpha 1$. This drags the Aa8-Ca1 loop region closer towards the C-C interface (Figure 5EF). This movement of the Aa8-Ca1 loop could, in turn, communicate the conformational change from the G6Pbinding site to the A-A interface and the active site, providing a plausible mechanism by which the allosteric regulator elicits its effect.

Discussion

Consistent with its role as the major PK isozyme in P. aeruginosa, a pykA mutant was defective in growth on glucose and glycerol. These data raise the question of why *P. aeruginosa* encodes two distinct PK isozymes? This is intriguing, especially given that the gene encoding one of the isozymes (pykF)appeared to show negligible expression under any of the growth conditions tested. One likely solution is that *pvkF* appears to be part of a larger cluster of genes (PA1498-PA1502) that may be involved in the catabolism of compounds such as ethylene glycol (20). Therefore, artificially-expressed although pykF is

capable of rescuing the growth defect associated with the *pykA* mutant (Figure 1), its expression is likely to be induced only under very select conditions.

P. aeruginosa lacks phosphofructokinase (encoded by *pfk*) and is therefore incapable of carrying out the EMP pathway of glycolysis (13). Instead, and in common with around 13% of species for which a genome sequence is available, it is exclusively reliant on the EDP for glucose oxidation (21). In this pathway, pyruvate kinase is the major regulatory enzyme. PykA activity was activated primarily by G6P, F6P, G3P and by intermediates of the reductive PPP. In many organisms, the upper arm of the reductive PPP is "fed" by the conversion of 6phosphogluconate into ribulose 5phosphate; a reaction catalysed by 6phosphogluconate dehydrogenase (*gnd*) (22-24). However, and like many other bacteria that exclusively utilise the EDP, P. aeruginosa does not encode a homologue of gnd (13, 25). As a consequence, carbon can only flow into the PPP from G3P or from gluconeogenesis-derived F6P (Figure 6) (26). Not surprisingly, given the lack of redundancy in input pathways, many of the PPP-encoding genes are essential in P. aeruginosa (26). It is therefore logical that P. aeruginosa has evolved to coordinate glucose oxidation with the supply of biosynthetic precursors by titrating both gluconeogenic and PPP intermediates. These two anabolic pathways are tightly inter-twined; as shown in Figure 6, EDPderived G3P, and F6P derived from gluconeogenesis also play a central role in the cyclical series of sugar phosphate interconversions which comprise the reductive PPP. High levels of these, and other key PPP intermediates (R5P, X5P and physiological RL5P) presumably are indicators that biosynthetic precursors are abundant (27), and that carbon flux can be redirected towards energy production via PykA activation. Similarly, high levels of G6P indicate sufficiency of either glucose supply or of gluconeogenic flux, so it makes good metabolic sense that this too should feed forward to stimulate PykA, and thence, carbon catabolism.

The only other bacterial PK structure containing bound regulatory ligands is the PK from *M. tuberculosis* (28). That work revealed that AMP binds in the heart of the C domain (Figure 4A), and that G6P binds at the interface between the A and C domains. However. Ρ. aeruainosa PvkA was insensitive to AMP, and G6P was found to be bound to the same site as that occupied by AMP in the Mtb structure. This indicates that although the allosteric binding pocket is conserved as a cavity in PykA, there is considerable evolutionary plasticity in regulator binding. That different enzymes from the same family (albeit, from different species) have evolved to bind the same allosteric regulator (G6P) in different sites is remarkable, especially from the perspective of allosteric transitions. Moreover, and given the radically-different structures of ring-sugar phosphates such as G6P and F6P compared with "configurationally locked" straight-chain forms such as RL5P and X5P, this suggests that the latter may well bind PykA at other, vet-to-be-characterised sites.

