The Trypanocidal Drug Suramin and Other Trypan Blue Mimetics Are Inhibitors of Pyruvate Kinases and Bind to the Adenosine Site

Ehrlich’s pioneering chemotherapeutic experiments published in 1904 (Ehrlich, P., and Shiga, K. (1904) Berlin Klin. Wochenschrift 20, 329–362) described the efficacy of a series of dye molecules including trypan blue and trypan red to eliminate trypanosome infections in mice. The molecular structures of the dyes provided a starting point for the synthesis of suramin, which was developed and used as a trypanocidal drug in 1916 and is still in clinical use. Despite the biological importance of these dye-like molecules, the mode of action on trypanosomes has remained elusive. Here we present crystal structures of suramin and three related dyes in complex with pyruvate kinases from Leishmania mexicana or from Trypanosoma cruzi. The phenyl sulfonate groups of all four molecules (suramin, Ponceau S, acid blue 80, and benzothiazole-2,5-disulfonic acid) bind in the position of ADP/ATP at the active sites of the pyruvate kinases (PYKs). The binding positions in the two different trypanosomatid PYKs are nearly identical. We show that suramin competitively inhibits PYKs from humans (muscle, tumor, and liver isoenzymes, $K_i = 1.1–17 \, \mu M$), $T. \, cruzi$ ($K_i = 108 \, \mu M$), and $L. \, mexicana$ ($K_i = 116 \, \mu M$), all of which have similar active sites. Synergistic effects were observed when examining suramin inhibition in the presence of an allosteric effector molecule, whereby $IC_{50}$ values decreased up to 2-fold for both trypanosomatid and human PYKs. These kinetic and structural analyses provide insight into the promiscuous inhibition observed for suramin and into the mode of action of the dye-like molecules used in Ehrlich’s original experiments.

Therapies against trypanosomatid-borne diseases have evolved from the historic work of Paul Ehrlich (1), who was the first to provide a rationale for a chemotherapeutic approach in the treatment of infectious diseases. He used a series of naphthale dyes related to trypan red and trypan blue (see Fig. 1a) and demonstrated that they had trypanocidal activity. Trypan blue successfully cleared trypanosomatid infections in mouse models but were not successful in other mammals. Ehrlich noted in his mouse trials where he tested over 50 related diazo dyes that mice treated with trypan red retained a red color for weeks and even months. This side effect was a significant drawback to its use in human therapy, and the search for a colorless analog was pursued by chemists at Bayer, who synthesized Bayer 205, now known as suramin (see Fig. 1c), which is still used in the treatment of human African trypanosomiasis (sleeping sickness) caused by the parasitic protozoan Trypanosoma brucei (2). Additional uses for suramin have been explored in the treatment of human cancers (3) and HIV infection (4), where it was the first drug to show antiviral activity (5). It is evident from the mass of literature that suramin is a promiscuous inhibitor of many enzymes and receptors and shows a range of interesting clinically relevant effects (6, 7). Despite following none of the currently accepted criteria for drug-likeness (8), suramin and a number of diverse analogues (9, 10) are still of clinical interest in an ever growing repertoire of diseases (11–13).

Suramin has been characterized as an inhibitor of $T. \, brucei$ and mammalian glycolytic enzymes (14–16) and has been shown to inhibit all seven of the $T. \, brucei$ glycolytic enzymes isolated from glycosomes (peroxisome-like organelles specific for kinetoplastid protists that harbor the first seven glycolytic enzymes found in the cytosol (phosphoglycerate mutase, enolase, and pyruvate kinase (PYK)) (17)) were not examined in these experiments. The importance of glycolytic enzymes for the viability of $T. \, brucei$ has been confirmed by RNAi knockdown of glycosomal and cytosolic enzymes (including PYK) with consequent death of the parasite (18, 19).

The abbreviations used are: PYK, pyruvate kinase; $Lm$PYK, $L. \, mexicana$ PYK; $Tc$PYK, $T. \, cruzi$ PYK; $LPYK$, human liver PYK isoenzyme; $M1$PYK, human muscle PYK isoenzyme; $M2$PYK, human embryonic/tumor PYK isoenzyme; $F16BP$, fructose 1,6-bisphosphate; $F26BP$, fructose 2,6-bisphosphate; BDS, benzothiazole-2,5-disulfonic acid; PTS, 1,3,6,8-pyrenetetrasulfonic acid; DMSO, dimethyl sulfoxide; r.m.s., root mean square; h, human.

‡ The on-line version of this article (available at http://www.jbc.org) contains supplemental text, Table S1, and Figs. S1–S4.

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PYK catalyzes the final reaction of glycolysis in which P-enolpyruvate and ADP are converted into pyruvate and ATP. All trypanosomatid PYKs share a high degree of sequence identity (73–80%), with highly conserved active and allosteric effector binding sites (supplemental Fig. S1). We have previously characterized the structure of PYK from the trypanosomatid parasite, *Leishmania mexicana* (*Lm*PYK), a homotetrameric enzyme (20) in which each monomer of 498 residues is composed of four domains (see Fig. 2a). Adjacent C-domains form the C-C or “small” interface, and bordering A-domains form the A-A or “large” interface. The B-domain acts as a mobile lid at one end of the (α/β)₅-barrel A-domain, and the active site lies in the cavity between them.

