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Single point mutations in ATP synthase compensate for mitochondrial genome loss in trypanosomes

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Viability of the tsetse fly-transmitted African trypanosome Trypanosoma brucei depends on maintenance and expression of its kinetoplast (kDNA), the mitochondrial genome of this parasite and a putative target for veterinary and human antitrypanosomatid drugs. However, the closely related animal pathogens T. evansi and T. equiperdum are transmitted independently of tsetse flies and survive without a functional kinetoplast for reasons that have remained unclear. Here, we provide definitive evidence that single amino acid changes in the newly encoded F1F0-ATPase subunit γ can compensate for complete physical loss of kDNA in these parasites. Our results provide insight into the molecular mechanism of compensation for kDNA loss by showing F0-independent generation of the mitochondrial membrane potential with increased dependence on the ADP:ATP carrier. Our findings also suggest that, in the pathogenic bloodstream stage of T. brucei, the huge and energetically demanding apparatus required for kDNA maintenance and expression serves the production of a single F1F0-ATPase subunit. These results have important implications for drug discovery and our understanding of the evolution of these parasites.

dourine | surra | dyskinetoplastic | RNA editing | mitochondrial DNA

Salivarian trypanosomes are extracellular protozoan parasites that cause important diseases in humans (human African trypanosomiasis) and their livestock (nagana). They predominantly infect the blood but, depending on the (sub)species, also other organs, such as the CNS. Transmission typically occurs through the saliva of blood-sucking insect vectors during feeding. The life cycle of African trypanosomes, such as Trypanosoma congolense, T. brucei brucei, T. b. gambiense, and T. b. rhodesiense, is fully dependent on the tsetse fly vector and is the pathogenic bloodstream form of T. brucei, the huge and highly complex (1). Thus, these parasites are restricted to areas inhabited by the tsetse fly (i.e., sub-Saharan Africa).

However, some pathogenic trypanosome species have adapted to efficient tsetse-independent transmission, abandoning any developmental stages associated with that vector, and therefore, they were able to escape from the African tsetse belt. T. evansi is transmitted mechanically by biting flies when the insect’s blood meal on an infected host is interrupted and a second host is bitten with trypanosome-contaminated mouth parts shortly thereafter (2). T. evansi infects various mammalian animals, including livestock, and it is the pathogenic trypanosome with the widest geographical distribution. The disease caused by this parasite, therefore, has many different names but is known as surra in large parts of Asia. The second species, T. equiperdum, causes a sexually transmitted horse disease called dourine and predominantly infects genital tissues (2). T. evansi and T. equiperdum are morphologically indistinguishable from each other and T. b. brucei, and their status as independent species has been questioned (3, 4).

Mitochondrial DNA (mtDNA) of trypanosomatids is organized as the kinetoplast (kDNA), a gigantic network of concatenated, circular DNA molecules (5). The second key feature distinguishing T. evansi and T. equiperdum from T. b. brucei is that they are dyskinetoplastic [DK; i.e., lacking all (akinetoplastic or kDNA) or critical parts (kDNA')] of their mitochondrial DNA] (6). T. brucei kDNA contains two types of molecules. The ~23-kb maxicircle, present in 20–50 copies, contains a typical set of rRNA and protein-coding genes, most of which encode subunits of respiratory chain complexes (6). Most trypanosomatid mitochondrial mRNAs require a unique form of posttranscriptional editing before they can be translated into functional proteins (7). The second kDNA component is a highly diverse set of thousands of ~1-kb minicircles, which encode guide RNAs required for editing. Maintenance and expression of kDNA require numerous essential enzymes and have been suggested as a target for existing and novel drugs for T. brucei and other trypanosomatids (8). Indeed, antitrypanosomatid therapeutics, such as pentamidine and ephedrine, have been shown to directly affect kDNA (9, 10). T. equiperdum strains typically have retained their maxicircle—in some cases with substantial deletions—but have lost their minicircle diversity (4, 6). T. evansi strains do not have a maxicircle and either show minicircle homogeneity or are akinetoplastic. Consequently, both species are incapable of functional mitochondrial gene expression.