Unfortunately, we were unable to obtain crystals of apo-PykA that diffracted bevond ca. 4Å. We therefore based our structural comparisons on the PykF apo-enzyme from E. coli, which does not contain bound substrate or regulatory ligands. In the PykF structure, the allosteric pocket is open, and is uncapped by the ring loop (29). This unbound configuration allows the Ca4 helices to form prominent interactions across the C-C interface. However, when the allosteric site becomes occupied by G6P, the ring loop is pulled down towards the ligand. This partially displaces $C\alpha 4$ and firmly closes the allosteric pocket through interactions with both the ligand and the phosphate loop. The new configuration of the allosteric site seems to loosen up the interactions from $C\alpha 4$, concomitantly recruiting other structures to the interface, most notably $C\alpha 1$ and the $A\alpha 8$ -Cal loop (Figure 5). Thus, in PykA, the Aa8contacts two Ca1 loop protomers simultaneously via the A-A interface and the C-C interface. Given the proximity of the A-A interface with the active site, this suggests a likely mechanism by which G6P binding may influence catalysis. Figure 7 illustrates this proposed mechanism. G6P-binding to the allosteric pocket leads to movement of the part of the active site). This proposed mechanism contrasts with the rigid-body "rock-shape-lock" mechanism proposed for Mtb PK (28).

Materials and Methods

For primers and bacterial strains see Table S1.

Construction of mutants

The *pykA* and *pykF* mutants used were from the Washington Mutant Library (identities PW8308 (pykA-B04::ISlacZ/hah, which carries a Tn insertion at position 903/1452) and PW3705 (pykF-C02::ISphoA/hah, which carries a Tn insertion at position 213/1434), respectively). The insertion site of the transposon in each mutant was confirmed by PCR using the primers recommended by the library curators. To generate the pykA pykF double mutant, we first had to remove the Tet^R marker from the pykA mutant. This was done by introducing plasmid pFLP2-cre into the pykA mutant by electroporation (to enable site-specific cre-mediated excision of the transposon) followed by plasmid curing on LBA plates containing 5% sucrose (pFLP2-cre carries sacB as a counterselectable marker). Site-specific excision of the Tet^R marker leaves behind a 63 codon scar within the target gene, and this was confirmed using PCR (Figure S3). The Tet^S *pykA* mutant was then used as a recipient for the Tet^R marker from a pykF donor, introduced by generalized transduction with ϕ PA3 as a vehicle (30). The *pykA pykF* double mutant was confirmed using (i) PCR to demonstrate the presence of the 63 codon scar in the pykA ORF (Figure S3), (ii) the absence of a PCR-amplifiable pykF fragment in the transductants (Figure S4), and (iii) the Tet^R phenotype of the transductants.

Construction of complementing plasmids

To complement the *pykA* mutant, a region encompassing the entire *pykA* ORF and 500 bp upstream of this were PCR-amplified using primers *pykA* -pLP170-FD and *pykA*pET19m-RV. The amplified DNA fragment was digested with EcoRI and BamHI and introduced into similarly-digested pUCP20, yielding $ppykA^+$. To complement the pykFmutant, we used a different strategy. The pykF ORF is predicted to form an operon with an adjacent glycerate kinase-encoding gene, PA1499. Therefore, we PCR amplified the entire region spanning from upstream of PA1499 to the 3' end of pykF (using primers PA1499-pLP170-FD and pykF-pET19m-RV primers). The amplified DNA fragment was digested with EcoRI and HindIII and introduced into similarly-digested pUCP20, yielding $ppykF^+$.

Bacterial growth

The indicated strains/mutants were grown in LB overnight and washed twice in PBS. Cultures were inoculated to an initial OD₆₀₀ of 0.05 in 50 mL of minimal media containing the indicated carbon sources and/or antibiotics. The cultures were grown at 37°C with vigorous aeration. Samples were taken every hour for optical density measurement. Each growth curve was carried out in triplicate.

