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# The kinetic characteristics of human and trypanosomatid phosphofructokinases for the reverse reaction

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41

## 42 ABSTRACT

43 Eukaryotic ATP-dependent phosphofructokinases (PFKs) are often considered unidirectional 44 enzymes catalysing the transfer of a phospho moiety from ATP to fructose 6-phosphate (F6P) to 45 produce ADP and fructose 1,6-bisphosphate (F16BP). The reverse reaction is not generally 46 considered to occur under normal conditions and has never been demonstrated for any eukaryotic 47 ATP-dependent PFKs, though it does occur in PPi-dependent PFKs and has been experimentally 48 shown for bacterial ATP-dependent PFKs. Evidence is provided via two orthogonal assays that all 49 three human PFK isoforms can catalyse the reverse reaction in vitro, allowing determination of kinetic 50 properties. Additionally, the reverse reaction was shown possible for PFKs from three clinically 51 important trypanosomatids; these enzymes are contained within glycosomes in vivo. This 52 compartmentalisation may facilitate reversal, given the potential for trypanosomatids to have an 53 altered ATP/ADP ratio in glycosomes compared to the cytosol. The kinetic properties of each 54 trypanosomatid PFK were determined, including the response to natural and artificial modulators of 55 enzyme activity. The possible physiological relevance of the reverse reaction in trypanosomatid and 56 human PFKs is discussed.

#### 57 INTRODUCTION

58 Phosphofructokinase (PFK) catalyses the phosphorylation of fructose 6-phosphate (F6P) to fructose 59 1,6-bisphosphate (F1,6BP) and plays a central role in the glycolytic pathway of prokaryotes and 60 eukaryotes. The enzymatic step catalysed by PFK is conserved in most organisms from Eukarya, 61 Bacteria and Archaea. Despite an enzyme mechanism that has been conserved for over 2 billion 62 years, different PFK families have evolved interestingly diverse regulatory mechanisms associated 63 with considerable differences in protein sequence and architecture. There is increasing interest in 64 PFK as a drug target in human diseases, including diabetes (1) and cancer (2,3). Additionally, 65 glycolysis is a valid therapeutic target for killing pathogens that rely exclusively on glucose catabolism 66 for their ATP supply; previous work has shown the effectiveness of this approach against 67 trypanosomatid parasites (4). PFKs from the three trypanosomatid species that cause significant 68 mortality and morbidity: Trypanosoma brucei (Human African Trypanosomiasis), T. cruzi (Chagas 69 disease), and Leishmania spp. (Leishmaniasis) are all potential drug targets.

70

71 Two main evolutionary groups of PFK are distinguished by their phospho-donor substrates. The 72 pyrophosphate-dependent group uses inorganic pyrophosphate (PPi) as the phospho donor and is 73 found in plants and certain protists, including amoeba and bacteria. The second group uses ATP as 74 the phospho donor and is found in many other bacteria and protists, plants, and all vertebrates (note 75 that plants usually contain PFKs from both groups). Phylogenetic and structural analyses 76 demonstrate that the two groups evolved from a common ancestor, though amino acid sequence 77 identities are low (~25%) (5). Despite these differences, there are similarities in catalytic mechanism; 78 in one interesting case the PP<sub>i</sub>-PFK of Entamoeba histolytica, which has a 10<sup>6</sup> fold preference for 79 inorganic pyrophosphate over ATP, could be converted to an ATP-dependent enzyme by a single 80 mutation (6). This supports the idea that the ATP-dependent PFKs are the primordial form from which, 81 on multiple occasions, the PPi-dependent enzymes evolved. However, the evolutionary path is 82 complex, with amino acid sequence comparisons suggesting the ATP-dependent PFK found in 83 trypanosomatids developed through an ancestral PPi reliant stage before switching back to ATP as a 84 substrate (7,8).

85

86 A biochemically important distinction between the two families is that all PPi-PFKs readily carry out 87 both the forward and reverse enzyme reactions under physiological conditions. The biological 88 consequence is that organisms that use PPi-dependent PFKs for the most part do not require 89 fructose-1,6-bisphosphatase enzymes to carry out the reverse dephosphorylation step required in the 90 gluconeogenic pathway. Possibly because of its ability to carry out the reaction in both directions, the 91 PPi-PFK family shows little evidence of allosteric regulatory mechanisms controlling enzyme activity, 92 though this is not universal (9). In contrast, ATP-dependent PFKs have evolved a wide range of 93 allosteric mechanisms, with associated differences in protein chain lengths and architecture.

94

95 The active form of bacterial ATP-dependent PFKs is a homo-tetramer with subunits of approximately
96 35 kDa. In yeasts, a gene duplication/fusion event yielded double-size chains and these subsequently

97 underwent additional duplications to give an octamer comprised of homologous catalytic and 98 regulatory subunits each with a molecular mass of 110-120 kDa (10). Mammals have tetrameric 99 PFKs, with subunits of 85 kDa. The N-terminal half of the double enzyme was constrained to retain 100 the catalytic function, while the substrate binding sites of the C-terminal half evolved into effector 101 binding sites (11). Higher levels of regulation in mammals are also achieved by three distinct isoforms 102 with varying properties (denoted as PFK-M, PFK-L and PFK-P) which are expressed in a tissue-103 specific pattern (12). Trypanosomatid PFKs are intermediate in size (~55kDa); these are strictly ATP 104 dependent but have amino acid sequences closer to the PPi family and can therefore be regarded as 105 chimeras. X-ray structures of trypanosomatid PFKs show major differences compared with other ATP-106 dependent PFKs; in particular, the C-terminal extensions can form long helices, acting as reaching 107 arms to hold the tetramer together (8).