T. evansi and T. equiperdum cannot develop in the tsetse fly, probably because ATP production in that environment requires oxidative phosphorylation (11) and, therefore, the capacity to express numerous mitochondrial genes. They can only survive as bloodstream forms (BFs), which produce ATP exclusively through glycolysis; however, they still require a mitochondrion, because it hosts other essential activities (12–14). A key process underpinning mitochondrial function is the maintenance of an electrochemical potential, Δψ, across the inner mitochondrial membrane (15). BF T. brucei, which lacks the proton pumping respiratory chain complexes III and IV, generates Δψ using the mitochondrial F1F0-ATP synthase complex functioning in reverse to pump protons from the matrix into the intermembrane space (16–18). Subunit a of the membrane-embedded F0 part is critical for proton translocation (Fig. S14) and kDNA-encoded, and its pre-mRNA requires substantial RNA editing (19). DK trypanosomes are incapable of expressing subunit a, because they lack either the gene itself or most, if not all, guide RNAs. The puzzling fact that these organisms are viable was hypothesized to involve compensatory mutations in the nucleocytoplasmic Δψ generation (4, 17).

We tested this hypothesis by generating BF T. b. brucei that express mutated γ subunits and investigating their response to kDNA loss. Our results show that a single amino acid change in subunit γ fully compensates for complete loss of kDNA and provide insight into the molecular mechanism of this compensation. This finding has important consequences for our understanding of...
the mitochondrial function in these organisms, their evolution, and the suitability of kDNA as a drug target.

Results

We first investigated the significance of the F1γ L262P mutation identified in the T. b. brucei 164DK cell line (17). These cells had lost their kinetoplast after several months of in vivo selection with aclaravine (Acr), a DNA-intercalating compound (20) (Table 1 lists the mutations investigated, and Table S1 lists the trypanosome cell lines and strains used in this study). We first determined whether this mutation confers Acr resistance in a standard 3-d drug sensitivity assay (21). We combined introduction of a single ectopic subunit γ allele bearing the L262P mutation (γL262P) into T. b. brucei 427 with a single (sKO) or double KO (dKO) of the endogenous γ gene, resulting in cell lines sKO + γL262P and dKO + γL262P. Cell lines expressing an ectopic WT allele (γWT) were generated as controls. We then compared the Acr sensitivity of these cell lines with parental T. b. brucei 427, T. evansi Antat 3/3 (which has the A281del mutation) (Table 1), and T. b. brucei 164DK (the source of the L262P mutation).

For T. b. brucei 164DK, the average EC50 value for Acr was sevenfold greater than for the T. b. brucei 427 WT strain (Fig. 1A). T. evansi Antat 3/3 had an intermediate EC50 value. Strains expressing γL262P showed considerable resistance, similar to the resistance level of the 164DK cell line, in which this mutation had originally been identified. Although the EC50 for sKO cells was slightly lower than for dKO cells, this result was not statistically significant (P value = 0.062; unpaired two-tailed Student t test). In contrast, cells expressing the ectopic γWT retained the 427 parental sensitivity. These results show that the L262P mutation is sufficient to confer a level of Acr resistance that is similar to the one observed for T. b. brucei 164DK.

Next, we investigated if F1γ mutations identified in DK trypanosomes (Table 1) enable long-term survival of parasites in the presence of 20 nM Acr, a concentration well above the 3-d parental sensitivity. These results show that the L262P mutation is sufficient to confer a level of Acr resistance that is similar to the one observed for T. b. brucei 164DK.

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Table 1. ATPase subunit γ sequence variations tested in this study