Enzyme assays

Cultures were grown in 50 mL of minimal media containing the indicated carbon sources and/or antibiotics. The cultures were grown at 37°C with vigorous aeration. After the indicated period of growth, the cells were harvested by centrifugation (3220 \times g, 15 min, 4°C) and resuspended in lysis buffer (50 mM Tris-HCl, 0.1 M NaCl, pH 7.5) containing a protease inhibitor cocktail tablet (Roche, 1 tablet per 50 mL buffer). The cells were lysed by sonication on ice and the lysates were clarified by centrifugation. Protein was determined using the Bradford assay. PK activity was measured using a lactate dehydrogenase (LDH) coupled assay. The reaction mixtures contained 50 mM Tris-HCI pH 7.5. 10 mM MgCl₂. 10 U rabbit muscle LDH, 0.2 mM NADH, 5 mM PEP and 2 mM ADP. Reactions were initiated by the addition of an equal volume of cell lysate (normalised for protein concentration) and allowed to proceed at 37ºC. NADH consumption was monitored at 340 nm. Each measurement was carried out in triplicate.

Kinetics of purified PykA

Pyruvate kinase activity was measured at 37ºC in 1 mL reactions containing 50 mM Tris-HCl, 10 mM MgCl₂, 10 U of rabbit muscle LDH, 0.2 mM NADH (pH 7.5). For the PEP titrations, [ADP] was kept constant at 2 mM. For the ADP titrations, [PEP] was kept constant at 5 mM. Reactions were initiated by addition of purified PykA to a final concentration of 0.2 µg/mL. Reaction monitored progress 340nm. was at Regulator screens were carried out using 1 mM final concentration of the indicated regulator candidates and fixed а concentration of either 2 mM ADP and 0.3 mM PEP (when screening for activators), or 2 mM ADP and 2 mM PEP (when screening for inhibitors). The effect of the "hits" identified this way on the detailed kinetics were re-measured with 1 mM of each potential regulator and variable concentrations of PEP and ADP, as above. The only exception to this was that F6P, R5P and X5P were added at 0.2 mM. 0.15 mM and 0.5 mM final concentration, respectively, in all experiments. Graphpad prism 7 was used to analyse the data and to extract the kinetic constants. All experiments were carried out in triplicate.

Western blotting

Anti-PykA and anti-PykF antibodies were raised against purified PykA and PykF proteins, respectively (Biogenes.De). The PykA and PykF antisera were pre-absorbed onto an acetone extract of whole lysate protein from a *pykA* mutant or from a *pykF* mutant, respectively. The primary anti-PykA and anti-PykF antibodies were used at 1:2,000 dilution and 1:3,000 dilution, respectively. HRP-conjugated goat antirabbit (1:10,000) was used as a secondary antibody. The *iso*citrate dehydrogenase isozyme, ICD, was used as a loading control. Antibodies raised against ICD has been described previously (31).

