Intracellular Positioning of Isoforms Explains an Unusually Large Adenylate Kinase Gene Family in the Parasite Trypanosoma brucei*

Received for publication, December 8, 2004, and in revised form, January 14, 2005 Published, JBC Papers in Press, January 18, 2005, DOI 10.1074/jbc.M413821200

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Adenylate kinases occur classically as cytoplasmic and mitochondrial enzymes, but the expression of seven adenylate kinases in the flagellated protozoan parasite Trypanosoma brucei (order, Kinetoplastida; family, Trypanosomatidae) easily exceeds the number of isoforms previously observed within a single cell and raises questions as to their location and function. We show that a requirement to target adenylate kinase into glycosomes, which are unique kinetoplastid-specific microbodies of the peroxosome class in which many reactions of carbohydrate metabolism are compartmentalized, and two different flagellar structures as well as cytoplasm and mitochondrion explains the expansion of this gene family in this parasite. The three isoforms that are selectively built into either the flagellar axoneme or the extra-axonemal parflagellar rod, which is essential for motility, all contain long N-terminal extensions. Biochemical analysis of the only short form trypanosome adenylate kinase revealed that this enzyme catalyzes phosphotransfer of γ-phosphate from ATP to AMP, CMP, and UMP acceptors; its high activity and specificity toward CMP is likely to reflect an adaptation to very low intracellular cytidine nucleotide pools. Analysis of some of the phosphotransfer network using RNA interference suggests considerable complexity within the homeostasis of cellular energetics. The anchoring of specific adenylate kinases within two distinct flagellar structures provides a paradigm for metabolic organization and efficiency in other flagellates.

The continuous production of ATP, its delivery from intracellular sites of synthesis to sites of energy consumption, and the homeostasis of adenine nucleotide pools are fundamental to cellular viability. Adenylate kinase (ADK) and phosphagen kinases, such as creatine kinase and arginine kinase, assume pivotal roles in these fundamental processes (1). Phosphagen kinases catalyze reversible phosphotransfer between ATP and the guanidine acceptor (e.g. arginine or creatine), with the phosphagen providing both a temporal and a spatial reserve of high energy phosphate. ADK, on the other hand, catalyzes the transfer of the γ-phosphate group from either ATP or GTP to the phosphoryl acceptor AMP.

Disruption of the single copy ADK gene in Escherichia coli or Schizosaccharomyces pombe is lethal (2, 3), but in Saccharomyces cerevisiae the ADK null phenotype reveals a compensating metabolic plasticity. S. cerevisiae survives because the broad substrate specificity of its uridylate kinase (Ura6p) compensates for the deficiency in cytoplasmic ADK activity (4). However, a small proportion of the yeast ADK (Aky2p) is localized in the mitochondrial intermembrane space (5), where it may contribute to efficient translocation of ATP from the mitochondrial matrix into the cytoplasm (1). Uridylate kinase is not present in yeast mitochondria, and hence AKY2-deficient S. cerevisiae exhibit a petite phenotype and are unable to grow under nonfermentative conditions (6). In skeletal muscle of transgenic AKT+/− knock-out mice, metabolic reprogramming and ultrastructural reorganization have been observed (7, 8). Here, glycolytic flux, creatine kinase activity, and mitochondrial volume are increased to support muscle function, albeit with reduced energy economy and efficiency under stress (7–9). The focus of this study is the unflagellated protozoan parasite Trypanosoma brucei, the causal agent of human African sleeping sickness (see, on the World Wide Web, www.who.int/tdr/diseases/tryp/). There are many interesting and unusual aspects to trypanosome metabolism (10). A single aerobically functioning mitochondrion and microbodies of the peroxisome class are present. However, in the mammalian host, the mitochondrial energy metabolism of the parasite is largely re-
pressed; neither cytochromes nor tricarboxylic acid cycle are present. Glycolysis is the sole energy-generating pathway, and pyruvate is excreted as the major end product of this glucose catabolism. Uniquely, the first seven glycolytic enzymes occur solely in peroxisomes. Peroxisomal targeting of glycolytic enzymes and certain other enzymes utilized for carbohydrate metabolism (10) is characteristic of all trypanosomatids and reflects the description of these trypanosomatid microorganisms as glycosomes (10, 11).

