



Oxidative Phosphorylation Is Required for Powering Motility and Development of the Sleeping Sickness Parasite Trypanosoma brucei in the Tsetse Fly Vector

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ABSTRACT The single-celled parasite Trypanosoma brucei is transmitted by hematophagous tsetse flies. Life cycle progression from mammalian bloodstream form to tsetse midgut form and, subsequently, infective salivary gland form depends on complex developmental steps and migration within different fly tissues. As the parasite colonizes the glucose-poor insect midgut, ATP production is thought to depend on activation of mitochondrial amino acid catabolism via oxidative phosphorylation (OXPHOS). This process involves respiratory chain complexes and F₁F₀-ATP synthase and requires protein subunits of these complexes that are encoded in the parasite's mitochondrial DNA (kDNA). Here, we show that progressive loss of kDNA-encoded functions correlates with a decreasing ability to initiate and complete development in the tsetse. First, parasites with a mutated F₁F₀-ATP synthase with reduced capacity for OXPHOS can initiate differentiation from bloodstream to insect form, but they are unable to proliferate in vitro. Unexpectedly, these cells can still colonize the tsetse midgut. However, these parasites exhibit a motility defect and are severely impaired in colonizing or migrating to subsequent tsetse tissues. Second, parasites with a fully disrupted F₁F_o-ATP synthase complex that is completely unable to produce ATP by OXPHOS can still differentiate to the first insect stage in vitro but die within a few days and cannot establish a midgut infection in vivo. Third, parasites lacking kDNA entirely can initiate differentiation but die soon after. Together, these scenarios suggest that efficient ATP production via OXPHOS is not essential for initial colonization of the tsetse vector but is required to power trypanosome migration within the fly.

IMPORTANCE African trypanosomes cause disease in humans and their livestock and are transmitted by tsetse flies. The insect ingests these parasites with its blood meal, but to be transmitted to another mammal, the trypanosome must undergo complex development within the tsetse fly and migrate from the insect's gut to its salivary glands. Crucially, the parasite must switch from a sugar-based diet while in the mammal to a diet based primarily on amino acids when it develops in the insect. Here, we show that efficient energy production by an organelle called the mitochondrion is critical for the trypanosome's ability to swim and to migrate through the tsetse fly. Surprisingly, trypanosomes with impaired mitochondrial energy production are only mildly compromised in their ability to colonize the tsetse fly midgut. Our study adds a new perspective to the emerging view that infection of tsetse flies by trypanosomes is more complex than previously thought.

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he protist parasite *Trypanosoma brucei* is the causative agent of human African trypanosomiasis, also called sleeping sickness (1). The life cycle of *T. brucei* is complicated in that it involves at least nine distinct stages within the mammalian bloodstream and tissues and the tsetse fly vector (2-4). The distinct biology of these life cycle stages is closely linked to changes in energy metabolism and regulation of the activity of respiratory chain and F₁F₀-ATP synthase complexes in the inner membrane of the parasite's single mitochondrion. As in other eukaryotes, these complexes are composed of both nuclearly and mitochondrially encoded subunits, and regulation of their expression therefore requires tight coordination between these organelles. In kinetoplastids such as T. brucei, structure and expression of the mitochondrial genome, termed kinetoplast DNA (kDNA), are extraordinarily complex. T. brucei kDNA comprises dozens of maxicircles (the equivalent of mitochondrial DNA in other eukaryotes) and thousands of minicircles (5). The latter encode guide RNAs (gRNAs) for remodeling of 11 of the 18 maxicircle-encoded mRNAs by uridylyl insertion and deletion, a maturation process called RNA editing (6).

In the proliferative long slender bloodstream form (BSF), ATP production is nonmitochondrial and occurs via glycolysis, exploiting the glucose-rich environment within the mammalian bloodstream (7, 8). The BSF mitochondrion lacks cyanide-sensitive oxygen uptake: respiratory complexes III (ubiquinol:cytochrome c oxidoreductase) and IV (cytochrome oxidase) are absent, and the corresponding maxicircle- and nucleusencoded transcripts are present at only very low levels (9-13). The BSF mitochondrion also lacks or has less developed cristae, which are invaginations of the inner mitochondrial membrane that are ultrastructural hallmarks of oxidative phosphorylation (OXPHOS) (14, 15). Despite the absence of cytochrome-mediated respiration, BSF parasites are dependent on oxygen for survival, since an alternative oxidase (AOX) in the inner mitochondrial membrane is critically involved in maintaining the redox balance for glycolysis (16). BSF cells use the F_1F_0 -ATP synthase complex in reverse: ATP hydrolysis by the F_1 part provides the energy for pumping protons from the mitochondrial matrix through the F_0 part into the intermembrane space. This activity generates a mitochondrial membrane potential ($\Delta\Psi$ m), which is essential for protein import and exchange of metabolites and therefore fundamental for mitochondrial function (17-19).