<table>
<thead>
<tr>
<th>Mutation</th>
<th>Source (origin, host, year of isolation)</th>
<th>Source for mutation; isolate</th>
</tr>
</thead>
<tbody>
<tr>
<td>L262P</td>
<td>T. b. brucei 164DK (USA, mouse, 1971)</td>
<td>17; 20</td>
</tr>
<tr>
<td>CTT→CCT</td>
<td>T. evansi BoTat1.1 (Morocco, horse, 1924)</td>
<td>3</td>
</tr>
<tr>
<td>A273P</td>
<td>T. equiperdum BoTat1.1 (Morocco, horse, 1924)</td>
<td>3</td>
</tr>
<tr>
<td>GCG→CGG</td>
<td>T. equiperdum BoTat1.1 (Morocco, horse, 1924)</td>
<td>3</td>
</tr>
<tr>
<td>A281del</td>
<td>T. equiperdum ATCC30019 (France, horse, 1903)</td>
<td>3</td>
</tr>
<tr>
<td>TCTGCTATG→TCT—ATG</td>
<td>T. equiperdum ATCC30023 (France, horse, 1903)</td>
<td>3</td>
</tr>
<tr>
<td>T. evansi Antat 3/3 (South America, capybara, 1969)</td>
<td>This work; 31</td>
<td></td>
</tr>
<tr>
<td>T. evansi C13 (Kenya, camel, 1982)</td>
<td>This work; 31</td>
<td></td>
</tr>
<tr>
<td>T. evansi CPOgz1 (China, water buffalo, 2005)</td>
<td>This work; 4</td>
<td></td>
</tr>
<tr>
<td>T. evansi E110 (Brazil, capybara, 1985)</td>
<td>This work; 44</td>
<td></td>
</tr>
<tr>
<td>T. evansi E9/CO (Columbia, horse, 1973)</td>
<td>This work; 51</td>
<td></td>
</tr>
<tr>
<td>T. evansi SS143M (Philippines, water buffalo, 2006)</td>
<td>This work; 4</td>
<td></td>
</tr>
<tr>
<td>T. evansi SS143M (Philippines, water buffalo, 2006)</td>
<td>This work; 4</td>
<td></td>
</tr>
<tr>
<td>T. evansi STIB805 (China, water buffalo, 1985)</td>
<td>4; 4</td>
<td></td>
</tr>
<tr>
<td>T. evansi STIB807 (China, water buffalo, 1979)</td>
<td>This work; 4</td>
<td></td>
</tr>
<tr>
<td>T. evansi STIB810 (China, water buffalo, 1985)</td>
<td>4; 4</td>
<td></td>
</tr>
<tr>
<td>T. evansiStock Kazakh (Kazakhstan, camel, 1995)</td>
<td>This work; 4</td>
<td></td>
</tr>
<tr>
<td>T. evansi KETRI2479 (Kenya, camel, 1981)</td>
<td>4; 51</td>
<td></td>
</tr>
</tbody>
</table>

In all strains investigated, the L262P and A273P mutations are homozygous, whereas the A281del and M282L mutations are heterozygous.