Here we report the x-ray structures of four chemically related dye-like molecules including suramin in complex with *L. mexicana* or *Trypanosoma cruzi* PYKs. These structures all share similar binding modes, overlapping with the site of the adenine ring of ADP/ATP at the active site. The structural data in combination with detailed biochemical studies of the interaction of suramin with three human PYK isoenzymes and with two parasite PYKs improve our understanding of the polypharmacology of suramin (7) and provide potential scaffolds for the development of more potent isoform-specific analogues. The structures provide an interesting link to Ehrlich's groundbreaking work over a century ago and show one way in which the phenylsulfonate dyes used in the original work interact with a trypanosomatid protein. Indeed, despite the ubiquitous use of such dyes as biological markers and the uninterrupted use of suramin as a prescription drug since 1916, there has previously been very little information on the nature of the interactions of this important class of phenylsulfonates with biological macromolecules. The structural and binding data presented here are consistent with the phenylsulfonate moieties acting as adenosine analogues.

**Experimental Procedures**

**Expression, Purification, Crystallization, and Data Collection**—*Lm*PYK was overexpressed in *Escherichia coli* and purified as described previously (20), and *Tc*PYK was overexpressed in *E. coli* and purified as described (see supplemental text T1). Purified *Lm*PYK and *Tc*PYK samples (30 mg ml⁻¹ in 20 mM triethanolamine-HCl buffer (pH 7.2)) were diluted to 7–15 mg ml⁻¹ using 20 mM triethanolamine-HCl buffer (pH 7.2). For co-crystallization experiments, the following concentrations of ligands were added to the protein sample: (a) *Lm*PYK-1,3,6,8-pyrenetetrasulfonic (PTS) acid (1 mM, final concentration is 0.5 mM in the crystallization drop) and suramin (2 mM, final concentration is 1 mM in the crystallization drop); (b) *Lm*PYK-acid blue 80 ((*Lm*PYK-AB80) ~250 μM, based upon 40% purity, final concentration is ~125 μM in the crystallization drop); (c) *Lm*PYK-benzothiazole-2,5-disulfonic acid (1 mM, final concentration is 0.5 mM in the crystallization drop); (d) *Lm*PYK-Ponceau S and *Lm*PYK-acid blue 25 (1 mM Ponceau S and 500 μM acid blue 25 (based on 50% purity), the final concentration in the crystallization drop is 500 and 250 μM, respectively); and (e) *Tc*PYK-Ponceau S (1 mM, final concentration is 0.5 mM in the crystallization drop). Crystals of both *Lm*PYK and *Tc*PYK complexes were obtained at 4 °C by vapor diffusion using the hanging drop technique. The drops were formed by mixing 1.5 μl of protein solution with 1.5 μl of a well solution, composed of 7–16% PEG 8000, 20 mM triethanolamine-HCl buffer (pH 7.2), 50 mM MgCl₂, 100 mM KCl, and 10–25% glycerol. The drops were equilibrated against a reservoir filled with 0.5 ml of well solution. Crystals grew to maximum dimensions after 1 week. Prior to data collection, crystals were equilibrated for 14 h over a well solution composed of 9–14% PEG 8000, 20 mM triethanolamine-HCl buffer (pH 7.2), 50 mM MgCl₂, 100 mM KCl, and 25% glycerol, which eliminated the appearance of ice rings. Intensity data were collected (φ scans were 0.2° over 100°) at the Diamond synchrotron radiation facility in Oxfordshire, UK on beamline IO3 from a single crystal flash-frozen in liquid nitrogen at ~17 °C. Data were then processed with MOSFLM (21) and scaled with SCALA (22).

**Structure Determination**—The *Lm*PYK structures were solved by molecular replacement using the program PHASER (23). A monomer from the previously determined tetrameric structure of *Lm*PYK (*IPKL (24)) was divided into two search regions. Region 1 (residues Pro-87–Pro-187, which is equivalent to a complete B-domain) and region 2 (residues 1–86, 188–481, and 489–498) served as search models. The resulting model was divided into three rigid body domains (B-domain = 87–187, A-domain = 1–86 and 187–356, C-domain = 358–498) and subjected to 10 cycles of rigid body refinement using the program REFMAC (25). The *Tc*PYK-Ponceau S structure was solved using an identical method to that of *Lm*PYK, and the sequence was adjusted using COOT to correspond with the published *Tc*PYK amino acid sequence (accession number EFZ25721.1). When appropriate, ligands and water molecules were added to the models. Models were then subjected to cycles of restrained refinement with manual adjustments to ligands and side chains using the program COOT (26). Figures were generated using PyMOL (27). A more detailed description of the refinement processes for each of the different *Lm*PYK and *Tc*PYK crystal structures can be found in the supplemental material.

The atomic coordinates of the *Lm*PYK-PTS/suramin (3PP7), *Lm*PYK-AB80 (3QV6), *Lm*PYK-BDS (3QV8), *Lm*PYK-Ponceau S (3QV7), and the *Tc*PYK-Ponceau S (3QV9) complexes have been submitted to the Protein Data Bank (PDB). Superpositions of PYK structures were performed using both PyMOL and CCP4 superpose (27, 28).