Long slender BSF T. brucei organisms differentiate to the cell cycle-arrested stumpy BSF upon reaching high parasite numbers, induced by quorum sensing (20-22). The stumpy form is preadapted to survival in the tsetse fly vector by being more resistant to pH and proteolytic stresses and by beginning to prepare its mitochondrial energy metabolism for the change in nutrient availability. This adaptation includes upregulation of mitochondrial ATP production capability via substrate-level phosphorylation and beginning of cristae formation (20, 23-25).

Once ingested with a blood meal, stumpy forms differentiate into proliferative procyclic forms (PCF) within the fly midgut. Mitochondrial ATP production is vital to the PCF (26-28). Its inner mitochondrial membrane is now invaginated to form prominent cristae and contains complexes I (NADH:ubiquinone oxidoreductase), II (succinate dehydrogenase), III, and IV of a classical cytochrome-mediated respiratory chain, along with complex V (F₁F₀-ATP synthase) (29, 30). In the tsetse fly midgut, proline and intermediates of the tricarboxylic acid cycle are thought to be the main energy sources (31-35), with glucose being only transiently abundant after blood meals (36). Under glucose-depleted in vitro growth conditions, which are similar to the tsetse fly midgut environment, ATP production via OXPHOS is essential (28, 37). Under glucose-rich in vitro growth conditions, only ATP production by mitochondrial substrate-level phosphorylation appears to be essential in PCF (26, 27); however, the electron transport chain remains essential under these conditions (27, 38). Complexes III and IV, despite



being structurally divergent (9, 39-43), are thought to have classical enzymatic roles in the PCF respiratory chain: by pumping protons from the mitochondrial matrix into the intermembrane space, they generate a proton motive force and a $\Delta\Psi m$ upon the uptake of oxygen (44, 45). Under glucose-rich conditions, it is possible that this generation of $\Delta\Psi$ m becomes the sole essential function of complexes III and IV. Regardless of whether production of mitochondrial ATP involves OXPHOS or substrate-level phosphorylation, it requires the ADP/ATP carrier (AAC) to function in the direction of ATP export from the mitochondria to fuel processes in other parts of the cell (46).

By around day 3 to 7 postinfection, PCF are found in the ectoperitrophic space of the midgut, which, according to a long-standing model, they have reached by crossing the peritrophic matrix (PM) in the anterior midgut (4). However, recent findings suggest that T. brucei colonizes the proventriculus' (PV) ectoperitrophic space by crossing the immature, fluid-like PM before it hardens (47). Within this revisited model, procyclic trypanosomes may colonize first the anterior midgut or establish parallel infections at both the anterior midgut and PV (47-51). Regardless of the topology of the infection, after approximately 2 weeks, parasites differentiate into long epimastigotes within the PV lumen, a process that involves the reversal of the positions of the kinetoplast and nucleus (52). Long epimastigotes then divide asymmetrically into long and short epimastigotes, aiding the delivery of short, less motile epimastigotes to the salivary glands (53, 54). Few parasites are able to complete this complex migration (55), and only a small proportion (~10%) of experimentally infected tsetse flies are found with infected salivary glands (56); in the field, salivary gland infection rates are even lower than experimentally obtained ones (<1%) (57–59). Once in the salivary glands, epimastigotes attach to the epithelium, proliferate, and thus colonize these tissues (4, 60). Epimastigotes undergo further asymmetric divisions to continuously yield trypomastigotes that are able to develop into cell cycle-arrested metacyclics around 3 to 4 weeks postinfection (61). After injection into the mammalian host upon a tsetse bite, metacyclics develop into BSFs to begin the life cycle again. Much less is known about potential changes in mitochondrial energy metabolism after the PCF parasites have left the midgut, owing to the difficulty of culturing other insect stages in the laboratory. Recently, it was discovered that overexpression of the regulatory RNA binding protein RBP6 in PCF triggers efficient differentiation into metacyclic forms in vitro (62). Using this experimental system, a switch from cytochrome-based to AOX-mediated respiration and a general retrogression of mitochondrial structure in metacyclic forms were demonstrated (62, 63).