*T. brucei* is transmitted between mammalian hosts by its insect vector (*Glossina* sp., tsetse fly). Procyelic trypanosomes (found in the tsetse fly midgut) have an aerobic metabolism and use a variety of carbon sources (12–14), and alternative oxidase and cytochrome c oxidase constitute the two terminal enzymes of an essential branched respiratory chain (14, 15). Although oxidative phosphorylation is an important facet of procyelic metabolism, pyruvate kinase and an acetate:succinate CoA-transferase/succinyl-CoA synthetase cycle catalyze important substrate phosphorylations. The cytoplasmic phosphotransfer reaction catalyzed by the former enzyme appears to be essential, except perhaps under conditions where increased substrate availability can result in increased mitochondrial amino acid (proline) metabolism (12, 13, 16). Acetate:succinate CoA transferase is restricted in its evolutionary distribution to trypanosomatid mitochondria, anaerobic helminth mitochondria, and protist hydrogenosomes; it catalyzes formation of acetate from acetyl-CoA. This reaction, rather than entry into the tricarboxylic acid cycle, is the predominant pathway used for acetyl-CoA metabolism; acetate is excreted from the cell as a major end product of metabolism, and the succinyl-CoA product is recycled by mitochondrial succinyl-CoA synthetase, generating ATP (15). Deletion of acetate:succinate CoA-transferase initially results in a severe growth defect, before compensatory metabolic rearrangements allow cells to adapt to loss of the enzyme (17).

Here, we report that the complex stage-regulated metabolism of *T. brucei* is buttressed by the expression of seven ADK-like genes. The expression of seven ADKs within the context of a single cell has not been reported previously, but in this protozoan parasite it reflects the requirement to target ADK to glycosomes and two distinct flagellar structures as well as more classical intracellular sites such as cytoplasm and mitochondrion. Molecular interrogation of the ADK gene family, while demonstrating metabolic adaptation, also indicates that there is likely to be considerable complexity within the relationship between phosphotransfer activity and homeostasis of cellular energetics.

**MATERIALS AND METHODS**

**Reagents**—All chemicals were purchased from Sigma unless otherwise stated. Restriction enzymes and DNA-modifying enzymes were purchased from Roche Applied Science.

**Cells**—Procyelic *T. brucei*, stock 247, was maintained at 27°C in SDM-79 medium supplemented with 10% (v/v) heat-inactivated fetal calf serum as described previously (18). RNA interference (RNAi) experiments were conducted using the procyelic 29-13 cell line, which has been genetically modified in order to express the tetracycline repressor protein and bacteriophage T7 RNA polymerase (19). Stable integration of ADKA, ADKB, and ADKE, each containing a single Ty1 epitope at their respective C termini, the plasmids pTAFADKA::Ty, pTAFADKB::Ty, and pTubADKE::Ty were constructed. For the expression of the GFP:ADKD fusion protein, pVsg4GFP:ADKD was constructed. This fusion protein was expressed under the control of an RNA polymerase I-transcribed EPI promoter following stable integration of pVsg4GFP:ADKD into a minichromosomal locus. For expression of recombiant ADK in *E. coli*, pThAKGr was constructed.

**Generation of ADKC**—**Null Mutants**—PCR amplification of hygromycin and bleomycin resistance markers was achieved using primer combinations ADKck01 and ADKck02 (template pGad8-tubulin (24)) or ADKck03 and ADKck04 (template pRM481 (25)), respectively (see Supplemental Table I for primer sequences). The 5′ region of each primer contained ~50 bp of homology to the ADK locus, facilitating gene disruption by homologous recombination; PCR products (3 μg) were used for transformation of procyelic 427 *T. brucei*. For second round transformation of hygromycin-resistant ADKC− heteroygotes, the bleomycin resistance marker was amplified from a first round heterozygote transformant using a primer combination of ADKck05 and ADKck06. The resulting PCR product was then used for stable integration of pVsg4GFP into the nucleus. PCR products (3 μg) and recombinant ADK were analyzed by PCR and Southern hybridization.