**Suramin Inhibition Assays**—*Lm*PYK inhibition was monitored by either a luciferase (as described previously (29)) or a lactate dehydrogenase-coupled assay.³ For the lactate dehydrogenase assay, the following was added to a 1-ml cuvette: 780 μl of assay mix (1× assay buffer (50 mM triethanolamine buffer, pH 7.2, 100 mM KCl, 3 mM MgCl₂, 10% glycerol), 0.2 mM NADH (Roche Applied Science, 128023), 3.2 units/ml lactate dehydrogenase (Sigma, 61309)), 1.6 units/ml *Lm*PYK (one unit will convert 1.0 μmol of P-enolpyruvate to pyruvate per min at pH 7.0 at 25 °C in the absence of fructose 2,6-bisphosphate (F₂BP)), 10% DMSO, and 200 μl of an inhibitor solution (made up with 1× assay buffer and 10% DMSO). The mixture was incubated for 5

³ Assay mix and working stocks of ligands (including inhibitor dilution series) were made fresh on the same day the assays were performed.
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min before the reaction was started by the addition of 20 μl of 20 mM P-enolpyruvate and 10 mM ADP (final concentration = 0.4 mM P-enolpyruvate (Sigma, 79430) and 0.2 mM ADP (Sigma, A4386), made up with 1× assay buffer and 10% DMSO). The substrate concentrations used for the inhibitor assay were subsaturating, unlike in the standard activity assay in which 2.5 mM P-enolpyruvate and 2.0 mM ADP are used (30). The mixture was gently agitated, and the decrease in absorbance at 340 nm was measured for 2 min (using LAMBDA Bio, PerkinElmer Life Sciences).

The initial rate was then calculated using the UV kinlab software module. A control rate with no inhibitor present was also determined. The rate for each inhibitor assay was expressed as a percentage of the control assay. Despite considerable light absorbance at 340 nm resulting from high concentrations of suramin (>100 μM), the data still showed inhibition of 22 and 40% at 50 and 100 μM concentrations, respectively. \(K_i\) values were calculated using the IC\(_{50}\)-to-\(K_i\) server (31). The results provided are for competitive inhibition with ADP using the following parameters: for \(Lm\)PYK and \(Tc\)PYK, \(c\) = 0.1 mM, \(S_{ADP} = 0.1\) mM, \(K_{m}(ADP) = 0.26\) mM, IC\(_{50} = 160\) μM for \(Lm\)PYK and 150 μM for \(Tc\)PYK; for human PYKs (hPYKs), \(c\) = 0.1 mM, \(S_{ADP} = 0.1\) mM, \(K_{m}(ADP) = 0.1\) mM, IC\(_{50} = \)values shown in Fig. 3, \(c\) and \(d\). IC\(_{50}\) values were determined by fitting the data to Equation 1 (a four-parameter logistic model) using the KaleidaGraph version 3.6 software (Synergy Software, Reading, PA).

\[
a + (b - a)/(1 + 10^{(x - c) \times d}) \quad \text{(Eq. 1)}
\]

where \(a\) is the maximum response; \(b\), is the baseline response; \(c\) is the drug concentration that provokes a response halfway between baseline and maximum; and \(d\) is the slope. The concentrations of PYK samples were determined at \(A_{280}\) using \(ε = 20480\ M^{-1}\ cm^{-1}\). Suramin was screened against lactate dehydrogenase in the absence of PYK to avoid a false positive.

RESULTS AND DISCUSSION

Identification of Phenyl Sulfonate Dye-like Molecules as Ligands of Trypanosomatid PYKs—An analysis of the ATP binding sites observed in the x-ray structures of PYKs from a number of different species showed conserved positions for bound sulfate ions (32), corresponding to the phospho positions of ATP bound to R-state \(Lm\)PYK (20). A pharmacophore-searching approach was used as a drug discovery strategy to identify small molecules with a similar distribution of charged groups. Searching the small-molecule database EDULISS (33) of ~5 million compounds identified a series of sulfonof groups-containing molecules, many of which are reminiscent of trypan blue and other trypanocidal azo dyes used in Ehrlich’s original mouse experiments (1) (Fig. 1). One particular family of acid blue compounds selected by the search (e.g. acid blue 120, 129, and 161) has a substructure similar to Cibacron blue (Fig. 1f), which has previously been used to purify PYKs by affinity chromatography (34). Co-crystallization studies of four of the molecules identified in the search (Fig. 1, b–e) are described below. The designed synthesis of the drug suramin in 1916 was based on the molecular scaffold of the trypan blue series. We therefore asked the question whether \(Lm\)PYK (which has 74% sequence identity with \(T.\) brucei PYK (\(Tb\)PYK)) would also bind suramin.

Trypanosomatid PYKs Co-crystallized with Phenyl Sulfonate Dye-like Molecules Adopt Inactive Conformations—Co-crystallization trials of \(Lm\)PYK with trypan blue (Fig. 1a) yielded only microcrystals; however, trials with AB80 (Fig. 1e) resulted in crystals that diffracted to 2.85 Å resolution. One of the smaller sulfonof group compounds, benzothiazole-2,5-disulfonic acid (BDS, Fig. 1b), was also successfully co-crystallized. In addition, both \(Tc\)PYK and \(Lm\)PYK were crystallized in the presence of the diazo dye, Ponceau S (a stain commonly used for the reversible staining of protein bands on nitrocellulose membranes) (Fig. 1d). \(Lm\)PYK in the presence of 1 mM suramin (Fig. 1c) yielded poor quality crystals that diffracted weakly to a resolution of 5–7 Å and contained at least one exceptionally long (>700 Å) cell edge, similar to that seen for the \(Lm\)PYK-F26BP complex (20). The addition of 1,3,6,8-pyrenetetrasulfonic acid has been shown to improve the \(Lm\)PYK crystal quality by providing a helpful non-covalent cross-link within the crystal lattice (35), and 0.2 mM PTS was therefore included in the crystallization conditions with suramin. The structure of the \(Lm\)PYK, PTS, and suramin complex was determined at 2.35 Å resolution (see Table 1 for data collection and refinement statistics). See supplemental text T2 and supplemental Fig. S4 for more detailed descriptions of the sulfonof acid complexes.