The kDNA encodes essential components of the multisubunit complexes involved in OXPHOS or in the translation of these components: eight subunits of complex I (ND1 to -5 and ND7 to -9), subunit apocytochrome b of complex III, three subunits of complex IV (COXI to -III), subunit A6 of complex V, two hydrophobic protein subunits of the mitoribosome (S12 [RPS12] and S3 [MURF5]), and the two mitoribosomal rRNAs (64, 65). As discussed above, complex V is essential in both BSFs and PCF, and complexes III and IV are essential in PCF. As a consequence, mitochondrial gene expression is critical for survival of both BSF and PCF T. brucei, although with PCF, this has been formally assessed only in vitro (66-69). (The role of complex I in energy metabolism in either the BSF or PCF is uncertain; this has been debated extensively elsewhere [24, 70-77].)

Intriguingly, "akinetoplastic" or "dyskinetoplastic" BSF T. brucei cells, which lack all kDNA (kDNA⁰) or critical parts of kDNA (kDNA⁻), respectively, occur in nature as the subspecies Trypanosoma brucei evansi and Trypanosoma brucei equiperdum and have also been generated in the lab (64, 78-80). In nature, these kDNA mutants are directly transmitted from mammal to mammal, either by mechanical transfer via successive fly bites or sexually, i.e., without any cyclical development or proliferation in an insect vector. This apparent contradiction regarding the essentiality of kDNA in BSF T. brucei was resolved by the discovery of point mutations in the nucleus-encoded F₁-ATPase subunit γ in akinetoplastic and dyskinetoplastic forms, for example, a change of leucine in position 262 to proline (L262P). When introduced into wild type BSFs, these mutations



TABLE 1 Mutations affecting trypanosome mitochondrial functions that were assessed for their effects on tsetse infectivity in this study^a

Gene and mutation (TriTrypDB gene ID)	Expected consequence of mutation(s)	References
kDNA ⁰ (NA)	Complete absence of essential subunits of respiratory complexes I, III, IV, and of the F_o part of complex V. Note that kDNA o can be tested only in combination with a compensatory mutation, in this study the F_1 - γ L262P.	GenBank entry MK584625; 66
F ₁ subunit γ (Tb927.10.180), heterozygous or homozygous L262P mutation (WT/L262Pγ, L262P/L262Pγ)	F_1 -ATPase reaction: appears to lower K_m for ATP and increase rate of ATP hydrolysis; F_1F_0 -ATP synthase reaction: (partial?) uncoupling of F_0 - γ rotation and F_1 activity, resulting in complete or partial loss of OXPHOS in homozygous mutant	17, 66, 84, 85, 88, 89
F _o subunit Tb1 (Tb927.10.520), null mutant (Tb1 ⁻ /Tb1 ⁻)	Loss of F_1F_0 . ATP synthase complex, accumulation of F_1 , resulting in complete loss of OXPHOS	107, 109

^aSee Table S1 for details on the corresponding cell lines. NA, not applicable.

can compensate for kDNA loss by enabling cells to manufacture $\Delta\Psi$ m independently of the F_o part of the ATP synthase (17, 66, 81). Therefore, the F_o subunit A6 is no longer essential (and neither are, consequently, the mitoribosomal protein and rRNA subunits), and the cells can survive as BSFs without any kDNA gene products.

The molecular mechanism of the compensatory mutations in subunit γ is not fully understood, but they enable F_0 -independent generation of $\Delta \Psi m$ via the electrogenic action of AAC-mediated ATP⁴⁻/ADP³⁻ exchange across the inner mitochondrial membrane (17, 24, 66, 82). Kinetic studies with yeast expressing mutated γ subunits have demonstrated a lowered K_m value for ATP in the F₁-ATPase reaction, suggesting that the mutant complex has a higher affinity for its substrate (17, 83). More efficient F₁mediated ATP hydrolysis, perhaps in the vicinity of the AAC, might support this essential electrogenic ATP⁴⁻/ADP³⁻ exchange (17, 83).