**Expression and Analysis of Recombinant ADK**—Following transformation of pThAKGr into *E. coli* XL-Blue (Stratagene), expression of recombinant ThADKG was induced by the addition of isopropyl-β-thio-galactopyranoside (80 μM). Recombinant protein was purified by Ni2+-nitrilotriacetic acid chromatography according to the manufacturer’s protocol (Qiagen). The activity of the recombinant enzyme was measured as described previously by following the rate of NDF formation from NTP and NMP substrates in a coupled system that required pyruvate kinase and lactate dehydrogenase (26). The standard reaction mixture contained Tris-HCl (0.1 M, pH 7.6), KCl (80 mM), MgCl2 (1.4 mM), phosphoenolpyruvate (400 μM), NADH (200 μM), pyruvate kinase (500 U/ml), lactate dehydrogenase (10 U/ml) and NTP (1.8 mM), NMP (2.3 mM), and recombinant ADKG (85.8 μM). One unit of enzyme activity was defined as 1 μmol of NAD+ formed/min.

**Immunoblotting and Immunofluorescence**—Polyclonal antiserum against recombinant ADKG was prepared by Eurogentech. Polyclonal rabbit antibodies raised against *T. brucei* aldolase were used as described previously (27). SDS-PAGE and immunoblotting were carried out according to standard methods (21). Immunofluorescence of formaldehyde- and methanol-fixed detergent-extracted cytoskeletons using BB2 and ROD-1 (28) monoclonal antibodies was performed as described previously (29, 30).

**RESULTS**

**Seven ADK-like Genes Are Expressed in T. brucei**—Using NMPPK genes characterized from *S. cerevisiae*, *S. pombe*, *H. sapiens*, *E. coli*, and *B. subtilis* as query sequences, we analyzed the *T. brucei* genome data bases at Sanger and the Institute for Genome Research for the presence of NMPPK genes. We found open reading frames homologous to the guanulate kinase and thymidylate kinase from *S. cerevisiae* and *H. sapiens* (31–33), but no hits to cytidylate and uridylate kinase family genes that are unique to bacteria (34–36) (data not shown). We also found a *T. brucei* homologue of a very recently discovered distinct class of ADK genes that is conserved in eukaryotes and Archaea; in *S. cerevisiae* and humans, this novel ADK family member is found in the nucleus (37). How-
ever, through a combination of database searching and PCR cloning, we identified seven different sequences that shared homology with the cytoplasmic and mitochondrial ADK genes from yeast and humans (Fig. 1). Now that the *T. brucei* genome project is nearing completion, all of the open reading frames that we identified can now be viewed at the Sanger Centre Web site (www.genedb.org/genedb/tryp/index.jsp). With the exception of the neighboring ADKC and ADKF, these genes are not clustered in the genome. We also demonstrated that each of the ADK genes was transcribed into mRNA using real time PCR to monitor the incorporation of SyberGreen into specific ampli-
cons. This analysis revealed that each gene was expressed in both procyclic and long slender bloodstream trypanosomes. We hypothesized that the expression of seven different ADK genes

![Sequence alignment of seven adenylate kinases from *T. brucei*. Sequences were initially aligned in Clustal, and then the alignment was adjusted manually. The asterisks and dots represent residues that are identical or conserved in a majority (six) of the sequences, respectively. The characterized *S. cerevisiae* enzymes Aky2p (accession number M18455), Pak3p (accession number X65126), and Ura6p (accession number M31455) were also included in the alignment. The highly conserved ATP-binding P-loop is boxed, and the length of the LID domain is indicated by the dashed line. Other residues known to be important either in substrate binding or catalysis are highlighted by red and yellow shading, respectively. The blue shading reveals two conserved hydrophobic residues that form part of the essential C-terminal amphipathic $\alpha$-helix, which makes hydrophobic contact with the purine ring of ATP.