The A- and C-domains (residues 18–86 and 188–480) of the \(Lm\)PYK-PTS/suramin, \(Lm\)PYK-AB80, and \(Lm\)PYK-BDS tetraramers were superimposed onto the inactive \(Lm\)PYK 1PKL (24) structure (chains E, F, G, and H), resulting in r.m.s. fits for all Cα atoms of 0.83, 0.96, and 0.76 Å, respectively, indicating that all sulfonof acid complexes were in a similar inactive (T-state) conformation (20).

Suramin, AB80, Ponceau S, and BDS Binding Sites Overlap with the ATP Sites of Trypanosomatid PYKs—Within the complex (\(Lm\)PYK-PTS/suramin), electron density for suramin was observed in both active sites of the asymmetric unit, although only one of the two monomers exhibited strong and unambiguous difference density, which could be modeled as the naphthalene-1,3,5-trisulfonic acid group of one-half of the symmetrical suramin molecule (Fig. 2b), with the remaining disordered portion pointing outward toward the solvent. The interactions between the naphthalene-1,3,5-trisulfonic acid group of suramin and the A-domain of \(Lm\)PYK are shown in Fig. 3a. A comparison of suramin binding with that of ATP binding observed in \(Lm\)PYK-APT/oxalate/F26BP (PDB ID = 3HQP) (Fig. 3e) shows that the two ligand binding sites overlap. Although \(F_o - F_e\) electron density was observed in both active sites, only suramin bound to chain A was well defined. In the active site of chain B, the \(F_o - F_e\) electron density was ambiguous, indicating that suramin may be bound in more than one conformation. Of the three negatively charged sulfonof groups of suramin bound to the active site, only two provide hydrogen-bond interactions (Fig. 3a), although there is a general area of positive charge surrounding the sulfonof groups within the binding cleft (Fig. 3f). \(Lm\)PYK forms a single interaction with the sulfonof group (ii) through Lys-335 (Fig. 3a), but the majority of interactions involve the sulfonof group (i), through three hydrogen bonds. In
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FIGURE 1. Trypan dyes and other sulfonic acid compounds have similar structural features. a, two-dimensional representation of one symmetrical half (as indicated by the dashed symmetry line) of trypan blue and trypan red (trypan red contains one additional sulfono group, indicated by red lettering). b, BDS bound in the active site of LmPYK (i). The BDS molecule is shown with an unbiased $F_{o} - F_{c}$, electron density map contoured at 3 $\sigma$ (green). Hydrogen bonds are shown as dashed red lines, and stacking interactions are shown as dashed purple lines. c, one symmetrical half of the colorless trypan analog, suramin. d, co-crystal (i) of Ponceau S bound in the active site of TcPYK (ii), e, co-crystal (i) of LmPYK and the AB80 molecule. f, a superimposition between dye-like molecules and suramin. Common groups superimpose onto the structure of suramin, highlighting the chemical relationships within the series of molecules. This figure highlights two-dimensional chemical similarities and not necessarily three-dimensional binding similarities (as illustrated by the shaded ellipses in panels a–e). g, a superimposition between Cibacron blue (a molecule commonly used for affinity purification of PYK) and AB80. Shaded ellipses in panels a–e highlight key binding characteristics observed in the LmPYK-sulfonic acid structures obtained: site S1, pink ellipse, a sulfono group ($S_{(o)}$) is observed bound in a near identical position in all complexes; site S2, green ellipse, a stacking interaction with His-54 is conserved in all structures; site S3, green ellipse, a second stacking interaction with Tyr-59 and Pro-29 is observed for suramin, Ponceau S, and AB80; site S4, blue ellipse, a hydrogen bound acceptor/donor is also commonly observed. In the LmPYK-Ponceau S, the azo group forms a hydrogen bond via the hydroxyl group of Tyr-59. Only microcrystals were obtained with trypan blue, but the common PYK binding characteristics are also observed. suramin and trypan blue are symmetrical molecules, and only half the molecule is shown (as indicated by the dashed symmetry lines).

addition, His-54 stacks above the ring system of the naphthalene-1,3,5-trisulfonic acid group with the NH group of the side chain pointing toward the sulfono group (ii). The remaining hydrophobic ring of suramin is held in place through stacking interactions with Pro-29 and Tyr-59 (Fig. 3a), found outside the deep binding cleft (Fig. 3f).

Within the LmPYK-AB80 structure, only the active site of one of the two monomers exhibited strong and unambiguous difference ($F_{o} - F_{c}$) density corresponding to one complete AB80 molecule (Fig. 2c). B-factors for the AB80 molecule were similar to those observed for the protein (Table 1). In the LmPYK-BDS structure, $F_{o} - F_{c}$, electron density corresponding to the BDS molecule was clear (supplemental Fig. S2), and the BDS molecule was clearly observed in all active sites within the asymmetric unit. Ponceau S was bound in near identical positions within the active sites of both LmPYK and TcPYK (Fig. 1d and supplemental Fig. S2d), with the sulfono groups bound to site S1 occupying a similar position to that observed in all other structures described (Fig. 1, pink ellipse).