On the other hand, the mutations appear to impair the function of the F₁F₀ enzyme when it is required to operate in the direction of ATP synthesis. Physiological and structural studies in yeast have indicated that similar mutations in the yeast ATP synthase γ subunit alter its interactions with the catalytic F_1 - β subunit by partially uncoupling the F₁ moiety from the F_o moiety (84–89). In the presence of an active electron transport chain, impaired coupling of F_1 and F_o action will result in some dissipation of $\Delta\Psi m$ and, consequently, diminished efficiency of the ATP synthesis reaction.

In this study, we investigated the consequences of compensatory $F_1-\gamma$ mutations and kDNA loss for life cycle progression of T. brucei in the tsetse fly vector. Available evidence points to kDNA being essential for PCF viability and fly transmissibility (90). RNA interference (RNAi) knockdown of genes required for kDNA replication and division was lethal in PCF in vitro (69, 91-93), and chemically induced kDNA^o PCF cells were also nonviable in vitro (68, 94). Similarly, from a mixed population of kDNA⁺ and kDNA^o parasites (the latter were obtained by treatment of kDNA⁺ parasites with acriflavine), only kDNA⁺ cells were able to establish a tsetse midgut infection (95). Finally, T. b. evansi and T. b. equiperdum isolates have been documented as being unable to transform into PCF in vitro (81, 96, 97). A caveat with these studies was that they had been performed either with dying parasites unable to survive as BSFs, or with "monomorphic" strains that were unable to develop into the differentiation competent stumpy forms. The ability of viable, "pleomorphic" kDNA^o strains to infect and potentially develop in the tsetse fly vector has never been formally assessed.

Here, we built on our development of a range of pleomorphic BSF mutants that represent various defects in the expression of kDNA-encoded functions (Table 1) to assess the role of kDNA in stumpy-to-PCF differentiation, in the proliferation of PCF in vitro, and in the completion of the parasitic life cycle in tsetse flies.

RESULTS

Ability of F₁F_o-ATP synthase mutants and kDNA^o cells to differentiate to PCF in vitro. To investigate the requirement for functional kDNA and a fully functional F₁F_o-ATP synthase in the differentiation of *T. brucei* from stumpy BSF to PCF, we generated a set of mutant cell lines (Table 1; also, see Table S1 in the supplemental material). In



the pleomorphic cell line EATRO 1125 (AnTat1.1 90:13), we first replaced one allele or both alleles of the nucleus-encoded F_1F_0 -ATP synthase subunit γ with either a wild-type version (WTy, serving as a control) or a version with the L262P mutation (L262Py), as detailed in reference 24. This resulted in the cell lines WT/WTy, WT/L262Py, and L262P/ L262Py, respectively (Table 1 and Table S1). The L262Py mutation, in a homozygous or heterozygous configuration, enables slender BSF T. brucei to survive and proliferate without kDNA (66). We generated two cell lines lacking kDNA (kDNA^o 1 and 2) by treating two distinct clones of genotype WT/L262Pγ (clones 1 and 2) with the kDNA-intercalating dye acriflavine (24). The absence of kDNA in these two cell lines and the lack of an in vitro growth phenotype in the absence of kDNA have been shown previously (24). Moreover, T. brucei kDNA^o parasites can still differentiate to the stumpy BSF (24).

To test the requirement of kDNA for differentiation to PCF, we harvested stumpy kDNA⁰ and control cells from infected mice and induced differentiation into PCF in vitro by adding 6 mM cis-aconitate (CA) to the medium and shifting the temperature from 37°C to 27°C for 24 h (98-102). By microscopic inspection, no intact, motile kDNA^o cells were visible at this point, with only nonmotile cells and debris remaining. This contrasted with all other cell lines, which remained motile, although some debris was present. We then harvested cultures by centrifugation and assessed lysates by Western blotting for expression of EP procyclin, an early marker for PCF differentiation (103–105) (Fig. 1A). Compared to WT/WT γ and WT/L262P γ lysates, the EP procyclin signal was much reduced in material recovered from kDNA^o cultures. These results suggest that kDNA⁰ BSF cells differentiated to PCF but died soon afterward, which implies that the requirement for kDNA in PCF cells cannot be compensated for by the L262P γ mutation.