**FIG. 1.** Sequence alignment of seven adenylate kinases from *T. brucei*. Sequences were initially aligned in Clustal, and then the alignment was adjusted manually. The asterisks and dots represent residues that are identical or conserved in a majority (six) of the sequences, respectively. The characterized *S. cerevisiae* enzymes Aky2p (accession number M18455), Pak3p (accession number X65126), and Ura6p (accession number M31455) were also included in the alignment. The highly conserved ATP-binding P-loop is boxed, and the length of the LID domain is indicated by the dashed line. Other residues known to be important either in substrate binding or catalysis are highlighted by red and yellow shading, respectively. The blue shading reveals two conserved hydrophobic residues that form part of the essential C-terminal amphipathic $\alpha$-helix, which makes hydrophobic contact with the purine ring of ATP.
The Short Form "Cryptic Kinase" Has Broad Substrate Specificity—TbADKG encodes the only trypanosome short form ADK; the distinction between long and short ADK isoforms referring to the length of the Lid domain (38). The Lid contributes to the overall folding of NMPPks and changes conformation following the binding of substrate to the enzyme. In mammalian cells, the major cytoplasmic form of ADK is a short form enzyme, and mitochondrial activity is associated with longer ADK isoforms (38). In yeast, cytoplasmic and mitochondrial ADK activities are provided by long form ADK enzymes (3, 39). Eukaryotic UMP/CMP kinases share primary sequence homology with ADK and are always of the short kind (38).

Comparison of the trypanosome ADK family with homologues reflects a molecular radiation to fill particular metabolic niches. In this regard, they resemble the cytoplasmic and mitochondrial ADK isoforms of other eukaryotes. By contrast, ADKF possess no unusual or structurally distinctive characteristics. In this regard, they resemble the cytoplasmic and mitochondrial ADK isoforms of other eukaryotes. From other organisms indicated that T. brucei ADKC and ADKF possess no unusual or structurally distinctive characteristics. In this regard, they resemble the cytoplasmic and mitochondrial ADK isoforms of other eukaryotes. By contrast, our attention was focused by unusual primary sequence characters in each of the remaining five T. brucei enzymes.

The Substrate Specificity of T. brucei ADKG

ADKG was expressed in S. cerevisiae (Fig. 1), proteins Aky2p and Ura6p (Fig. 1), confirmed using antibodies against known stage-regulated proteins, as described previously (21). To probe the subcellular localization of the three trypanosome ADKs with unusually long leader sequences upstream of the conserved P-loop, we engineered TbADKA to encode proteins that were tagged at their respective C termini with a single Ty epitope. Recombinant genes were then integrated in this manner into the genome. Expression of the Ty-tagged proteins was determined by immunoblotting and immunofluorescence using the anti-Ty BB2 antibody (29). Immunofluorescence images for ADKA:Ty and ADKB:Ty, or ADKE:Ty but not in the untransfected parental cell line, and this signal indicated that each protein was present in the trypanosome flagellum. Important, flagellum-associated antibody staining was retained in detergent-extracted trypanosomes (Fig. 3, A–C), indicating that each protein was incorporated into a flagellar cytoskeletal structure. In Fig. 3, A and B, the images shown are of cells double-labeled with the anti-tag.
was seen only in the case of with the growth rates of noninduced controls, reduced growth double-stranded RNA-mediated decay process. By comparison and ADKA either post-induction the growth of cell lines in which mRNA for tion approaches require. Thus, we tracked for up to 14 days sequences of rapidly silencing the expression of each ADK gene in tetracycline-inducible RNAi system to investigate the conse-

localization of GFP fluorescence with the glycosomal marker

its N terminus to green fluorescent protein (GFP), and co-
localization of ADKD through the expression of ADKD, fused at
and exhibits a pI of 10.1. We demonstrated the glycosomal
ADKD gene that we considered likely to encode a glycosomal ADK was
ture of many glycosomal matrix proteins is a high pI. The only
N-terminal region of the targeted protein. An additional fea-
ture of many glycosomal matrix proteins is a high pl. The only
gene that we considered likely to encode a glycosomal ADK was
ADKD; the predicted protein contains a C-terminal -SKL motif and exhibits a pl of 10.1. We demonstrated the glycosomal
localization of ADKD through the expression of ADKD, fused at
its N terminus to green fluorescent protein (GFP), and co-
localization of GFP fluorescence with the glycosomal marker
aldolase.