108

Structural differences between ATP-dependent PFKs derive from varying requirements for allosteric regulation. The smaller bacterial ATP-dependent PFKs are activated by ADP and GDP alone (13) with *Escherichia coli* PFK used in a definitive study by Monod and co-workers to support the now classic allosteric model of enzyme kinetics (14). In trypanosomatid PFKs AMP is the only known activator while in human PFKs the non-catalytic C-domain of each isoform binds the allosteric activators AMP, ADP and fructose 2,6-bisphosphate (F26BP) (15).

115

116 The evolution of these tightly regulated allosteric effector systems in the ATP-dependent PFK family 117 contrasts with the less regulated bi-directional activity of the PPi-dependent family. For the ATP-118 dependent PFKs the forward enzymatic reaction is favoured under physiological conditions, often 119 being regarded as an essentially irreversible reaction under normal conditions. Indeed, the reverse 120 reaction (F16BP + ADP  $\Rightarrow$  F6P + ATP) has never been demonstrated for any eukaryotic ATP-121 dependent PFK, including any of the human isoforms, in vitro or in vivo. In this paper we demonstrate 122 that the reverse reaction is possible under experimental conditions for all three human PFK isoforms 123 and the three trypanosomatid PFK orthologues, and present the kinetic properties for these reactions. 124 The results suggest that the reverse reaction could occur under certain physiological conditions.

#### 125 METHODS

126 The identity of all recombinant PFKs was confirmed by SDS-PAGE showing highly pure PFKs of 127 expected molecular weights (Figure 1). Additionally, western blots and MALDI-TOF mass 128 spectrometry using a Bruker Ultraflex instrument confirmed protein identities (data not shown).

129

#### 130 Production of trypanosomatid phosphofructokinases

131 N-terminally His<sub>6</sub>-tagged trypanosomatid PFK DNA sequences, codon optimised for *E. coli* 132 expression, were inserted into pET28a or pDEST17 expression plasmids. The recombinant plasmids 133 were used to transform chemically competent E. coli cells which were grown on LB agar plates with 134 corresponding antibiotic (Table 1). Single colonies were inoculated into 500ml media in 2L conical 135 flasks and grown in a shaking incubator at 250 rpm and 37°C to an OD<sub>600nm</sub> 0.8-0.9, then cold 136 shocked at 4°C for 30 min. PFK expression was induced with 1mM isopropyl β-D-1-137 thiogalactopyranoside (IPTG) for 16h at 100 rpm and 18°C before harvesting the cells via 138 centrifugation and removal of supernatant.

139

140 Cell pellets from 1L cultures were suspended in 50ml lysis buffer (50 mM TEA, 5 mM MgCl<sub>2</sub>, 50 mM 141 KCI, 10% glycerol, pH 7.4), supplemented with Roche cOmplete™ EDTA-free Protease Inhibitor 142 Cocktail and ~5mg bovine pancreas deoxyribonuclease [Sigma-Aldrich D5025]) and lysed with 143 Constant Cell Disruption Systems at 25kPsi and centrifuged. Filtered supernatant was loaded onto a 144 cobalt-charged HiTrap 1ml FF immobilized metal affinity chromatography (IMAC) column (GE 145 Healthcare) equilibrated in wash buffer (50 mM triethanolamine (TEA), 300 mM NaCl, 20 mM 146 imidazole, 10% glycerol, pH 8.0) in a GE Healthcare ÅKTA purifier system at 6°C. Impurities were 147 removed by further wash buffer steps with gradually increasing imidazole concentrations before PFK 148 eluted with elution buffer (50 mM TEA, 300 mM NaCl, 500 mM imidazole, 10% glycerol, pH 8.0). T. 149 brucei and T. cruzi PFK (TbPFK and TcPFK) eluates were loaded onto a HiPrep SephacryI™ S-200 150 16/60 column (GE Healthcare), pre-equilibrated with gel filtration buffer (20 mM TEA, 5 mM MgCl<sub>2</sub>, 50 151 mM KCl, 10% glycerol, pH 7.4) and tetrameric fractions eluted with 1.5 column volumes (CV) of gel 152 filtration buffer. Leishmania infantum PFK (LmPFK) IMAC eluates were loaded onto a HiPrep 26/10 153 Desalting column (GE Healthcare) pre-equilibrated with gel filtration buffer and eluted using 1.5 CV 154 gel filtration buffer. Samples were concentrated to 1mg/ml with Vivaspin® 20ml 30,000kDa Molecular 155 Weight Cut-Off (MWCO) spin concentrators. Aliquots were flash-frozen and stored at -80°C until 156 required. Tag removal was attempted but only partially successful (Figure 1A); tagged protein was 157 therefore used.