We performed a time course to compare the kinetics of EP procyclin expression after induction of differentiation with CA in cells with and without kDNA. We collected samples from each cultured cell line at 0, 1, 2, 3, 6, 16, and 24 h after CA induction (+CA), stained them with mouse anti-EP procyclin antibody and anti-mouse IgG Alexa Fluor 488, and analyzed them by flow cytometry (Fig. 1B). Parental, WT/WTγ, and WT/ L262Pγ cells containing kDNA expressed EP procyclin with similar kinetics; 55 to 70% of cells expressed EP procyclin by 6 h postinduction. Two clones (clones 1 and 2) of the L262P/L262Py cell line initially showed slower kinetics of EP procyclin expression (Fig. 1B, inset), but after 6 h, the percentage of EP procyclin-positive cells was similar to that in other cell lines with kDNA. kDNA^o cells, on the other hand, showed a delayed rate of EP procyclin expression, with the proportion of cells expressing the protein increasing to a maximum of 55% by 16 h. As the vast majority of kDNA⁰ cells had died by 24 h after CA induction, this figure included EP procyclin-positive cell debris, which was not gated out of this analysis.

To test if freshly differentiated PCF cells expressing one or two of the mutated $L262P\gamma$ alleles could reenter the cell cycle and proliferate under conditions that require efficient OXPHOS, we resuspended cells in SDM80 medium after 24 h of CA treatment. SDM80 lacks added glucose and contains N-acetyl p-glucosamine to inhibit uptake of residual glucose from fetal calf serum (FCS); it is in this low-glucose environment that PCF depend on OXPHOS to generate ATP (28, 37). Here, day 0 (d0) was defined as being after 24 h in HMI-9 at 27°C plus CA and at 24 h in SDM80 at 27°C. Under these conditions, freshly differentiated WT/WTy and WT/L262Py cells were able to proliferate (Fig. 1C), and analysis of the number of kinetoplasts (K) and nuclei (N) per cell confirmed that they could progress through the cell cycle (Fig. 1D). Cells with two kinetoplasts and one nucleus (2K1N) or with two kinetoplasts and two nuclei (2K2N) were consistently seen in culture across the time course, indicating actively dividing cells. Homozygous L262P/L262Pγ cell lines, however, did not grow in this medium. Although these cells survived under these conditions, they did not thrive and merely maintained the population density. Observation by microscopy revealed that these cells were motile but that forward motility appeared to be impaired compared to that in the Dewar et al. mBio