**Identification of the Glycosomal ADK**—In trypanosomes, ADK activity has been found in purified glycosomes (47–49). The evolutionary connectivity of glycosomes and peroxisomes extends to the conservation of targeting and import mecha-
nisms between both organelles (11). Two canonical recognition signals for the import machinery are a peroxisome targeting signal 1, which is a C-terminal tripeptide motif (typically -SKL), or a peroxisome targeting signal 2, which is a nonapep-
tide sequence (consensus -(R/K)(L/V/I)/X₅(R/K)(L/A/P)) at the N-terminal region of the targeted protein. An additional feature of many glycosomal matrix proteins is a high pl. The only
gene that we considered likely to encode a glycosomal ADK was
ADKD; the predicted protein contains a C-terminal -SKL motif and exhibits a pl of 10.1. We demonstrated the glycosomal
localization of ADKD through the expression of ADKD, fused at
its N terminus to green fluorescent protein (GFP), and co-
localization of GFP fluorescence with the glycosomal marker
aldolase.

**ADKD and ADKF Are Important for Growth**—We used a tetracycline-inducible RNAi system to investigate the conse-
quences of rapidly silencing the expression of each ADK gene in the absence of the longer selective pressure that gene disruption
approaches require. Thus, we tracked for up to 14 days post-induction the growth of cell lines in which mRNA for either ADKA, ADKB, ADKC, ADKD, ADKE, ADKF, ADKG, or ADKE and ADKF was specifically targeted for destruction by a double-stranded RNA-mediated decay process. By comparison with the growth rates of noninduced controls, reduced growth was seen only in the case of ADKD and ADKF knockdowns (Fig. 4). Ultrastructural analysis using electron microscopy indi-
cated that the slow growth phenotype resulting from RNAi against the glycososomal ADK was not accompanied by any ob-
vious change in glycosome morphology or number (data not shown).

Western blot analysis of RNAi cell lines for ADKG, encoding a presumably essential UMP/CMP kinase activity, indicated in at least one instance that the absence of phenotype may result from obtaining only a moderate knockdown of protein levels (Supplemental Fig. 1). However, in other cell lines producing no discernable phenotype, an assay of ADK activity in cytoske-

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**TABLE II**  
Kinetic properties of *T. brucei* ADK

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Substrate(s)</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Kₘ (µM)</td>
<td>ATP</td>
<td>0.18</td>
</tr>
<tr>
<td></td>
<td>AMP</td>
<td>3.2</td>
</tr>
<tr>
<td></td>
<td>CMP</td>
<td>0.17</td>
</tr>
<tr>
<td>Vₘₕ (units/mg)</td>
<td>AMP/ATP</td>
<td>77</td>
</tr>
<tr>
<td></td>
<td>CMP/ATP</td>
<td>78</td>
</tr>
</tbody>
</table>

* For Kₘ (ATP), the concentration of AMP was fixed at 2.3 mM.
* For Kₘ (NMP), the concentration of AMP was fixed at 0.18 mM.

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antibody BB2 and antibody ROD-1, which detects the paraflagellar rod (PFR), an extra-axonemal structure that runs alongside the classical “9 + 2” axoneme from the point where the flagellum exits the flagellar pocket to its distal tip. Co-
localization of the BB2 and ROD-1 antibody patterns indicated, therefore, that ADKA and ADKB are not associated with the axoneme but rather with the PFR. On the other hand, BB2-
detected ADKE::Ty did not co-localize with ROD-1 antibody (not shown), and the immunofluorescence extended to a posi-
tion close to the kinetoplast (Fig. 3C). The kinetoplast is phys-
ically attached to the flagellar basal body from which the ax-
oneme extends (46); our immunofluorescence data were therefore consistent with the assembly of an ADK isofrom
within the trypanosome axoneme. A spectrophotometric en-
zeyme assay confirmed the presence of ADK activity in both cytoskeletal fractions and purified flagella (45).