158

## 159 Production of human phosphofructokinases

160 Plasmid pJJH71 (16) containing yeast codon optimised cDNA for His<sub>6</sub>-tagged PFK-M1, PFK-L1, or

161 PFK-P1 was used to transform PFK-deficient S. cerevisiae (16) via electroporation which were

162 subsequently grown on YPDA (yeast extract, peptone, dextrose, adenine broth) agar plates. Colonies

163 were transferred to 2L conical flasks containing 500ml YPDA medium with 50µ/ml carbenicillin and

164 cultures grown using an Infors HT Multitron standard shaking incubator at 30°C and 250rpm before165 harvesting the yeast via centrifugation and removal of supernatant.

166

167 Cell pellets from 2L cultures were suspended in lysis buffer (50mM TEA, 300mM KCI, 10mM 168 imidazole, 1mM TCEP, 1mM ATP/F6P) supplemented with Roche cOmplete™ EDTA-free Protease 169 Inhibitor Cocktail and ~5mg bovine pancreas deoxyribonuclease) at 8% w/v and lysed with Constant 170 Cell Disruption Systems at 40kPsi and centrifuged. Filtered supernatant was loaded onto a nickel-171 charged HiTrap 1ml FF IMAC column equilibrated in wash buffer (50mM TEA, 300mM KCI, 10mM 172 imidazole, 1mM TCEP (tris(2-carboxyethyl)phosphine), 1mm ATP/F6P, 10% glycerol) in an ÄKTA 173 purifier system at 6°C. Impurities were removed by further wash buffer steps with gradually increasing 174 imidazole concentrations before PFK eluted with elution buffer (50mM TEA, 300mM KCI, 500mM 175 imidazole, 1mM TCEP, 1mM ATP/F6P). PFK-M was further purified using a GE Healthcare HiPrep 176 Sephacryl S300 16/600 size-exclusion chromatography column pre-equilibrated with gel filtration 177 buffer (50mM TEA, 500mM KCI, 5mM MgCl<sub>2</sub>, 1mM TCEP, 1mM ATP/F6P, 10% glycerol). For PFK-L 178 and PFK-P a GE Healthcare Superose 6 10/300 size-exclusion column was used. Samples 179 corresponding to tetrameric protein (340kDa) were pooled and concentrated using a pre-equilibrated 180 20ml 30,000kDa MWCO spin concentrator to above 0.3mg/ml. Aliquots were flash-frozen and stored 181 at -80°C until required. All buffers were at pH 8, except for PFK-M purifications (pH 7.4).

182

## 183 Demonstration of reverse reaction using an endpoint assay

184 The Promega Kinase-Glo ATP assay system (V6713) was used to measure ATP production in the 185 reverse PFK reaction. Energy of the ATP produced was converted to light via the luciferase/luciferin 186 reaction in an endpoint reaction. 10µl of PFK at 2µg/ml was added to 100µl assay buffer (50mM TEA, 187 10mM MgCl<sub>2</sub>, 0.1% w/v BSA, 0.005% TWEEN20, 1% DMSO, pH 7.4) containing 5mM ADP in a white 188 non-binding 96-well plate. Incubation was carried out at 4°C for 20min, followed by 10min at room 189 temperature. 5mM F16BP (final concentration) was added and the plate was then centrifuged at 190 1000rpm for 30 seconds, and further incubated at room temperature for 60 min. 25µl of the "Kinase-191 Glo reagent" was added to each well for a final incubation period of 30min. Assay output was 192 measured as luminescence using a Molecular Devices Spectramax M5 Multi-Mode Plate Reader and 193 converted to ATP concentrations using control data from a constructed ATP titration curve.

194

### 195 Determination of kinetic characteristics using an enzyme-linked kinetic assay

196 Conversion of F16BP (Sigma F6803) and ADP (Sigma A4386) to F6P and ATP was measured using 197 an enzyme-linked assay. F6P was converted to glucose 6-phosphate (G6P) by phosphoglucose 198 isomerase (PGI), and subsequently to 6-phosphogluconolactone by glucose-6-phosphate 199 dehydrogenase (G6PD), with concurrent reduction of NAD<sup>+</sup> to NADH. Formation of NADH was 200 measured via absorbance of UV at 340nm. ATP produced was re-converted into ADP by glycerol 201 kinase (GK) (in the presence of glycerol) to keep the ATP/ADP ratio low.