Dramatically different Acr sensitivities were observed for T. b. brucei expressing F1γ with mutations identified in field isolates of T. evansi and T. equiperdum from various geographical areas (Table 1). Cells expressing γM282L behaved like WT and were dead by 72 h (Fig. 1B and Fig. S2B); in contrast, the γA273P-expressing cells continued to grow like γL262P-expressing cells (Fig. 1B). The result for γA281del-expressing cells (the mutation present in the vast majority of field isolates; see Table 1) was more complex. Although some clones died as quickly as the negative controls, others persisted, and a small number of live cells were still observed after 7 d (Fig. 1B and Fig. S2B, asterisks). After transfer to Acr-free medium, these cells recovered, and when reexposed to Acr, no lag in growth was observed (Fig. 1E), suggesting that they had undergone a secondary adaption. Like the γL262P-expressing cells, all cell lines that had survived Acr treatment were kDNA−17. Expression of functional F1γ-ATPase was broadly equivalent (Fig. S3), ruling out variations in expression level as an explanation for the differing Acr sensitivities. The antibody that we used detects the 15 kDa N-terminal domain of the α subunit, which in trypanosomatids, is proteolytically cleaved off from the rest of the polypeptide but remains associated with the F1 complex (22). Interestingly, the β-barrel domain seems to have been lost or altered in T. evansi Antat 3/3, because here, the 15 kDa band was absent (Fig. S3A, lane 8). Whether this absence is a general feature of the T. evansi and T. equiperdum F1γ-ATPase and if it is of functional significance remains to be investigated. Loss of the domain is not a direct consequence of kDNA loss, because the dKO + γL262P cells retained the domain after Acr treatment (Fig. S3A, compare lane 3 with lane 4). Growth rates for sKO or dKO cells expressing a mutated γ subunit in the absence of Acr were very similar to WT (Fig. S2 A and B), suggesting that, under standard culture conditions and within the time frame observed, the mutated subunits were fully functional. The γA281del-expressers seemed to not be adapted as well to life without kDNA as cells expressing γL262P or γA273P, which was indicated by the slightly reduced growth rate in Acr-free medium (Fig. S4).
These results suggested that strains of *T. evansi* and *T. equiperdum* with an A281del or A273P mutation depend on these mutations to be viable. To test this suggestion further, we inductively expressed either γWT or γL262P in *T. evansi* Antat 3/3 (these parasites are heterozygous for the A281del mutation; see Table 1). Expression of γWT caused a strong growth defect after 48 h, whereas expression of γL262P had no effect (Fig. 2A), suggesting that replacement of sufficient endogenous (A281del-mutant) γ subunits in the ATP synthase complex with WT subunits severely impairs the viability of these DK cells. This observation also suggested that the L262P mutation from the laboratory-induced DK strain of *T. b. brucei* and the A281del mutation present in *T. evansi* Antat 3/3 are, at least to some extent, functionally interchangeable. In another experiment, we introduced expression of an ectopic γL262P allele in *T. b. brucei* 427 cells and then treated the cells with 20 nM Acr. As predicted, these cells were resistant to Acr and rapidly lost their kinetoplast. Subsequent repression of the γL262P allele forced these DK cells to rely on endogenous γWT alone and caused a severe growth defect after 24 h (Fig. 2B). In contrast, the growth rate of cells with maintained expression of γL262P remained constant. These results confirm that, after expression of a mutated F1γ subunit has permitted loss of kDNA, the DK trypanosomes now depend on continued expression of this mutated subunit to remain viable. Because specific F1γ mutations are able to compensate for kDNA loss, all genes exclusively involved in kDNA biogenesis or expression would be predicted to become dispensable. RNA editing ligase 1 (REL1) is essential in the kinetoplast mRNA editing process and its knockdown lethal (23). To determine whether the L262P mutation can compensate for REL1 loss, the γL262P gene or the γWT control was constitutively expressed in *T. b. brucei* 427 engineered for inducible REL1 RNAi. Contrasting with γWT cells, γL262P-expressing trypanosomes showed no growth effect after RNAi-mediated knockdown of REL1 (Fig. 3). Interestingly, γL262P-expressing *T. b. brucei* was not viable after ablation of mitochondrial topoisomerase II (Top2mt), a protein essential for kDNA replication (24). To validate this observation, we ablated Top2mt expression in *T. evansi* Antat 3/3 and observed a growth defect after 48 h (Fig. S5A). Secondary effects resulting from faulty kDNA replication (for example, potentially toxic accumulation of kDNA) are unlikely to be responsible, because the Antat 3/3 strain used for this study is kDNA*−* (Fig. S5B) (17), probably as a result of long-term in vitro culture (25). The most likely explanation for the growth defect in *T. evansi* is, therefore, that Top2mt has an additional important function outside of its role in kDNA replication. Nevertheless, together with the experiments investigating chemically induced kDNA loss, these genetic data show that specific point mutations of ATPase γ are fully sufficient to compensate for loss of kDNA or its gene expression.

The proton-translocating function of the membrane-embedded F0 part of the ATP synthase involves subunit α, which is thought to be the only ATPase subunit encoded in the kinetoplast (Fig. S1A). To test directly whether a mutated ATPase γ is necessary for generating ΔΨm in kDNA*−* cells, we measured ΔΨm in BF trypanosomes over the course of Acr treatment. For cells expressing only γWT, ΔΨm decreased after 24 h and was completely abolished after 48 h (Fig. 4A), preceding cell death by at least 24 h. In contrast, ΔΨm of the γL262P-expressing trypanosomes was not affected by Acr-induced kDNA loss (Fig. 4B). The slight decrease in ΔΨm during Acr exposure is probably the result of kDNA-independent Acr toxicity, because no difference in ΔΨm could be observed after removal of Acr from the medium (Fig. S6). The current model for ΔΨm generation in BF trypanosomes proposes that increased ATP hydrolysis by a mutated F1 part facilitates the electronegative exchange of cytosolic ATP94 into mitochondrial ATP93 by the ADP/ATP carrier (AAC) (17, 26). We measured sensitivity of our cell lines to oligomycin, an inhibitor of the coupled F1F0-ATPase, and the AAC inhibitor bongkrekic acid. Trypanosomes expressing mutant ATPase γ showed similar levels of oligomycin resistance before and after Acr-induced kDNA loss.