**Fig. 3. Localization of ADKA, ADKB, and ADKE to the trypanos-
me flagellum and ADKD to glycosomes. A and B, detergent-
extracted cytoskeletons from cells expressing either ADKA (A) or ADKB (B) proteins tagged at the C terminus with a single Ty epitope were used in double-labeling immunofluorescence with the anti-Ty tag anti-
body BB2 (anti-Tag; green) and the ROD-1 antibody (anti-PFR; red), which recognizes a minor component of the extra-axonemal paraflagel-
lar rod. 4',6-Diamidino-2-phenyindole-stained nuclei and mitochondrial DNA (kinetoplast) are shown in blue (DAPI). C, immunofluorescence with antibody BB2 of a detergent-extracted cytoskeleton from a cell that was expressing ADKE tagged at the C terminus with a Ty epitope and was also elongating a new flagellum (an asterisk marks the tip of this new flagellum). A white arrowhead shows the position of one kineto-
plast, which is physically attached to a flagellar basal body, and the yellow arrowhead indicates the start of an immunofluorescence signal close to this structure. Control experiments using an unmarked procy-
celic cell line and immunofluorescence using only secondary antibodies confirmed the authenticity of ADKA, ADKB, and ADKE in the trypano-
some flagellum. D, co-localization of a GFP::ADKD fusion protein with the glycosomal marker aldolase in immunofluorescence using anti-
aldolase-specific antibodies. The merged fluorescence also includes the phase-contrast image.**
Conservation, we identified orthologues of Tb ADKs from several trypanosomatid parasite genomes. Based on sequence identity and syntenic data, these orthologues are of which succeed in environments different from that encountered by T. brucei, which uses the ADK gene family in the latter species was a common feature of trypanosomatid biology. The results of this survey are summarized in Table III. Based on sequence identity and syntenic conservation, we identified orthologues of Tb ADKs and Tb ADKs from the genomes of T. cruzi and Leishmania major. Orthologues of Tb ADKs are also present in T. cruzi, L. major, and Leishmania infantum, although intriguingly putative orthologues from both Leishmania species are characterized by a series of short deletions (Fig. 5). We also identified a putative ADK isoform that is conserved between L. major and L. infantum but absent from the Trypanosoma gene databases. Targeting of two distinct ADK isoforms to the PFR appears to be restricted to African trypanosome species (T. brucei and Trypanosoma vivax).

**Discussion**

Our characterization of the extended ADK gene family in T. brucei provides insight into an additional layer of complexity to our understanding of energy metabolism in this important protozoan parasite. Remarkably, the ADK family in trypanosomes is the largest identified so far in any organism; at least seven distinct ADKs are expressed by bloodstream and procyclic trypanosomes. An obvious question arises as to why the ADK family of trypanosomes is so large. The sequence and distribution within the genome of each ADK gene indicate that this gene family has not arisen as a consequence of recent gene duplication events. Insight into why seven distinct ADKs are needed comes from the revelation that individual isoforms are required in the many diverse cellular compartments that occur in trypanosomes, including its flagellum and the unique glycosomes. A general absence of introns within protein-coding genes, polycistronic transcription, and a lack of regulation at the level of transcript initiation preclude the use of alternative splicing or alternative transcription start sites to generate the diversity of targeting signals required to partition different products of a single ADK gene between diverse cellular compartments. In other eukaryotic microbes, the use of such molecular mechanisms to differentially target enzymes has been documented (e.g., see Ref. 51). Our attention was focused on those T. brucei ADKs with unusual structural properties.

**Substrate Specificity of ADKG**—Sequence analysis of the T. brucei ADKs indicated that six enzymes are of the so-called long variety, whereas the seventh enzyme is a short form kinase. Bioinformatic analysis placed this short form enzyme, ADKG, with UMP/CMN kinases of other organisms, but our biochemical analysis of recombinant ADKG revealed that the enzyme was significantly more active using an AMP phosphate acceptor than using UMP. However, it is not without precedent for UMP/CMP kinases to show activity toward AMP; indeed, the yeast orthologue Ura6p has been reported to be more active with an AMP phosphate acceptor than with UMP. The activity of ADKG toward CMP may be critical in maintaining CDP/CTP levels in trypanosomes, but failure to observe growth phenotypes in procytic trypanosomes, but failure to observe growth phenotypes when the expression of other ADKs were subject to RNAi reflected an inability to completely ablate gene expression.
(likely in the case of ADKG) and/or was suggestive of functional redundancy in culture. Although the notion of metabolic redundancy or flexibility is evident from various molecular interrogations of gene function in trypanosomes (e.g. see Refs. 12 and 15), the subtle metabolic restructuring that is observed in transgenic AK1/H11002/H11002 and ScCKmit/H11002/H11002/ M-CK/H11002 mice provides a specific paradigm for complex compensatory phenotypes that can arise following the loss of adenylate or creatine kinase activity, respectively (7, 8, 54). Only when tissues were placed under metabolic challenge was the importance of the ADK-catalyzed reaction for muscle function and energetic efficiency realized (7, 9, 55). The ability to observe a mutant phenotype is obviously a consequence of the interrogation conditions; any "no phenotype" statement can merely reflect an inability to apply the correct interrogation conditions.