202

203 15μl of assay buffer (50mM TEA, 100mM KCl, 10mM MgCl<sub>2</sub>, 1mM TCEP, 10% glycerol, pH 7.4) was
204 added to 40μl of assay mix (NAD<sup>+</sup> (Sigma NAD100-RO), G6PD (Sigma G8529), PGI (Sigma P9544),

- and GK (Sigma G6142), then 20µl of ADP titration (or 15mM ADP stock) in a clear 96-well plate. The
  plate was incubated at 25°C for 2min. 5µl of 0.2mg/ml PFK was added and the reaction initiated with
  20µl F16BP titration (or 25mM F16BP stock). UV absorbance at 340nm was measured at 13s
  intervals for 15min in a Molecular Devices Spectramax M5 Multi-Mode Plate Reader at 25°C.
- 209
- Time-dependent absorbance change was converted into rate of NADH oxidation (µM.s<sup>-1</sup>) or specific activity (µmol.min<sup>-1</sup>.mg<sup>-1</sup>) using the Beer-Lambert law (molar extinction co-efficient of NADH 6.22mM<sup>-1</sup>). Reaction rates for each well were calculated using an 8-point (104s) rolling average. Kinetic parameters of the steady-state stage of the reaction were determined with GraphPad Prism 7. Nonlinear regression analysis was performed on substrate titration data, with curves fitted using allosteric sigmoidal models, enabling determination of kinetic values.

- 216 **RESULTS**
- 217

#### 218 Proof of concept for reversal of direction of PFK reaction

219 Proof of concept for reversal of the direction of the PFK reaction was established with the 'Kinase-220 Glo'® assay. Figure 2 shows that addition of PFK to F16BP and ADP enabled much greater 221 production of ATP than in control samples alone. Conversion from concentrations in µg/ml to molar 222 concentrations shows that human and trypanosomatid PFKs produce similar amounts of ATP per 223 mole. The yield of ATP from control experiments may derive from spontaneous conversion of ADP 224 into ATP occurring at high ADP concentrations (2ADP \IGHT ATP + AMP). Potential contamination of 225 ADP stocks with ATP was prevented by using Ultrapure ADP (>99% purity; Promega). Minimal 226 concentrations of ATP were present in PFK stocks, as shown by control experiments without F16BP 227 or ADP; amounts were insufficient to confound results (data not shown).

228

## 229 Kinetic properties for reverse activity by trypanosomatid PFKs

230 An enzyme-linked assay was used to measure F6P production from the reverse reaction catalysed by 231 PFK isoforms. Michaelis-Menten curves using allosteric sigmoidal models were generated for ADP 232 titrations (Figure 3A) and F16BP titrations (Figure 3B). TbPFK is the most active isoform, with the lowest  $K_{0.5}^{ADP}$  and  $K_{0.5}^{F16BP}$  ( $K_{0.5}$  defined as concentration of substrate at which half maximal enzyme 233 234 velocity is reached, analogous to Michaelis-Menten constant [K<sub>M</sub>]). TcPFK has similar kinetic 235 parameters to TbPFK. LmPFK has a slightly lower V<sub>max</sub> and markedly lower affinities for both 236 substrates, in keeping with known lower activities in the forward reaction (17). Full kinetic parameters 237 are listed in Table 2. Control experiments without PFK did not produce significant quantities of ATP 238 (data not shown).

239

Both TcPFK and TbPFK reverse reaction kinetics were consistent with allosteric sigmoidal models for
 F16BP titrations and for ADP titrations, with LmPFK reaction kinetics showing no statistical difference
 between the Michaelis-Menten and allosteric sigmoidal models; the allosteric sigmoidal model was
 preferred to retain consistency.

244

# 245 The reverse PFK reaction can be allosterically modulated in trypanosomatid PFKs

AMP is a known activator, and sole physiological allosteric effector, of the forward PFK reaction in trypanosomatids (15). The effect of AMP on the reverse activity of trypanosomatid PFKs was assessed using the enzyme-linked kinetic assay described above. All three trypanosomatid PFKs were activated in the presence of 0.5mM AMP, with the  $k_{cat}/K_{0.5}^{F16BP}$  value increasing two-fold for TbPFK and TcPFK, and ten-fold for LmPFK (Supplementary Figure 1 and 2). F26BP, a potent activator of the forward reaction in many eukaryotic ATP-dependent PFKs (but not trypanosomatids) and bi-directional PPi-dependent plant PFKs (18) did not have any effect at 1mM concentration.

253

A series of allosteric inhibitors deriving from a compound with anti-TbPFK activity (4) were tested for effects on the reverse PFK reaction catalysed by trypanosomatid PFKs. These compounds inhibited the reverse trypanosomatid PFK reaction with similar potencies to the forward reaction (results not shown).

258

# 259 The reverse PFK reaction in human PFK isoforms

260 Kinetic parameters for human PFK isoforms were determined using the enzyme-linked kinetic assay. 261 Michaelis-Menten curves using allosteric sigmoidal models were generated for ADP titrations (Figure 262 4A) and F16BP titrations (Figure 4B). PFK-M and PFK-L have similar V<sub>max</sub> values, but PFK-P is much 263 less active. However, PFK-L has much lower affinities for ADP and F16BP than the other isoforms, with the highest  $K_{0.5}^{ADP}$  and  $K_{0.5}^{F16BP}$  ( $K_{0.5}^{ADP}$  for PFK-P not determined). Kinetic parameters are listed 264 265 in Table 3. Human PFK isoforms were purified in the presence of 1mM ATP, likely degrading to ADP 266 over time; human PFK stocks thus contributed 10-40µM extra nucleotide (ATP or ADP) to the assay. 267 Each reaction was therefore normalised against a control reaction that included PFK isoform sample 268 but not exogenous substrate.