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**Fig. 1.** Mutations in ATPase γ allow BF *T. b. brucei* to survive kDNA loss. (A) Acr sensitivity of γL262P-expressing and control BF trypanosomes given as EC50 values. Error bars are SEM; n ≥ 3. (B) Cumulative growth in 20 nM Acr of cells ectopically expressing WT F1γ or an L262P, A273P, A281del, or M282L mutated copy in an sKO background. Numbers indicate independent clones. Parental WT strain 427, *T. b. brucei* 164DK, and *T. evansi* Antat 3/3 were also analyzed. Fig. S2A and B shows growth curves without Acr and for dKO cells. (C) Differential interference contrast (DIC) and fluorescence microscopy of DAPI-stained dKO + γL262P trypanosomes before and after exposure to 20 nM Acr. White arrowhead in pre-Acr exposure images indicate the kinetoplast. (Scale bars: 5 μm.) (D) PCR assay for presence of kDNA-encoded genes A6, ND4, ND7, and NDS in dKO + γL262P cells before and after Acr exposure. The faint band observed with ND4 primers post-Acr treatment is a result of nonspecific amplification, which is shown by its larger size. The nuclearly encoded dihydrodipicolinate dehydrogenase gene (*LipDH*) was assayed as a positive control. (E) Cumulative growth in the presence of 20 nM Acr of previously Acr-treated (and therefore, DK) γ-A281del clones 2 and 9 after they had been allowed to recover in Acr-free medium (dashed lines; B and Fig. S2B show the initial response of these clones to Acr exposure). The same clones but without any prior Acr exposure were included in the analysis (solid lines). The parental *T. b. brucei* 427 strain was assayed for comparison.
activities, even in kDNA evolve from T. evansi mutations in yeast T. evansi cells. Such a supercomplex might in mtDNA L262P mutation allows BF γ by the AAC (17, 18). The L262P mutation was expressed in T. brucei cells, and kDNA loss was triggered by exposure to 20 nM Act for 7 d. The culture was split, and expression of L262P in one subculture was repressed by transfer to Tet-free medium (0 h). Cumulative cell growth in the presence (dashed line and open circles) or absence (solid line and closed circles) of Tet was determined.

(Fig. 4C), suggesting that these mutations fully uncouple the F₁ and Fₒ activities, even in kDNA cells. In support of the proposed importance of the AAC for ∆Ψm generation in the absence of a proton-pumping F₀F₁-ATPase, cells expressing L262P showed increased sensitivity to bongkrekic acid. Again, this change in sensitivity was irrespective of the presence or absence of kDNA (Fig. 4D), sustaining the hypothesis that, even in the presence of a functional Fₒ, the mutation results in a complete switch to the alternative mode of ∆Ψm generation.

Discussion

The apparently conflicting observations that BF T. brucei depends on kDNA for survival (23, 27, 28), whereas the closely related species T. evansi and T. equiperdum as well as certain laboratory-generated T. brucei strains are viable, despite the loss of all or critical parts of their kDNA (6), have generated a puzzling conundrum. In this report, we show that specific single amino acid mutations in the nuclearly encoded γ subunit of the mitochondrial ATP synthase complex are fully sufficient to compensate for loss of kDNA-encoded gene products in the BF parasite. The underlying biochemical mechanism involves uncoupling of the F₁ and Fₒ parts of the enzyme and increased dependence on a highly active AAC. Some field isolates seem to depend on additional adaptations that remain to be identified.