Cloning, expression and purification of PykA and PykF

The ORFs encoding PykA (PA4329) and PykF (PA1498) were PCR-amplified from *P. aeruginosa* genomic DNA using primers *pykA*-pET19m-FD and *pykA*-pET19m-RV (for *pykA* amplification) and *pykF*-pET19m-FD and *pykF*-pET19m-RV (for *pykF* amplification). The amplified ORFs were cloned into pET-19m, generating pET-19m (pykA) and pET-19m (pykF), respectively. Expression from pET-19m yields a TEV protease-cleavable N-terminally His₆-tagged protein product. For protein expression, E. *coli* BL21 (DE3) containing pET-19m (*pykA*) or pET-19m (*pykF*), as appropriate, was grown in 1L of LB containing 50 µg/ml carbenicillin and 34 µg/ml chloramphenicol at 37°C until the OD₆₀₀ reached 0.5. The temperature was then decreased to 20°C and protein expression was induced by adding isopropyl- β -D-thiogalactopyranoside (IPTG) to a final concentration of 1mM. Following overnight growth, the cells were harvested by centrifugation (4000 \times g, 30 min, 4°C) and re-suspended in 50 mL icecold lysis buffer (50 mM Tris-HCl, 200 mM NaCl, 10% (v/v) glycerol, 10 mM imidazole (pH 8.0)) containing a dissolved EDTA-free protease inhibitor cocktail tablet (Roche). The resuspended cells were lysed by sonication on ice and the sample was then clarified by sedimentation $(10,000 \times q, 30)$ min, 4°C). The clear supernatant was loaded onto an Ni-NTA column that had been previously pre-equilibrated with lysis buffer. The column was washed overnight with lysis buffer. The bound protein was eluted in 50 mM Tris-HCl, 200 mM NaCl, 10% (v/v) glycerol, 250 mM imidazole (pH 8.0). The eluted protein was mixed with His6-tagged TEV protease (1 mg per 20 mg target protein) and loaded into a dialysis bag (10,000 MWCO). The sample was dialysed overnight against 20 mM Tris-HCl, 100 mM NaCl, 5% (v/v) glycerol, 1 mM DTT, 0.1 mM EDTA (pH 7.5). After dialysis, the protein was transferred into a Falcon tube and incubated end-over-end for 2 hr at 4°C with a slurry of Ni-NTA that had been previously pre-equilibrated in dialysis buffer. Unbound protein was concentrated by ultrafiltration (Vivaspin column, 30,000 MWCO) then aliquoted and flash frozen in liquid nitrogen. The protein concentration was determined by spectrophotometry using a calculated molar extinction coefficient of 24,410 M⁻¹cm⁻ ¹ for PykA and 25,440 M⁻¹cm⁻¹ for PykF.

Analytical ultracentrifugation (AUC)

Purified PykA was dialyzed against 20 mM Tris-HCl, 100 mM NaCl and 0.1 mM EDTA (pH 7.5) for glycerol removal. The centrepieces of the Epon double-sector of a Beckman Optima XL-I (AN-60 Ti rotor) were filled with 400 μ L of purified protein or blank buffer. The sample was sedimented at 29,160 \times *g*, 24 h, 20°C. Absorbance data (280 nm) were acquired every 2 min with interference scans collected every minute. Data analysis and the calculations of buffer viscosity, protein partial specific volumes and frictional rations were performed using SEDFIT and SEDNTERP (32, 33).

Amino acid sequence analysis

Amino acid sequences were extracted from UniProt in FASTA format, aligned by Clustal Omega (34, 35) and displayed by ESPript (36).

Crystallization of PykA

PykA was crystallized using the sitting drop vapour diffusion method. PykA (22 mg/mL in 20mM Tris-HCl, 100 mM NaCl, 5% (v/v) glycerol, 1 mM DTT, 0.1 mM EDTA, 20 mM MgCl₂, 2 mM G6P, 2 mM PEP (pH 7.5)) was mixed 1:1 with the reservoir buffer (200 nL each), which contained 20% (w/v) PEG3350, 0.1 M BIS-TRIS propane, and 0.2 M disodium malonate (pH 7.5). Crystals grew within one week. The crystals were mounted on nylon loops and cryoprotected in mother liquor supplemented with 40% (v/v) glycerol before being flash frozen in liquid N₂.

X-ray diffraction and structure refinement

Diffraction data were collected at the Diamond Light Source (Didcot, UK) on beamline IO4 (MX14043-47). The PykA structure was obtained from molecular replacement with Phaser (37) using the pyruvate kinase of T. brucei (4HYV) as a structural template. Coot (38) was used for building, and refinement was model performed by BUSTER (39). Table 2 shows the data collection and refinement statistics. Structural coordinates were deposited in the PDB with accession code (6QXL). PDBePISA (40) was used for analysis of the tetramer and ligand interfaces. Figures were generated using CCP4mg and PyMOL (41, 42).

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Author contributions

YA carried out the laboratory work and drafted the manuscript; PB assisted in phasing and solving the crystal structure; JG, XC and TR carried out the computational screening and analyses; MW conceived of the study and assisted in preparation of the manuscript.