269

All reverse reaction kinetics are consistent with allosteric sigmoidal models both for F16BP titrations and ADP titrations; PFK-M ADP titrations did not show any statistically significant difference between the two models so the allosteric sigmoidal model was preferred to retain consistency. The reaction obeys allosteric sigmoidal models at ADP concentrations up to 2.5mM.

274

275 PFK-L has low affinity for ADP and F16BP, indicating a reduced propensity for the reverse reaction 276 compared to other isoforms. The sequence identities between the three isoforms range from 68 to 277 71%, with the ATP binding site being 85-90% identical (the F6P binding site has not been fully 278 characterized as yet). It is challenging to interpret these differences in reverse kinetic properties given 279 the lack of precise information about isoform specific tissue locations, with much of the original 280 published data being measured from relatively crude tissue extracts. However, assuming the original 281 conclusions reached in the 1970-80s (19) are broadly accurate in stating that PFK-L is highly 282 expressed in liver tissues (hence the name: PFK-Liver) then the relative inability of PFK-L to catalyse 283 the reverse reaction may relate to the highly gluconeogenic - and FBPase rich - environment in the 284 human liver. The relatively low activity of PFK-P was also observed in the forward reaction (not 285 shown) and may be an intrinsic property resulting from its susceptibility to time and concentration 286 dependent inactivation probably caused by dissociation of the active tetrameric form (20),

287

# 288 Substrate inhibition of the reverse PFK reaction in human PFK isoforms

There is an inhibitory effect at higher concentrations of ADP (above 2mM) for PFK-M, but the effect is not as clearly demonstrated for the other isoforms (Figure 4A). This effect can be seen more clearly when higher concentrations of ADP are investigated (Figure 5). A similar effect is observed at higher concentrations of F16BP (above 5mM) for PFK-M (Figure 4B). Standard Michaelis-Menten or allosteric sigmoidal kinetic models should be used with caution when substrate inhibition is present; however, more complex models did not increase accuracy of data fitting. It is likely that substrate inhibition of the reverse PFK reaction is also present in PFK-L and PFK-P at higher substrate

- 296 concentrations than were used experimentally but to varying degrees, in a similar way that ATP-
- 297 dependent inhibition differentially affects the forward reaction in each isoform (21). The rationale for
- substrate inhibition of the reverse reaction remains uncertain. It may be analogous to substrate
- inhibition by ATP in the forward reaction, which enables finer control of the glycolytic flux (22), though
- 300 it would be surprising if such an effect had been evolutionary advantageous given the presumed small
- effect on fitness of precise control of the reverse reaction. High ADP/ATP ratios may enable sufficient
   ATP production to allow the forward reaction to occur simultaneously; this possibility was reduced –
- 303 but necessarily eliminated by adding an excess of glycerol kinase in the presence of glycerol.

- 304 **DISCUSSION**
- 305

306 The experiments presented in this paper provide the first kinetic data on the reverse PFK reaction by 307 eukaryotic ATP-dependent phosphofructokinases. The basis for the doctrine that PFK acts only in a 308 forward direction derives from  $\Delta G^{\circ}$  being highly favourable for the forward reaction.

309

310 [1] F6P + ATP = F1,6BP + ADP

311

 $\Delta G^{\circ} = -3.4 \text{ kcal.mol}^{-1}$  as calculated from the free energies of formation for the products under standard conditions and 1M concentrations (23). The free energy of the reaction under different cellular conditions can be estimated from [2], where the reaction quotient Q is the ratio of concentrations of available products and reactants [3].

316

317 [2]  $\Delta G = \Delta G^{\circ} + RT \ln Q$ 

- 318 [3] Q = [ADP].[F16BP]/[ATP].[F6P]
- 319

320 Estimated cellular concentrations of reactants and products for human and trypanosomes are given in 321 Table 4 and can be used to calculate the reaction quotients (Q), the free energy ( $\Delta G$ ), and the 322 equilibrium constant (K), for the PFK reaction in each of the tissue types. For the reverse PFK 323 reaction [1] to take place,  $\Delta G$  needs to be greater than 0: ( $\Delta G^{\circ}$  + RT ln Q > 0). Substituting 324 appropriate values ( $\Delta G^{\circ} = -3.4 \text{ kcal.mol}^{-1}$  and RT= 0.543 kcal.mol<sup>-1</sup>) we can solve for reaction 325 quotient Q ([ADP].[F16BP]/[ATP].[F6P]) to show that Q must be greater than 500. In other words, a 326 net reverse reaction will only be energetically favourable when the relative concentration of products 327 (F16BP & ADP) is at least 500-fold greater than that of the substrates (F6P & ATP).