Four different candidate mutations have been identified to date (Table 1). We found that the L262P and A273P mutations are fully sufficient to permit normal growth of BF parasites in the absence of kDNA. Neither chemically induced loss of the kinetoplast nor loss of kDNA expression by RNAi resulted in any lag in growth that would have indicated the requirement for additional adaptations. This finding also rules out a potential kDNA replication checkpoint for cell cycle completion (29). More than 40 y after the generation of the only surviving DK T. b. brucei strain (20), its viability can now be explained. Likewise, the A273P mutation present in certain field isolates of T. equiperdum strains is fully sufficient to explain why these strains are viable, despite the lack of mtDNA. One might postulate that it uncouples the F₁ and Fₒ ATPases of the enzyme. This scenario is reminiscent of mgi mutations in yeast (Fig. S1 B–D), although in that organism, it is not clear whether uncoupling is directly linked to viability without a mitochondrial genome (31). A reduced Kₘ for ATP was also suggested as part of the mechanism for yeast mgi mutations (32), and the effects of the trypanosome γ mutations on F₁-ATPase kinetics remain to be investigated. Both mechanisms could result in increased ATPase activity and be necessary to provide sufficient ADP⁺ for an efficient electrogenic exchange with ATP⁺ by the AAC (17, 26, 32). Our finding of bongkrekic acid hypersensitivity in cells expressing the mutant γ shows an increased importance for the role of the AAC in these cells. There is evidence in yeast and Leishmania for an AAC/Fₐₐₜ-ATPase supercomplex (33, 34), which might explain the apparent preference for the ATPase as the source of ADP⁺ in mtDNA cells. Such a supercomplex might generate localized zones of increased ∆Ψm and, in association with the protein import machinery, exploit them for protein import, similar to the supercomplex of Tim21 and respiratory complexes III and IV reported for yeast (35). A number of protists, including many important parasites, lack classical mitochondria and instead, harbor related organelles classified as hydrogenosomes or mitosomes that usually lack an organellar genome. How these organelles generate a ∆Ψm is unclear (36, 37), and trypanosomes promise to be an excellent experimental system to further investigate this important problem in cell biology.

How did T. evansi and T. equiperdum evolve from T. brucei, and can the findings reported here help to address this question?

Fig. 3. The L262P mutation allows BF T. b. brucei to survive inhibition of mitochondrial gene expression. (A) Cumulative growth of T. b. brucei after RNAi-mediated knockdown of REL1 (dashed lines). Cells expressed either γWT (open circles) or γL262P (closed squares). REL1 RNAi was induced with 1 μg/mL Tet; uninduced control cultures are shown as solid lines. (B) Western analysis of REL1 protein expression from whole-cell lysates taken at the 72-h time point in A; α-tubulin was used as a loading control (lower).
likely that isolates sharing the same mutation also share the same ancestor. Importantly, the largest group, characterized by the A281del mutation, contains isolates from Africa, Asia, and South America collected during a span of 100 y, and it contains both T. evansi and T. equiperdum isolates. Either these T. evansi isolates are T. evansi isolates that had been misclassified (3) or many isolates of these two species are descendants of the same evolutionary event (43, 44). Based on these considerations, we would propose that, contrary to what was proposed recently (4), the extant strains seem to be the result of a limited rather than large number of independent evolutionary events. It has been suggested that T. evansi/T. equiperdum are analogous to yeast petite mutants (4, 38), and, indeed, the mechanism that allows mtDNA loss in petite-negative yeast and T. brucei is strikingly similar (17). However, it is important to stress that the ability to survive without a mitochondrial genome is only one distinctive feature of these parasites, because the other one, efficient tsetse-independent transmission, has had such dramatic epidemiological consequences.