The four conserved ligand recognition features observed in the structures are highlighted in Figs. 1 and 2d by shaded ellipses. In site S1, a sulfono group (indicated by $S_{(o)}$) mimics the $\alpha$-phospho group of ATP, forming at least one hydrogen bond. In site S2, an aromatic ring system stacks above His-54. In site S3, aromatic rings form stacking interactions with Pro-29 and/or Tyr-59. In site S4, a hydrogen bond acceptor or donor forms a hydrogen bond via a water to the backbone carbonyls of Asn-51 and His-54 in the suramin structure (Fig. 3a).
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**TABLE 1**

Data collection and refinement statistics

*Values in parentheses are for highest resolution shell.*

| Data collection | LmPYK-suranin | LmPYK-AB80 | LmPYK-BDS | LmPYK-Ponceau S* | TcPYK-Ponceau S |
|-----------------|--------------|------------|-----------|------------------|-----------------|
| Space group     | I222         | C2         | I222      | C2               | C2              |
| Cell dimensions | a, b, c (Å)  | 123.8, 129.5, 165.8 | 130.1, 127.5, 118.2 | 120.6, 131.3, 167.6 | 182.1, 164.8, 123.4 |
|                | α, β, γ (%)  | 90.0, 90.0, 90.0 | 90.0, 116.2, 90.0 | 90.0, 90.0, 90.0 | 90.0, 115.8, 90.0 |
| Resolution (Å)  | 2.35         | 2.85       | 2.45      | 2.70             | 2.1             |
| Rmerge (%)      | 5.8 (35.2)   | 8.3 (54.1) | 9.4 (56.2) | 12.3 (54.2) | 9.2 (47.9) |
| lso (%)         | 22.0 (4.5)   | 11.8 (2.5) | 14.7 (4.6) | 10.2 (2.1) | 9.4 (2.4) |
| % of completeness | 98.2 (96.9) | 98.3 (97.6) | 100.0 (100.0) | 99.6 (99.0) | 98.2 (88.6) |
| Redundancy      | 6.5 (6.5)    | 4.6 (4.5)  | 7.3 (7.5)  | 3.7 (3.5) | 6.3 (3.9) |

**Refinement statistics**

| Resolution range (Å) | 44.7–2.35 | 86.1–2.85 | 213.4–2.45 | 21.4–2.7 | 52.2–2.1 |
|----------------------|------------|------------|------------|------------|------------|
| Rmerge/Rwork (%)     | 19.1 (23.1)| 23.5 (26.7)| 23.4 (26.6)| 21.5 (24.4)| 19.8 (24.9)|
| No. of atoms         | 7530       | 6745       | 6775       | 13502      | 6863       |
| Ligands              | 72         | 85         | 33         | 199        | 128        |
| K                   | 4          | 4          | 0          | 8          | 2          |
| Water                | 433        | 106        | 210        | 652        | 464        |
| Average B-factors (Å²) | 40.5     | 72.5       | 52.1       | 45.0       | 31.0       |
| Protein              | 106.8      | 67.9       | 92.9       | 43.5       | 29.9       |
| Active site ligand†  | 39.6       | 61.7       | 46.2       | 36.3       | 35.1       |
| Water                | 0.02       | 0.01       | 0.01       | 0.01       | 0.02       |
| r.m.s. deviations    | 1.69       | 0.87       | 1.26       | 0.86       | 1.92       |
| Bond lengths (Å)     | 97.0 (952/981) | 97.3 (853/877) | 96.4 (849/882) | 97.7 (1737/1778) | 96.9 (865/893) |
| Bond angles (deg)    | 99.8 (979/981) | 99.8 (875/877) | 99.8 (880/882) | 99.8 (1774/1778) | 99.8 (891/893) |
| Ramachandran         | 0.2 (2/981) | 0.2 (2/877) | 0.2 (2/882) | 0.2 (4/1778) | 0.2 (2/893) |

*S. typhi Ponceau S was crystallized in the presence of acid blue 25 (see supplemental text T1 and supplemental Figure S4).† B-factors are for ligands bound to the active site in a similar manner.

**Suramin Binds with Up to 2-Fold Higher Affinity to Effector-bound PYKs**—The strongly colored dye-like compounds AB80 and Ponceau S absorb light at wavelengths required for the available kinetic assays and make it impractical to determine IC50/Ki values accurately. For this reason, only suramin was taken forward for kinetic studies. We have measured suramin inhibition for three hPYK isoenzymes (with IC50 values of 20, 33, and 2.2 μM and Ki values of 10, 155, and 112 μM, respectively) and for two trypanosomatid enzymes, LmPYK (Fig. 3c) and TcPYK (supplemental Fig. S3) (with IC50 and Ki values of ~155 and 112 μM, respectively). To confirm these results, LmPYK inhibition was also monitored by an unrelated lactate dehydrogenase-based assay, which gave very similar results (supplemental Fig. S3).

The natural effector for hM2PYK and hLPYK is fructose 1,6-bisphosphate (F16BP), whereas the effector for LmPYK is F26BP (36–38). We observed a 2-fold reduction in the Ki value for suramin when effector is present for both hM2PYK (Ki decreases from 16.5 ± 0.4 to 8.5 ± 0.2 μM) in the presence of F16BP and LmPYK (Ki decreases from 115.6 ± 2.6 to 57 ± 1.2 μM) in the presence of F26BP. The presence of F16BP has no measurable effect on the Ki values of hLPYK (~1 μM) and hM1PYK (~9.5 μM) (Fig. 3d). The latter result is perhaps expected as hM1PYK is constitutively active and not regulated by occupancy of the effector site.