328

329 As shown in Table 4, the concentrations of substrates and products measured in different cell types 330 give estimated reaction quotients of 0.07-0.85 in human tissues and 0.4 in trypanosomatid 331 glycosomes. Additional experimental measurements of muscle from various species, including 332 insects, fish and mammals give ATP/ADP ratios ranging from 11:1 to 1:1 and an average reaction 333 quotient (Q) of 0.42 for their PFKs (24). These Q low values show that in most organisms PFK is 334 working far from the reaction's equilibrium. Further experimental studies using mass spectrometry to 335 estimate forward and reverse flux of each step in glycolysis in yeast, E. coli and a mammalian kidney 336 cell line (25) identified PFK as an almost exclusively forward driven glycolytic step in all cells. The 337 measured  $\Delta G = -3.2 \text{ kcal.mol}^{-1}$  for PFK in the kidney cell estimated the reverse flux of PFK to be less 338 than 0.7% of the forward flux, with the authors concluding that (under steady state conditions with 339 cells grown in high glucose media) "phosphofructokinase functions as a classic irreversible step". Our data show that for human cells, the measured  $K_{0.5}^{ADP}$  values for the reverse reaction (Table 3) 340 341 exceeds the cytosolic concentrations of ADP (Table 4), but as steady-state concentrations of F16BP are lower than the  $K_{0.5}^{F16BP}$  values, it seems likely that under steady state conditions the reverse 342 343 reaction would indeed be inefficient. However cells can be subjected to extreme conditions of stress

and nutrient deprivation and the PFK reverse flux may be very different in non-steady state, glucose poor conditions and Q values of 500 or above could easily be achieved if ATP or G6P were
 (transiently) depleted.

347

348 Despite the apparently high energetic barrier, the reverse reaction for ATP-dependent PFK is possible 349 in cells under certain conditions. Experimentally, the reverse reaction was first shown in vitro in E. coli PFK at high ADP and F16BP concentrations (26). Kinetic parameters were determined as K<sub>M</sub><sup>F16BP</sup> 350 of 398 $\mu$ M at 2mM ADP and K<sub>M</sub><sup>ADP</sup> 50 $\mu$ M at saturating concentrations of F16BP. The kinetic 351 352 parameters suggest generally tighter F16BP and ADP binding compared with the PFK isoforms 353 presented here (Tables 2 and 3) but are not far removed from the kinetic values for PFK-M and this 354 leaves open the question whether there are ever cellular conditions when the reverse reaction could 355 occur.

356

357 Metabolite concentrations vary widely between individuals (27) and these differences will be 358 exaggerated further under conditions of metabolic stress. Low concentrations of aldehyde severely 359 deplete ATP levels (28); cells infected with viruses (29) and necrotic cancer cells (30) have also 360 shown large variations in ADP:ATP ratios. Cells also undergo programmed responses when adapting 361 to new states invoked by events including cell division, apoptosis or nutritional stress which involve 362 concerted changes in activities for families of enzymes (so called 'allostatic changes') (31). The 363 transition between these states frequently necessitates large swings in metabolite concentrations. 364 One such example is the transition from glycolysis to gluconeogenesis triggered by large changes in 365 the ATP:ADP ratio. There will be a lag period whilst waiting for upregulation of FBPase (the 366 gluconeogenic protein that carries out the reverse PFK reaction but not coupled to ATP formation) by 367 increasing transcription/translation or by post-translational modification. During this lag period we 368 postulate that substrate concentrations may be sufficiently perturbed to permit PFK reversal, before 369 FBPase activity becomes available. The potential role of the PFK filament assemblies (35) and the 370 existence of the as yet poorly characterised glycosome complex (36) or G-bodies (32) that tunnel 371 substrates with potentially very high effective concentrations also provides a mechanism for achieving 372 non-equilibrium concentrations of substrates sufficient to push PFK into reverse.

373

374 Trypanosomatids have a uniquely interesting way of organising the glycolytic enzymes with the first 375 seven enzymes in the pathway (including PFK) sequestered in glycosomes (33). In the bloodstream 376 form of T. brucei, there can be over 60 of these organelles per cell, comprising about 5% of the cell 377 volume (34). Metabolism in the T. brucei parasite has been extensively studied and kinetic 378 parameters for the glycolytic enzymes have been incorporated into a sophisticated in silico, 379 experimentally validated metabolic model (35,36). Despite the good predictive properties of such 380 models, it remains difficult to determine the concentrations of individual metabolites. A potential 381 complication in parameterising in silico models is caused by compartmentalisation in eukaryotic cells, 382 as metabolite concentrations may vary significantly between different types of vesicles.

383

384 An experimental indication that the reverse PFK reaction may be physiologically relevant and play a role in gluconeogenesis in trypanosomes comes from <sup>13</sup>C labelled glucose LC-MS work in 385 386 bloodstream forms of T. brucei (37). This showed that hexose phosphates can indeed be derived 387 (albeit at a low rate of 2%) from F16BP despite no active form of FBPase being detectable in 388 glycosomes thus possibly indicating reversal of the normal PFK reaction direction. Further 389 biochemical studies suggest that gluconeogenesis occurs after complete knock-out of FBPase 390 (F.Bringaud, personal communication) and leaves open the possibility that the reverse PFK reaction 391 may contribute to this pathway. Gluconeogenesis is also carried out in *Leishmania*, but recent studies 392 (38) showed that Leishmania amastigotes in activated macrophages cannot use amino acids, instead 393 relying on glycolysis. It is unlikely that FBPase is active in amastigotes in vivo, because as this 394 would lead to ATP loss by futile cycling. A likely scenario is that these leishmanial forms have an 395 inactivated form of FBPase (e.g. by posttranslational modification), and like the bloodstream-form T. 396 brucei in the presence of abundant glucose, PFK may function in reverse. As for mammalian PFK, 397 the estimated physiological reaction quotient of 0.4 (Table 4) is far below the required reaction 398 quotient (500) for the reverse direction. Nevertheless, measuring accurate intra-glycosomal substrate 399 concentrations is technically difficult, meaning that the in vivo reaction quotient inside glycosomes 400 may be significantly different from whole cell data. An analogous situation occurs with the T. brucei 401 glycosomal glycerol kinase reaction, for which it is known that reversal occurs in vivo with a low 402 ATP/ADP ratio in the organelles (as created under anaerobic conditions) in the presence of glycerol 403 3-phosphate despite the  $\Delta G^{\circ}$  being even less favourable than for PFK reversal (39). Furthermore, 404 mutagenesis experiments revealed structural optimisation of this enzyme for catalysis of the reverse 405 reaction (40).