In the case of the African trypanosome, the niches within which it has evolved (tsetse fly midgut and salivary glands, mammalian bloodstream) are, at present, difficult to interrogate with experimentally tractable cultured parasites. For instance, whereas a classical tricarboxylic acid cycle plays no role in the energy generation of cultured parasites (56), it is not known whether a complete cohort of tricarboxylic acid cycle enzymes is necessary for energy generation at any point during a natural life cycle (14). It is conceivable that maintenance of ATP homeostasis at the lowest metabolic cost could be critical for successful transmission of T. brucei through tsetse. In this context, certain ADKs could therefore assume essential roles that cannot be compensated by an increase in high energy phosphoryl flux through other metabolic pathways. It is perhaps pertinent to note that arginine kinase activity has also been detected in procyclic extracts (57). Again, we have seen that RNAi against T. brucei arginine kinase yields no discernable phenotype in culture, although neither arginine kinase activity nor protein can be detected 96 h after induction of RNAi.2

Slow growth of the ADKF RNAi knockdown indicates loss of a compartmentalized phosphotransfer activity that is not compensable, but we can only speculate as to the cellular role of this isoform. Since we might expect the broad substrate specificity of ADKG to be sufficient, at least in culture, to compensate for the loss of any additional cytoplasmic ADK activity, we suggest that slow growth does not occur because of the loss of cytoplasmic ADKF. On the other hand, mitochondrial energy generation is critical for growth of procyclic T. brucei (12, 15), and in yeast the available data indicate that export of ATP generated in the mitochondrial matrix to the cytoplasm is dependant upon a mitochondrial intermembrane space ADK activity provided by Aky2p (4, 6). ADK activity has been detected previously in T. brucei mitochondrial fractions (49).

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**TABLE III**

<table>
<thead>
<tr>
<th>Gene</th>
<th>Location or function in T. brucei</th>
<th>Homologue in T. cruzi</th>
<th>Homologue in L. major</th>
<th>Homologue in T. vivax*</th>
</tr>
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<tbody>
<tr>
<td>ADKA</td>
<td>Flagellar</td>
<td>No</td>
<td>No</td>
<td>Yes</td>
</tr>
<tr>
<td>ADKB</td>
<td>Flagellar</td>
<td>Yes</td>
<td>Yes</td>
<td>Yes</td>
</tr>
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<td>ADKC</td>
<td>Flagellar</td>
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</tr>
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<td>ADKD</td>
<td>Glycosomal</td>
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</tr>
<tr>
<td>ADKE</td>
<td>Flagellar</td>
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<td>Yes</td>
</tr>
<tr>
<td>ADKF</td>
<td>ND</td>
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<td>Yes*</td>
</tr>
<tr>
<td>ADKG</td>
<td>UMP/CMP kinase</td>
<td>Yes</td>
<td>Yes</td>
<td>LmjP25.23702d</td>
</tr>
</tbody>
</table>

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* Only the homologues of flagellar enzymes are highlighted.

* ND, not determined.

* By comparison with homologues in Trypanosoma, this protein contains deletions in two Leishmania species (Fig. 5).

* A homologue is also present in L. infantum but is absent from the Trypanosoma.

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**FIG. 5.** Primary sequence alignment of ADKE homologues from T. brucei, L. major, and L. infantum highlights deletions within the Leishmania proteins. Sequences were initially aligned in Clustal, and then the alignment was adjusted manually. The P-loop is boxed, and the important structural and catalytic residues highlighted in Fig. 1 are shown as white letters on a black background. The asterisks and dots represent residues that are identical or conserved in all three sequences.

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2 M. Ginger, C. Pereira, and K. Gull, unpublished results.
analog with our understanding of yeast biochemistry, further studies could indicate whether ADKF is required for export of mitochondrial ATP in trypanosomes.