LmPYK-PTS/suramin is in the inactive conformation, which is inherently flexible when compared with that of the more rigid effector-bound active conformer, locked by the F26BP effector molecule (LmPYK-F26BP (20)). The flexibility of the inactive conformation is likely to contribute to the differential binding of suramin to the active sites (Fig. 3b). The relationship between enzyme flexibility and differential binding of active-site ligands has also been observed for oxalate and ATP binding in the absence of the effector molecule (20, 39), whereby ATP is either completely absent or bound in a different position within the active site. When the enzyme is stabilized by the effector molecule, it is inherently more rigid and adopts the optimal conformation for binding substrates. It is only within this rigidified effector-bound structure that all active-site components occupy near identical positions within all four active-site clefts of the PYK tetramer (20). The difference in conformational flexibility between effector-bound and unbound states also provides an explanation for the different Ki values observed for oxalate inhibition of human M2PYK, in the presence (Ki = 24 μM) and absence (Ki = 220 μM) of the effector molecule (37).

Consideration of in vivo conditions in human serum has important implications for PYK inhibitors acting like suramin. The binding affinity for suramin is lower (Ki = 1–17 μM) when compared with that measured for the natural substrate ADP (100 μM (40)). Furthermore, the ADP concentration in which PYK operates in vivo is ~50 μM (total cytosolic ADP = 300 μM (41) and free cytosolic ADP = 50 μM (42)), but suramin can achieve serum levels as high as 352 μM (43) and cellular levels (in certain compartments) in excess of 150 μM (44). Moreover, effector-bound PYK is a better representation of in vivo conditions as effector levels are likely to be saturating in trypanosomatids (where F26BP concentration = 15–50 μmol/g of protein (~85 μM) (45)) and in human cells (where F16BP concentration can reach 5 mM in cancer cells but is only 50 μM in non-cancerous cells (46)). The tighter binding found for both suramin and oxalate inhibitors in the presence of effector sug-
suggests that inhibitor assays for allosterically regulated forms of PYK should be carried out in the presence of effector.

Mode of Action of Suramin-like Phenyl Sulphonates—Suramin is known to interact with a diversity of targets, but x-ray crystal structures of suramin-target protein complexes have been determined for only five proteins (supplemental Table S1), three of which are enzymes. None of these enzymes binds ATP/ADP, which is unexpected as the chemotherapeutic effects of suramin are often attributed to its ability to inhibit the binding of ATP and other nucleotides (47). There is evidence to show that the trypanocidal action of suramin results from the inhibition of ATP production by glycolysis (48), an enzymatic pathway that has evolved to bind phosphometabolites. Although the binding mode has previously remained elusive, it is not surprising that suramin and the chemically related AB80 and BDS molecules are found to occupy ADP/ATP binding sites, which are present in many glycolytic enzymes, and may explain the promiscuous behavior of this compound class.

To examine binding modes, the LmPYK-PTS/suramin, LmPYK-PTS/AB80, and LmPYK-PTS/BDS monomers (residues 18–86 and 188–480, excluding the B-domain) were superposed onto the LmPYK-ATP/oxalate/F26BP (PDB ID = 3HQ) structure, with an r.m.s. fit for the Ca atoms of the A- and C-domain core of 0.50 Å. It is clear from Fig. 3e that suramin occupies the adenine binding site and blocks ADP/ATP binding. Similar positions are observed for AB80, Ponceau S, and BDS binding (supplemental Fig. S2c). In all PYK structures presented here, the adenine and ribose positions overlap with the dye molecules, but only the interactions of the α-phospho group are mimicked by one of the sulfono groups. The x-ray structure shows that the bulk of the suramin ring system sterically hinders the molecule from penetrating further into the active-site cleft and occupying the ATP-phospho position. The adenine moiety actually makes very few interactions with LmPYK (20), in part explaining the relatively weak value ($K_i = 116 \mu M$) and high B-factor (∼100 Å$^2$) of suramin. Both
AB80 (Fig. 2d) and Ponceau S (supplemental Fig. S2d) show stronger electron density and bind in a similar position to that of suramin, with key groups providing identical interactions with the protein (Fig. 1). The residues involved in suramin, AB80, Ponceau S, BDS, and ATP/ADP binding in \textit{Lm}\PYK{} are all fully conserved in both \textit{T. brucei} and \textit{T. cruzi} \PYK{}s (supplemental Fig. S1), suggesting that suramin would also inhibit both these PYKs. This was confirmed using the lactate dehydroge-

\begin{figure}[h]
\centering
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\caption{Suramin inhibition of PYK is enhanced in the presence of the activator molecule. \textit{a}, a schematic drawing showing the interatomic distances (given in Å, dotted black lines) for the interactions shown in Fig. 2b). The solid purple lines indicate stacking interactions. Residues involved in ATP binding (orange box) and not involved (green) have been indicated. Only one symmetrical half of the suramin molecule is shown (as indicated by the dashed symmetry line). \textit{b}, schematic representation of the inactive \textit{Lm}\PYK{}-suramin structure. Suramin is unambiguously (yellow lozenges) bound to one of the chains in the asymmetric unit (two chains outlined in black) but is disordered (blue square) in the adjacent chain, separated by the large (A-A) interface. A crystallographic two-fold axis running along the small interface generates two more chains (purple outlines), forming the biologically relevant tetramer structure. See Ref. 20 for schematics of the enzyme in its other conformations. \textit{c} and \textit{d}, concentration-response curves observed for the titration of suramin against PYK activity in the absence (\textit{c}) and presence (\textit{d}) of activator (F16BP for human PYKs, F26BP for \textit{Lm}PYK), using a luciferase-based assay. The values are expressed ± S.D. Suramin showed no inhibition in the control assay of luciferase activity alone. All assays were performed in triplicate. \textit{e}, the binding of suramin prevents ADP/ATP binding in trypanosomatid PYKs. The R-state \textit{Lm}PYK-ATP/oxalate/F26BP monomer (colored blue, PDB ID 3HQP) was superposed onto the inactive \textit{Lm}PYK-suramin structure (colored green). The A- and C-domains (residues 18 – 86 and 188 – 480) of both structures were superposed (r.m.s. fit of the C-\alpha atoms of domains A and C is 0.50 Å). The superpositions of the ATP and suramin molecules clash, indicating a clear mechanism of competitive inhibition. \textit{f}, the electrostatic surface of the suramin binding site, showing areas of positive charge (blue), which interact with the negatively charged sulfono groups of suramin. Two hydrophobic groups (Pro-29 and Tyr-59) provide further stability, helping to hold the molecule in the ATP/ADP binding site.