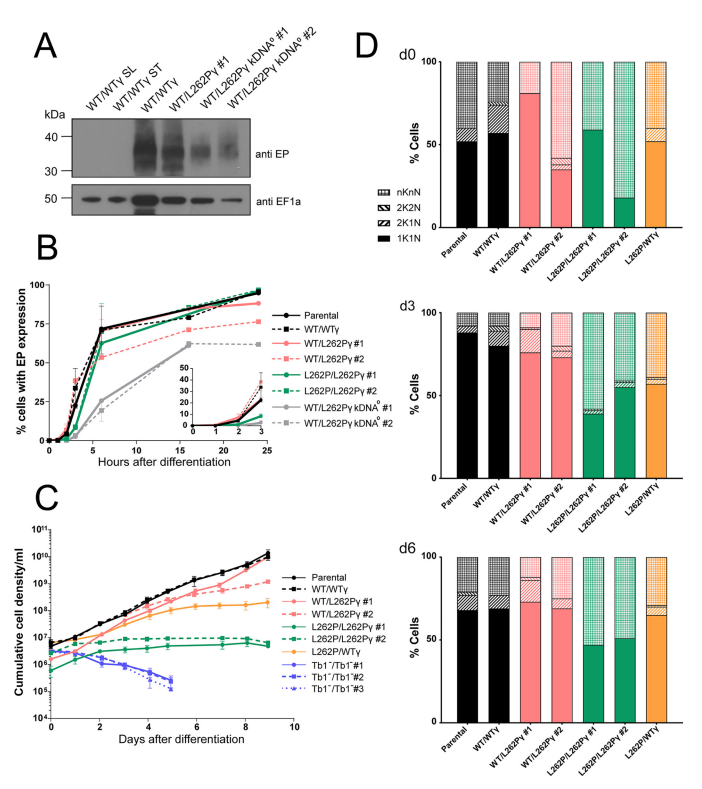


FIG 1 Ability of kDNA $^{\circ}$ cells and F $_1$ F $_0$ -ATP synthase mutants to differentiate into PCF *in vitro*. (A) Western blot to assess EP procyclin expression. Stumpy cells were treated with 6 mM *cis*-aconitate (CA) and incubated at 27 $^{\circ}$ C to induce differentiation. Cells were harvested after 24 h, 2 × 10 $^{\circ}$ cells were loaded per lane of an SDS-PAGE gel, and blots were probed with anti-EP procyclin (showing an expected smear due to heterogenous nature of EP protein [top]) and anti-EF1 α as a loading control (bottom). Slender WT/WT γ cells and stumpy WT/WT γ cells not exposed to CA were analyzed as negative controls for EP expression. (B) EP expression measured by flow cytometry. Stumpy cells were treated with 6 mM CA and incubated at 27 $^{\circ}$ C at 0 h. At each time point, cells were fixed and stained with mouse anti-EP procyclin and anti-mouse IgG Alexa 488 antibodies. Ten thousand cells were analyzed per sample; n = 2. (Inset) Magnification of the time points taken between 0 and 3 h after differentiation. Note that for kDNA $^{\circ}$ cells, numbers at 24 h represent large amounts of cell debris that stained positive for EP. No intact, motile cells were visible upon inspection by microscopy at this time point. (C) Growth of newly differentiated PCF *T. brucei.* Stumpy form cells were treated with 6 mM CA in HMI-9 medium for 24 h at 27 $^{\circ}$ C. Cells were transferred to SDM80 medium at 27 $^{\circ}$ C (0 h), and (Continued on next page)



other cell lines (Movies S1 to S4). The L262P/L262P γ cells were unable to progress through the cell cycle, with barely any 2K1N or 2K2N cell types detectable (Fig. 1D).

Cells in the category nKnN, which includes zoids (nK0N), "monsters" with more than two nuclei and/or kinetoplasts, and other dying cells, were visible in this analysis at every time point and with every cell genotype (Fig. 1D). This was possibly caused by cell swelling and/or nuclear fragmentation due to osmotic stress, which is a side effect of transferring freshly thawed stumpy cells from blood into liquid medium for the differentiation experiments. Notably, for the L262P/L262Py cell lines, the proportion of nKnN cells remained extremely high as the time course continued, consistent with the high proportion of dying cells in these populations.

To confirm that the L262P/L262Pγ genotype, and not a secondary deficiency, was responsible for the severe growth defect of these PCF cell lines, we generated an L262P/WT γ (WT γ add-back) cell line by replacing one subunit γ allele in cell line L262P/ L262P γ clone 2 with a WT subunit γ gene. The ability to maintain growth in SDM80 and progress through the cell cycle was partially restored in the resulting heterozygous γ cell line (Fig. 1C and D), thus confirming an effect specifically due to the L262P γ mutation.

We then asked whether 10 mM glucose in the medium (SDM79 medium) could rescue the growth defect of homozygous L262Pγ PCF cells, as ATP production via OXPHOS was reported to be inessential under these conditions (26, 27, 106), although this may be cell line dependent (107). Surprisingly, we observed an immediate, severe growth phenotype in all freshly differentiated cell lines, with L262Pγ homozygotes having the strongest growth defect (Fig. S1A). In WT/WT γ or heterozygous WT/L262P γ cells, this phenotype could be rescued by removal of glucose from the medium and restored by addition of glucose (Fig. S1B and C). This growth retardation was probably due to an impact of glucose signaling on metabolic adaptation in freshly differentiated PCF parasites (108).