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PEP titration	No	0.2 mM	0.15 mM	1 mM G3P	1 mM G6P	1 mM KDPG	0.5 mM X5P	1 mM RL5P	100 mM	100 mM	100 mM
	additive	F6P	R5P						KCI	NH₄CI	NaCl
S (mM)	0.670 +	0.150 ±	0.105 ±	0.207 ± 0.017	0.000 +	0.420 ±	0.000 +	0.060 +	1 107 ⊥	1 402 ± 0 084	1 402 ± 0 116
$S_{0.5}$ (IIIIVI)	0.070 ±	0.159 ±	0.195 ±	0.297 ± 0.017	0.290 ±	0.432 ±	0.209 ±	0.269 ±	1.107 ±	1.402 ± 0.064	1.492 ± 0.116
	0.058	0.021	0.017		0.026	0.036	0.011	0.012	0.086		
Hill Coeff (h)	2.14 ±	1.45 ±	1.65 ±	1.89 ± 0.21	1.78 ± 0.32	1.52 ± 0.18	1.87 ± 0.22	1.61 ± 0.12	2.36 ± 0.28	2.31 ± 0.22	2.42 ± 0.32
	0.34	0.31	0.25								
V _{max} (mM/min)	0.099 ±	0.096 ±	0.092 ±	0.092 ± 0.002	0.100 ±	0.100 ±	0.110 ±	0.110 ±	0.087 ±	0.130 ± 0.003	0.120 ± 0.004
	0.004	0.004	0.003		0.003	0.004	0.002	0.002	0.002		
L. (1)	101.0	440.0	101.1	101.1	400.0	400.0	470.0	470.0	070 0	507.0	500.0
$\mathbf{K}_{cat}(\mathbf{S}^{-1})$	431.9	418.8	401.4	401.4	436.3	436.3	479.9	479.9	379.6	567.2	523.6
k _{cat} /S _{0.5}	644.1	2626	2055	1348	1504	1008	2286	1778	319.8	404.6	350.9
(s ⁻¹ mM ⁻¹)											
ADP titration	No	0.2 mM	0.15 mM	1 mM G3P	1 mM G6P	1 mM KDPG	0.5 mM X5P	1 mM RL5P	100 mM	100 mM	100 mM
	additive	F6P	R5P						KCI	NH₄CI	NaCl
<i>K_M</i> (mM)	0.070 ±	N/A	N/A	N/A	N/A	N/A	N/A	N/A	0.147 ±	0.155 ± 0.026	0.146 ± 0.033
	0.015								0.020		
V _{max} (mM/min)	0.080 ±	N/A	N/A	N/A	N/A	N/A	N/A	N/A	0.089 ±	0.067 ± 0.003	0.078 ± 0.005
	0.005								0.004		
k _{cat} (s ⁻¹)	349.0	N/A	N/A	N/A	N/A	N/A	N/A	N/A	388.3	292.3	340.3
k _{cat} /K _M	4952	N/A	N/A	N/A	N/A	N/A	N/A	N/A	2633	1876	2321
(s⁻¹mM⁻¹)											

Table 1. The effect of metabolic regulators and monovalent ions on PykA activity.

Kinetic parameters were calculated using GraphPad Prism from best-fit nonlinear regression analysis of the data. The allosteric sigmoid function was used for the PEP titration and the Michaelis-Menten function was used for ADP titration. Abbreviations: F6P, fructose 6-phosphate; R5P, ribose 5-phosphate; G3P, glyceraldehyde 3-phosphate, G6P, glucose 6-phosphate; KDPG, 2-*keto*-3-deoxy-6-phosphogluconate; X5P, xylulose 5-phosphate; RL5P, ribulose 5-phosphate. N/A: not applicable (the regulator had no significant effect on PykA kinetics compared with the control (no additive present)). The values of k_{cat} were calculated using [E_t] = [PK monomer].