\end{figure}
nase-linked enzyme assay, whereby TcPYK was shown to be inhibited by suramin with an IC50 of 150 μM (supplemental Fig. S3). It is noteworthy that all the ligands described here share common characteristics and can all be adequately superimposed onto the structure of suramin (Fig. 1f). It may therefore be hypothesized that all the molecules are likely to bind to similar protein binding sites, which indeed appears to be the case for LmPYK.

The Trypanosomatid PYK Complexes Provide a Template for Inhibitor Design—Affinity values for suramin inhibition are noticeably different for human (K1 values are 10.0, 16.5, and 1.1 μM for M1PYK, M2PYK, and LPYK, respectively) and parasite (K1 value for LmPYK and TcPYK = ~112 μM) PYKs. Such differences in ligand affinities suggest that it may be possible to exploit the amino acid differences around this suramin/dye binding pocket (supplemental Fig. S2b) to design and develop species-specific small-molecule inhibitors. The efficacy of inhibitors could be further enhanced by the concomitant use of structural and kinetic results presented in this study show one aspect creating a general trypanosomatid inhibitor. It is now over 100 years since the discovery of the trypanocidal effects of azo dyes and the subsequent development of suramin (51). The structural and kinetic results presented in this study show one aspect of its remarkable polypharmacology and provide a template for the design of new generations of inhibitors with improved specificity.

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REFERENCES

1. Ehrlich, P., and Shiga, K. (1904) Berlin Klin. Wochenschrift 20, 329–362
2. Hawking, F. (1978) Adv. Pharmacol. Chemother. 15, 289–322
3. La Rocca, R. V., Stein, C. A., Danesi, R., and Myers, C. E. (1990) J. Steroid Biochem. Mol. Biol. 37, 893–898
4. Mahoney, C. W., Azzi, A., and Huang, K. P. (1990) J. Biol. Chem. 265, 5424–5428
5. Mitsuya, H., Popovic, M., Yarchan, R., Matsushita, S., Gallo, R. C., and Broder, S. (1984) Science 226, 172–174
6. Voogd, T. E., Vansterkenburg, E. L., Wilting, J., and Janssen, L. H. M. (1993) Pharmacol. Rev. 45, 177–203
7. McGreary, R. P., Bennett, A. J., Tran, Q. B., Cosgrove, K. L., and Ross, B. P. (2008) Mini Rev. Med. Chem. 8, 1384–1394
8. Lipinski, C. A., Lombardo, F., Dominy, B. W., and Feeney, P. J. (1997) Adv. Drug Deliv. Rev. 23, 3–25
9. Freissmuth, M., Boehm, S., Beindl, W., Nickel, P., Ijzerman, A. P., Hohenegger, M., and Nanoff, C. (1996) Mol. Pharmacol. 49, 602–611
10. Klinger, M., Freissmuth, M., Nickel, P., Stäbler-Schwarzbart, M., Kassack, M., Suko, J., and Hohenegger, M. (1999) Mol. Pharmacol. 55, 462–472
11. Braga-Silva, L. A., dos Santos, A. L., Portela, M. B., Souto-Padrón, T., and de Araújo Soares, R. M. (2007) FEMS Immunol. Med. Microbiol. 51, 399–406
12. De Clercq, E. (2009) Med. Res. Rev. 29, 571–610
13. Baghdiguian, S., and Fantini, J. (1997) Cancer J. 10, 31–37
14. Miao, O., and Oppedore, F. R. (1987) Eur. J. Biochem. 162, 493–500
15. Willson, M., Callens, M., Kunz, D. A., Peré, J., and Oppedore, F. R. (1993) Mol. Biochem. Parasitol. 59, 201–210
16. Iagielski, A. K., Kryskiewicz, E., and Bryla, J. (2006) Eur. J. Pharmacol. 537, 205–209
17. Oppedore, F. R. (1987) Annu. Rev. Microbiol. 41, 127–151
18. Drew, M. E., Morris, J. C., Wang, Z., Wells, L., Sanchez, M., Landfair, S. M., and Englund, P. T. (2003) J. Biol. Chem. 278, 46596–46600
19. Albert, M. A., Haanstra, J. R., Haanstra, V., Van Roy, J., Oppedore, F. R., Bakker, B. M., and Michel, P. A. (2005) J. Biol. Chem. 280, 28306–28315
20. Morgan, H. P., McNae, I. W., Nowicki, M. W., Hohenegger, M., Michel, P. A., Fothergill-Gilmore, L. A., and Walkinshaw, M. D. (2010) J. Biol. Chem. 285, 12892–12898
21. Potterrton, E., Briggs, P., Turkenburg, M., and Dodson, E. (2003) Acta Crystallogr. D Biol. Crystallogr. 59, 1131–1137
22. Evans, P. (2006) Acta Crystallogr. D Biol. Crystallogr. 62, 72–82
23. McCoy, A. J., Grosse-Kunstleve, R. W., Adams, P. D., Winn, M. D., Storoni, L. C., and Read, R. J. (2007) J. Appl. Crystallogr. 40, 658–674
24. Rigden, D. J., Phillips, S. E., Michel, P. A., and Fothergill-Gilmore, L. A. (1999) J. Mol. Biol. 291, 615–635
25. Murshudov, G. N., Vagin, A. A., and Dodson, E. J. (1997) Acta Crystallogr. D Biol. Crystallogr. 53, 240–255
26. Emsley, P., and Cowtan, K. (2004) Acta Crystallogr. D Biol. Crystallogr. 60, 2126–2132
27. Delano, W. L. (2010) The PyMOL Molecular Graphics System, version 1.8r1, Schrödinger, LLC, New York
28. Potterton, E., McNicholas, S., Krisine, E., Cowtan, K., and Noble, M. (2002) Acta Crystallogr. D Biol. Crystallogr. 58, 1955–1957
29. Jiang, J. K., Boxer, M. B., Vander Heiden, M. G., Shen, M., Skoumbourdis, A. P., Southall, N., Veith, H., Leister, W., Austin, C. P., Park, H. W., Inglese, J., Cantley, L. C., Auld, D. S., and Thomas, C. J. (2010) Bioorg. Med. Chem. Lett. 20, 3387–3393
30. Huantra, V., Vernal, C., Rigden, D. J., Fothergill-Gilmore, L. A., Oppedore, F. R., and Michel, P. A. (2002) FEBS Lett. 514, 255–259
31. Cer, R. Z., Mudunuri, U., Stephens, R., and Lebeda, F. J. (2009) Nucleic Acids Res. 37, W441–W445
32. Tulloch, L. B., Morgan, H. P., Huantra, V., Michel, P. A., Fothergill-Gilmore, L. A., and Walkinshaw, M. D. (2008) J. Mol. Biol. 383, 615–626
33. Hsin, K. Y., Morgan, H. P., Shave, S. R., Hinton, A. C., Taylor, P., and Walkinshaw, M. D. (2011) Nucleic Acids Res. 39, D1042–D1048
34. Subramanian, S. (1994) Crit. Rev. Biochem. Mol. Biol. 29, 169–205
35. Morgan, H. P., McNae, I. W., Hsin, K. Y., Michel, P. A., Fothergill-
Inhibitors of Pyruvate Kinase