406

407 Our work shows that LmPFK has significantly lower affinities for ADP and F16BP in the reverse 408 reaction compared to the other trypanosomatid PFKs, though it is activated much more by AMP. The 409 rationale for the differences in trypanosomatid PFK kinetic properties may derive from the differing 410 nutritional environments to which each parasite has become adapted. T. brucei is exclusively extra-411 cellular, usually confined to the haemolymphatic circulation and cerebrospinal fluid, whereas T. cruzi 412 is both (transiently) bloodstream and intracellular (cytosolic), infecting a wide variety of cells. 413 Leishmania parasites are predominantly restricted to macrophage phagolysosomes, where they may 414 become metabolically quiescent resulting in low growth rates (41). In this energy restricted 415 environment, the low-energy signal of rising AMP levels may be more important in stimulating 416 glycolytic or gluconeogenic activity, accounting for the greater sensitivity of LmPFK to AMP.

417

The kinetic data for the eukaryotic ATP-dependent PFKs presented in this paper help provide a more detailed understanding of the controls governing the glycolytic and gluconeogenic pathways. They will

420 also provide useful experimental data to feed into the increasingly detailed computational models

- 421 describing these pathways in mammals (42) and trypanosomes (43). The high reaction quotient (Q)
- 422 required for the reverse reaction can in principle be attained when ATP has been depleted (< 100µM)
- 423 in the presence of physiologically relevant cellular concentrations of ADP, F6P and F16BP. However,

- 424 for the human isoforms, the measured  $K_{0.5}$  values for F16BP are higher than the measured cellular 425 concentrations. To suggest any physiological relevance for the human isoforms would require the 426 existence of an 'apparent concentration' of F16BP up to ten-fold higher than has been measured; this 427 is potentially achievable by invoking substrate tunnelling or metabolon structures. The kinetic data 428 measured for the trypanosomatid PFKs would (as for the mammalian PFKs) require a significant 429 increase in ADP/ATP ratio for the reverse reaction, though the cellular concentration of F16BP would 430 (unlike the mammalian case) be above the  $K_{0.5}$  and sufficient to drive the reaction in the opposite 431 direction. Future more detailed metabolomics studies will continue to deliver more precise data on
- 432 time-dependent and organelle-dependent metabolite concentrations which will shed more light on the
- 433 potential physiological relevance of the reverse PFK reaction.
- 434

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- 440

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# TABLES

	DNA source	Expression	Expression	Antibiotic	<u>Media</u>
	<u>strain</u>	<u>plasmid</u>	<u>cell line</u>		
<u>T. b. brucei</u>	Lister 427	pET28a	C41 (DE3)	Kanamycin	2xYT broth
<u>T. cruzi</u>	CL Brener	pET28a	BL21 (DE3)	Kanamycin	Superbroth
L. infantum	JPCM5	pDEST17	C41 (DE3)	Carbenicillin	LB broth

Table 1 Expression conditions for trypanosomatid PFKs in E. coli

Table 2 Kinetic parameters for trypanosomatid PFKs in the reverse reaction. Values are mean averages (+/- standard error of mean; n = 3). Large standard errors for TcPFK data result from varying degree of fit of non-linear regression sigmoidal models, despite narrow error bars apparent in Figure 4.

	V <sub>max</sub> (μmoles/min.mg)	K <sub>0.5</sub> <sup>ADP</sup> (μΜ)	К <sub>0.5</sub> <sup>F16BP</sup> (µМ)	h (ADP)	h (F16BP)
TcPFK	4.77 (1.37)	1768 (2238)	1540 (153)	0.51 (0.14)	1.72 (0.22)
TbPFK	4.22 (0.28)	1382 (275)	1287 (202)	0.83 (0.07)	0.86 (0.06)
LmPFK	3.79 (0.69)	3137 (1084)	2495 (624)	1.42 (0.37)	1.71 (0.65)

Table 3 PFK-L has the lowest affinity for F16BP in the reverse reaction. (Mean average values with SEM values in brackets; n = 3).