Table 2. Crystallographic data collection and refinement statistics of PykA. Values in parentheses are for the highest resolution shell.

PDB code	6QXL				
Synchrotron/X-ray source	Diamond Light Source				
Beamline	IO4-1				
Data collection					
Wavelength (Å)	0.9159				
Resolution range (Å)	405.44-2.43 (2.49-2.43)				
Space group	P31 2 1				
Unit cell					
a, b, c, (Å)	182.48, 182.48, 405.04				
a, b, g (°)	90, 90, 120				
Total reflections	4318494 (226510)				
Unique reflections	292996 (21477)				
Multiplicity	14.7 (10.5)				
Completeness (%)	100.0 (100.0)				
Mean I/sigma(I)	11.1 (1.1)				
Resolution (Å) at mean l/sigma(l) > 2	2.65 Å				
Wilson B-factor	55.5				
R-merge	0.143 (2.210)				
R-meas	0.148 (2.324)				
CC1/2	0.999 (0.642)				
Refinement					
Resolution range (High res) (Å)	158.03 – 2.43 (2.49-2.43)				
Reflections used in refinement	289310 (21209)				
Reflections used for R-free	14331 (1071)				
R-work	0.227 (0.298)				
R-free	0.251 (0.315)				
Number of molecules in the ASU:	12				
Number of non-hydrogen atoms					
macromolecules	43,331				
Ligands	306				
Solvent	2268				
Protein residues					
RMS (bonds) (Å)	0.014				
RMS (angles) (°)	1.64				
Ramachandran favoured (%)	96.98				
Ramachandran allowed (%)	2.8				
Ramachandran outliers (%)	0.23				
Average B-factor					
macromolecules	55.65				
Ligands	62.51				
Solvent	50.59				

Figure legends

Figure 1: PykA is the dominant PK in P. aeruginosa. Upper panel. PK activity of the WT and indicated mutants grown in M9 minimal media supplemented with 20 mM glucose, 30 mM glycerol or 40 mM acetate. Where indicated, the pykA mutant, the pykF mutant, or the pykA pykF double mutant were complemented with a plasmid expressing pykA (denoted $ppykA^+$) or pykF(denoted $ppykF^{+}$). Cell cultures were collected for measurement of enzyme activity after 10 h of growth. Middle panel. Western blots showing expression of PykA and PykF (as indicated) in the different carbon sources. Isocitrate dehydrogenase (ICD) was probed using anti-ICD antibodies as a loading control in each blot. The doublet observed when anti-PykA antibodies were used to probe cell extracts of the pykF mutant expressing pykF in trans indicates weak cross-reactivity of the anti-PykA antibodies with PykF (the lower of the bands in the doublet being PykF). No such cross-reactivity was apparent with the anti-PykF antibodies. Lower panel. Growth of the indicated PK mutants (± complementation) in different carbon sources. Data represent means ± SD for three independent biological replicates.

Figure 2: Kinetic characterization of **PykA.** (A),(B) PykA kinetics with respect to PEP and ADP. The PEP titration was carried out using a saturating concentration of ADP (2 mM) whereas the ADP titration was carried out using a saturating concentration of PEP (5 mM). (C) Effects of monovalent and divalent cations on PykA Activity. MgCl₂ and KCl were added at 10 mΜ and 100 mΜ concentration, respectively. PEP and ADP were at saturating initial concentration (5 mM and 2 mM, respectively). (D) The effect of putative metabolic regulators on PykA activity at low [PEP] (0.3 mM). [ADP] was fixed at 2 mM. The rationale here was to identify potential activatory molecules. Potential regulators were added at a final concentration of 1 mM, except for F6P, R5P and X5P which were added at 0.2 mM, 0.15 mM and 0.5 mM concentration, respectively. Data of figures A, B, C and D

represent the mean and standard deviation of three independent experiments. (E) Lineweaver-Burk plots demonstrating that the indicated metabolic molecules primarily act to decrease $S_{0.5}$ of PykA compared with control (without regulators).