Gilmore, L. A., and Walkinshaw, M. D. (2010) Acta Crystallogr. Sect. F Struct. Biol. Cryst. Commun. 66, 215–218
36. Valentini, G., Chiarelli, L. R., Fortin, R., Dolzan, M., Galizzi, A., Abraham, D. J., Wang, C., Bianchi, P., Zanella, A., and Mattevi, A. (2002) J. Biol. Chem. 277, 23807–23814
37. Dombrauckas, J. D., Santarsiero, B. D., and Mesecar, A. D. (2005) Biochemistry 44, 9417–9429
38. Ernest, I., Callens, M., Opperdoes, F. R., and Michels, P. A. (1994) Mol. Biochem. Parasitol. 64, 43–54
39. Larsen, T. M., Benning, M. M., Rayment, I., and Reed, G. H. (1998) Biochemistry 37, 6247–6255
40. Boxer, M. B., Jiang, J. K., Vander Heiden, M. G., Shen, M., Skoumbourdis, A. P., Southall, N., Veith, H., Leister, W., Austin, C. P., Park, H. W., Inglese, J., Cantley, L. C., Auld, D. S., and Thomas, C. J. (2010) J. Med. Chem. 53, 1048–1055
41. Veech, R. L., Lawson, J. W., Cornell, N. W., and Krebs, H. A. (1979) J. Biol. Chem. 254, 6538–6547
42. Akerboom, T. P., Bookelman, H., Zuurendonk, P. F., van der Meer, R., and Tager, J. M. (1978) Eur. J. Biochem. 84, 413–420
43. La Rocca, R. V., Cooper, M. R., Stein, C. A., Kohler, D., Ulrich, M., Weinberger, E., and Myers, C. E. (1992) Ann. Oncol. 3, 571–573
44. Huang, S. S., Koh, H. A., and Huang, J. S. (1997) FEBS Lett. 416, 297–301
45. Van Schaftingen, E., Opperdoes, F. R., and Hers, H. G. (1987) Eur. J. Biochem. 166, 653–661
46. Eigenbrodt, E., Reinacher, M., Scheefers-Borchel, U., Scheefers, H., and Friis, R. (1992) Crit. Rev. Oncog. 3, 91–115
47. Avliyakulov, N. K., Lukes, I., Kajava, A. V., Liedberg, B., Lundström, I., and Svensson, S. P. S. (2000) Eur. J. Biochem. 267, 1723–1731
48. Fairlamb, A. H., and Bowman, I. B. R. (1980) Mol. Biochem. Parasitol. 1, 315–333
49. Ashizawa, K., Willingham, M. C., Liang, C. M., and Cheng, S. Y. (1991) J. Biol. Chem. 266, 16842–16846
50. Pal, A., Hall, B. S., and Field, M. C. (2002) Mol. Biochem. Parasitol. 122, 217–221
51. Kleine, F. K., and Fischer, W. (1922) Dtsch. Med. Wochenschr. 48, 1693–1696