	V <sub>max</sub> (µmoles/min.mg)	K <sub>0.5</sub> <sup>ADP</sup> (μΜ)	K <sub>0.5</sub> <sup>F16BP</sup> (μΜ)	h (ADP)	h (F16BP)
PFK-M	0.63 (0.05)	66.8 (13.4)	804 (66.1)	0.71 (0.22)	1.83 (0.23)
PFK-L	0.62 (0.01)	344.9 (36.4)	2100 (78.4)	1.39 (0.17)	2.49 (0.19)
PFK-P	0.22 (0)	ND	717.8 (58.1)	ND	1.35 (0.12)

Table 4: Concentrations of substrates and products of PFK in various mammalian tissues.  $\Delta G$  calculated from equation (2) using RT = 0.543 kcal.mol<sup>-1</sup> where R is the gas constant (1.987 cal.mol<sup>-1</sup>K<sup>-1</sup>) and T = temperature (293K). Equilibrium constant K is calculated from K = exp (- $\Delta G/RT$ ). References for concentrations (directly measured in non-italicised text, derived from models based on whole cell lysate data in italicised text): <sup>a</sup> (Minakami and Yoshikawa 1966); <sup>b</sup> (Beis and Newsholme 1975); <sup>c</sup> (Zalitis and Oliver 1967); <sup>d</sup> (Lowry and Passonneau 1964); <sup>e</sup> (Spolter, Adelman, and Weinhouse 1965); <sup>f</sup> (Rao and Oesper 1961); <sup>g</sup> (Hisanaga, Onodera, and Kogure 1986); <sup>h</sup> (Bakker, Westerhoff, and Michels 1995); <sup>i</sup> (Graven et al. 2014).

	Erythrocyte	Muscle	Brain	<i>T. brucei</i> glycosome
[ATP] μM	1850 <sup>ª</sup>	4990 <sup>b</sup>	3325 <sup>9</sup>	3870 <sup>′</sup>
[ADP] μ <b>M</b>	180 <sup>°</sup>	600 <sup>f</sup>	309 <sup>g</sup>	1315
[F6P] μM	15.7 <sup>a</sup>	110 <sup>c</sup>	27 <sup>d</sup>	2400 <sup>h</sup>
[F16BP] μM	7 <sup>a</sup>	32 <sup>e</sup>	200 <sup>d</sup>	1900 <sup>h</sup>
Q (derived)	0.04	0.03	0.69	0.27
⊿G (kcal/mol)	-5.09	-5.21	-3.6	-4.7

### FIGURES

Figure 1. SDS-PAGE gels (4-20%) showing A) trypanosomatid PFKs were produced at high purity but removal of tags was only partially successful and B) human PFK isoforms were produced at high purity but with anomalous migration of PFK-L. Tagged enzymes were used for all experiments due to incomplete tag cleavage and reduction of enzyme activity after cleavage (data not shown), likely secondary to conditions required to remove tag. Ladder markers are in kDa.



Figure 2. ATP is produced by PFKs from ADP and F16BP using an endpoint assay (ADP 5mM, F16BP 5mM, error bars are standard deviations; n = 2).



Figure 3.

A) Trypanosomatid PFKs have different kinetic responses for ADP titrations (F16BP 5mM).
B) Trypanosomatid PFKs have different kinetic responses for F16BP titrations (ADP 3mM).
(error bars are standard deviations, n = 3).



Figure 4.

A) Human PFK isoforms have different kinetic responses for ADP titrations (F16BP 5mM)
B) Human PFK isoforms have different kinetic responses for F16BP titrations (ADP 3mM)
PFK-P data are missing for 4A due to lack of enzyme stability under these assay conditions. (Error bars are standard deviations, n = 3, isoforms were individually normalised to highest specific activity).



[F16BP]µM

Figure 5. High concentrations of ADP inhibit the reverse reaction for PFK-M. (Error bars are standard deviations; n = 3).



[ADP] µM

#### **SUPPLEMENTARY FIGURE 1**

- A) AMP activates TcPFK with respect to ADP titrations (F16BP 10mM). 0mM AMP (solid triangles ▲ with solid line), 0.5mM AMP (open triangles △ with dotted line)
- B) AMP activates TbPFK with respect to ADP titrations (F16BP 10mM). 0mM AMP (solid squares with solid line), 0.5mM AMP (open squares □ with dotted line)
- C) AMP activates LmPFK with respect to ADP titrations (F16BP 20mM). 0mM AMP (solid circles
   with solid line), 0.5mM AMP (open circles with dotted line)

(n= 2, error bars are standard deviations, results for each enzyme normalised to highest specific activity without AMP present).





#### SUPPLEMENTARY FIGURE 2

- A) AMP activates TcPFK with respect to F16BP titrations (ADP 5mM). 0mM AMP (solid triangles
   ▲ with solid line), 0.5mM AMP (open triangles △ with dotted line)
- B) AMP activates TbPFK with respect to F16BP titrations (ADP 5mM). 0mM AMP (solid squares with solid line), 0.5mM AMP (open squares □ with dotted line)
- C) AMP activates LmPFK with respect to F16BP titrations (ADP 10mM). 0mM AMP (solid circles● with solid line), 0.5mM AMP (open circles with dotted line)

(n= 2, error bars are standard deviations, results for each enzyme normalised to highest specific activity without AMP present).