Figure 3: X-ray crystal structure of PvkA. (A) The PvkA homotetramer. The A-A and C-C interfaces are shown. Substrate and G6P binding sites are indicated in the top left protomer. (B) Domain organization of a PykA protomer showing malonate bound in the active site and G6P bound in the allosteric site. (C) Close-up of the PykA active site. The network of interactions involved in holding the malonate-Mg²⁺ (orange mesh, Fo-Fc map contoured at 3σ) and water molecules are shown. (D) Superposition of the side chains involved in malonate binding in PykA (green) and pyruvate binding in rabbit muscle PK (1F3W, light blue). (E) Superposition of the side chains involved in malonate binding in **PykA** aeruginosa with Ρ. (green) sidechains present in the active site of E.coli PykF (1PKY, yellow).

Figure 4: The G6P binding site in PykA.

(A) Superposition of the allosteric site in PykA from *P. aeruginosa* (light green) and PK from *M. tuberculosis* (5WSB, blue). The G6P in PykA is shown in coral whereas the AMP and G6P in Mtb PK are shown in pink and yellow, respectively. (B) Superposition of the allosteric pocket in G6P-bound PykA (light green) and unbound PykF from E. coli (yellow). The G6P is shown surrounded by an electron density map (orange mesh, Fo-Fc map contoured at 3σ). (C) Schematic of the interactions involved in binding G6P to PykA. (D) Interaction of the phosphate loop (light green) and ring loop (blue) in the G6P-binding site of PykA. The G6P is shown as magenta sticks.

Figure 5: The inter-protomer interfaces in PykA. Comparison between (A) the A-A interface and (B) the C-C interface in PykA and *E. coli* PykF. Interactions present in both PykA and PykF are shown as solid black lines. Interactions unique to PykA only are shown as black dashed lines. Interactions unique to PykF only are shown as yellow dashed lines. (C) Close-up view of the A-A interface in PykA showing Arg₂₉₆ oriented towards the active site. (D) Closeup view of the A-A interface in *E. coli* PykF showing Arg₂₉₂ (equivalent to Arg₂₉₆ in PykA) oriented away from the active site. (E) The C-C interface in PykA. G6P is depicted as large magenta spheres. (F) The C-C interface in *E. coli* PykF.

Figure 6: PykA is primarily regulated by PPP metabolites. Pathways of glucose metabolism in P. aeruginosa are indicated by black (EDP), blue (PPP) and vellow (EMP) arrows, respectively. Identified PykA regulators are boxed and shown in green with the respective $S_{0.5}$ indicated below each regulator. Fructose 6-phosphate and glyceraldehyde 3-phosphate are depicted as part of the PPP to emphasize the importance they play in this series of sugar interconversions. The S_{0.5} of PykA without regulators is shown in a violet box. Abbreviations: RpiA, ribose 5-phosphate isomerase; Rpe, ribulose phosphate 3epimerase: TktA. transketolase: Tal. transaldolase; Fda. fructose 1.6bisphosphate aldolase; Fbp, fructose 1,6bisphosphatase; KDPG, 2-keto-3-deoxy-6phosphogluconate.

Figure 7. Spatial disposition of key secondary structural elements in P. aeruginosa PykA and E. coli PykF. Note how the movement of the ring loop in PvkA leads to disruption of the Ca4-Ca4 interaction and concomitant outward movement of Ca1. This, in turn, tugs on the Aa8-Ca 1 loop, engaging it with the C-C interface. The resulting movement of Aa8 directly or indirectly affects the A-A interface by forming new interactions with $A\alpha 6$ - $A\alpha 6$ '. Given that $A\alpha 6$ ' forms part of the active site, this set of interactions provides a plausible mechanism by which G6P binding to the allosteric pocket in the structure can lead to altered catalysis.



Figure 1



Figure 2







Figure 4



Figure 5



Figure 6



Figure 7