Maintenance and expression of kDNA has been suggested as a target for existing and novel antitrypanosomatid drugs (8–10). In fact, the Acr compound used for kDNA elimination in the present study was originally developed with the aim of finding a cure for sleeping sickness (45). The fact that single nucleotide changes can make the parasite completely independent of kDNA and its expression suggests that this target needs to be treated with caution, at least for T. brucei. However, a few considerations are important. First, an uncoupled F_{0}F_{1} enzyme will be incapable of proton gradient-driven ATP synthesis, which is thought to be required for survival in the tsetse vector because of the low abundance of glucose in the insect’s midgut (11, 46). The mutation would, consequently be expected to lock the parasite in the mammalian host and prevent spread of resistant parasites. Second, other pathogenic trypanosomatids, including T. congolense, T. cruzi, and Leishmania spp., do not seem to be able to circumvent the need for functional kDNA, which therefore, remains a highly promising target in those parasites.

Another important conclusion from our study is that ATP synthase subunit a seems to be the sole kDNA product ultimately required for viability of BF T. brucei. A subunit of the mitochondrial ribosome, RPS12, may be encoded in kDNA (47), and it has been reported that the product of an alternatively edited mRNA, AEP-1, is required for kDNA maintenance (48); however, these proteins would also become dispensable, along with subunit a, in the presence of a compensatory ATP synthase γ-mutation. The same is true for the numerous nuclearly encoded proteins required exclusively for maintenance and expression of kDNA (5, 7, 49). Thus, a single amino acid mutation in BF T. brucei makes a large number of otherwise essential proteins redundant. The compensatory mutations reported in this paper offer an attractive tool for their identification and characterization.

Materials and Methods

Materials, Trypanosome Strains, and Culture Conditions. Details on materials can be found in SI Materials and Methods. All experimental work and culturing were carried out with BF trypanosomes only. ATP synthase γ (Trb927.10.180) genetic manipulations in T. b. brucei were conducted on the Lister 427 strain, except for inducible expression and RNAi experiments, which used the 427 single marker cell line (50). Inducible expression in T. evansi was conducted in a cell line expressing T7RNA and TET (17). Cell lines T. evansi Antat 3/3 (51) and T. b. brucei DK 164 (20) were included in growth experiments for comparison. SI Materials and Methods has details on culturing, plasmid construction, and transfection. See Table S2 for oligonucleotides.

Alamar Blue Dose–Response Assay. The Alamar Blue assay was performed essentially as described (21) with minor modifications. Briefly, test compound was doubly diluted in 100 μL Hirumi-modified Iscove’s medium 9/10% (vol/vol) PBS in a 96-well plate; an equal volume of medium containing 80 BF trypanosomes was added to give a final density of 5 × 10^5 cells/mL. The plate was incubated at 37° C and 5% (vol/vol) CO_{2} to 72 h, after which 20 μL 0.5 mM resazurin sodium salt in PBS were added to each well; then, the plate was incubated for another 4 h. Fluorescence was measured in a plate reader with excitation and emission filters of 544 and 590 nm, respectively. EC_{50}.
values were derived from a variable slope (four parameter) nonlinear regression using Prism 5 software (GraphPad).

DAPI Staining. Trypanosomes were washed in PBS, settled onto poly-L-lysine coated slides, and fixed in 3% (wt/vol) formaldehyde for 10 min before treating with excess cold methanol for at least 30 min. Slides were then rehydrated in PBS and mounted in glycerol containing 1 μg/mL DAPI before imaging using a Leica SP5WCS confocal laser microscope (blue diode laser at 405-nm wavelength) with Volocity version 5.2 image analysis software (PerkinElmer).

PCR and Western Analyses. Details are given in SI Materials and Methods.

**Ym Measurements.** Samples of trypanosome cultures, either exposed or unexposed to Acr, were incubated with 260 μM rhodamine 123 (Rh123) for 20 min at 37 °C. Cells were harvested by centrifugation at 1,300 × g for 10 min and washed once with 25 mM Hepes, pH 7.6, 120 mM KCl, 0.15 mM CaCl2, 10 mM KH2PO4/K2HPO4, pH 7.6, 2 mM EDTA, 5 mM MgCl2, and 6 mM β-Glucose. Fluorescence caused by Rh123 uptake was measured using a FACS caliber flow cytometer with CellQuest Pro software (Becton Dickinson). Baseline fluorescence was subtracted for each sample. The percentage of cells with 10 μM FCCP before adding Rh123: the FCCP concentration was maintained throughout the wash and flow cytometer steps.

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