**Phosphotransfer within the Flagellum**—In the case of the flagellar ADKs, their independent identification through a proteomic analysis of the trypanosome flagellum (45) and the detection of ADK activity in isolated flagella confirm the localization data presented in this paper. Whereas ADK activity has long been associated with purified eukaryotic flagella (e.g. see Refs. 58 and 59), the identity of bona fide flagellar proteins possessing ADK activity had not, until recently, been established. The biological function(s) of such activity remains enigmatic; ADK could contribute to the numerous signaling processes that regulate the coordination of axonemal dynein ATPases and flagellar beating, but equally ADK activity could be required for energetic efficiency within the flagellar compartment. Our discovery of an axonemal T. brucei ADK extends our previous identification of ADK in the PFR (45). A parsimonious explanation of such intriguing differential targeting within the flagellum is that ADK assumes a different role depending upon the flagellar structure to which it is targeted.

Integration or positioning within the close proximity of dynein arms is an obvious possibility for any flagellar ADK, and indeed an ODA6-dependant association of ADK with outer dynein arms in *Chlamydomonas reinhardtii* was recently reported (60). Intriguingly, however, a survey of the literature indicates that the availability of a flagellar energy-generating pathway may be conserved within flagellates and that the identity of this energy-generating pathway may be determined, at least in part, by the environment in which the flagellum beats. For instance, upon release into sea water, sea urchin spermatozoa derive energy for motility solely from mitochondrial fatty acid oxidation and oxidative phosphorylation; ATP is propagated along the axoneme to the dynein ATPases by a creatine kinase-catalyzed phosphocreatine shuttle (61, 62). On the other hand, within the microaerobic environment of the female reproductive tract, mammalian spermatozoa derive chemical energy from a glycolytic pathway that is compartmentalized through the tethering of hexokinase and glyceraldehyde-3-phosphate dehydrogenase to the fibrous sheath that surrounds the axoneme and outer dense fibers along the principal piece of the sperm tail (63). It is true that sperm are highly polarized cells containing little cytoplasm and perhaps a limited capacity for diffusional exchange of metabolites, but even in the case of a microbial flagellum, the recent characterization of a central pair multiprotein complex containing ADK and enolase subunits highlights the probability of a scaffold for assembly of an energy metabolism within *Chlamydomonas* flagella also (64). In trypanosomes, the first six or seven enzymes of glycolysis are compartmentalized within glycosomes, although we suggest that there is a theoretical possibility that cytoplasmic phosphoglycerate kinase- and pyruvate kinase-catalyzed phosphotransfer could also partition into the flagellum of procyclic trypanosomes. However, during tsetse transmission, glucose is considered to provide a relatively scarce nutrient, with demand for ATP satisfied through the mitochondrial metabolism of amino acid substrates. In this context, the distribution of two ADKs along the length of the PFR could therefore be required to maintain energy economy in much the same way as ADK maintains an essential energy economy in mouse muscle (7, 9). Are trypanosome flagellar ADKs perhaps representative of a more widespread cytoskeletal anchoring of metabolic pathways to maximize energetic efficiency in eukaryotic parasites? Interesting case studies would be flagellate protozoa *Giardia* and *Trichomonas*; both lack the capacity for oxidative phosphorylation and can rely only on substrate level phosphorylation to generate ATP.

Finally, there is the issue of how ADKs with similar N-terminal extensions are differentially targeted to either axoneme or PFR. We recently demonstrated that PFR targeting of ADKA requires its N-terminal extension (45). Although this N-terminal extension is also sufficient to confer flagellar targeting of GFP, it cannot facilitate detergent- or salt-resistant incorporation of GFP into PFR or axoneme. Anchoring of the ADKs within the trypanosome cytoskeletal architecture therefore requires additional structural elements, and it may be that such features also identify key interacting partners within either axoneme or PFR, respectively.

**Acknowledgments**—We thank Paul Michels (Brussels), Be Wieringa and his group (Nijmegen), Kevin Brindle (Cambridge), León A. Bouvier (Buenos Aires), and Bill Wickstead and Catarina Gadehla (Oxford) for useful discussions during the course of this work; Eleanor Barnwell for technical assistance; and Mike Shaw for electron microscopy. Sequence data for the *T. brucei* genome were obtained from the Institute for Genomic Research and the Sanger Institute; sequencing of the *T. brucei* genome was accomplished as part of the *Trypanosoma* Genome Network with support from NIAID, National Institutes of Health, and the Wellcome Trust.

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