Cells lacking kDNA and homozygous L262Py mutants differ in two key aspects: due to the absence of all kDNA-borne genes, the former cannot generate functional respiratory complexes III (cytochrome bc_1 complex) and IV (cytochrome c oxidase) or the F_0 part of the F_1F_0 -ATP synthase. In the latter, the F_1 and F_0 parts are thought to be functionally uncoupled, although the extent of uncoupling is unclear (Table 1). To further investigate the basis for the phenotypic differences between these cell lines, we generated a cell line lacking ATP synthase subunit Tb1 (Tb1-/Tb1-) (Fig. S2). Tb1 (systematic TriTrypDB ID Tb927.10.520) is the largest T. brucei F_o subunit and required to maintain the intact F₁F_o complex structure and ATP synthase activity; however, it is not required for F_1 -ATPase activity (107, 109). Like other F_o subunits, the gene becomes dispensable in BSF T. brucei in the presence of an L262P γ allele, which enabled us to generate mutants that completely lack Fo but possess all kDNA-borne genes (Fig. S2). Three independent Tb1-null clones (Tb1-/Tb1- clones 1, 2, and 3) were used in this study. This cell line was able to generate stumpy forms in vivo, but once differentiated to the PCF in vitro, they died rapidly (Fig. 1C). This suggests that ATP production by OXPHOS becomes critical upon differentiation into insect stage parasites and that the L262P γ mutation only partially uncouples the F₁F_o-ATP synthase, permitting cell survival, if not proliferation (Fig. 1C), via residual OXPHOS activity.

Mitochondrial ATP production capacity in F₁F_o-ATP synthase mutants. We next investigated the efficiency of mitochondrial ATP production pathways in the different cell lines. We added metabolic substrates that allowed ATP production by OXPHOS only (succinate) or a combination of OXPHOS and substrate phosphorylation (succinate plus pyruvate; α -ketoglutarate) to preparations of disrupted cells containing intact

FIG 1 Legend (Continued)

cell density was determined with a hemocytometer over 24 h. Counting was performed in triplicate with three separate blood cryostocks of each cell line thawed and differentiated at the same time. Error bars indicate standard deviations. kDNA^o cells were not included in this analysis due to their death within 24 h after induction of differentiation. (D) Cell cycle analysis of newly differentiated PCF T. brucei. Slides were prepared at day 0 (d0), d3, and d6 after differentiation and transfer into SDM80, and cells were blindly scored for cell cycle stage. Approximately 100 cells were assessed per strain and time point. The category nKnN includes zoids (i.e., cells with no nucleus), monsters (multiple nuclei), and dying (large, blurry, and/or fragmented nuclei) cells. kDNA^o and Tb1⁻/Tb1⁻ cells were not included in this analysis due to their nonviability after differentiation.



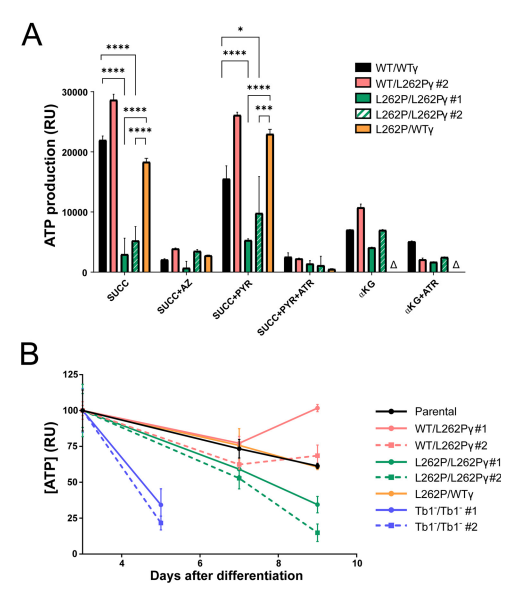


FIG 2 Mitochondrial ATP production capacity in F₁F_o-ATP synthase mutants. Cells were differentiated to PCF as described for Fig. 1 and maintained in SDM80 medium. (A) Mitochondrial ATP production assay using permeabilized differentiated PCF cells with intact mitochondria. The capacity for ATP production via OXPHOS (succinate [SUCC]), substrate-level phosphorylation via the acetate:succinate CoA transferase (ASCT)/succinyl-CoA synthetase (SCoAS) cycle (pyruvate [PYR]), and substrate-level phosphorylation via α -ketoglutarate dehydrogenase and SCoAS (lphaKG) was assessed. Azide (AZ) and atractyloside (ATR) are inhibitors of the F_1F_0 -ATP synthase and AAC, respectively. All assays were performed in triplicate at day 2 after differentiation. Error bars indicate standard deviations. A triangle indicates that the assay was not performed. P values were determined using ANOVA: *, <0.05; **, <0.01; ***, <0.001; ****, <0.0001. See Table S2 for details of statistical comparisons. (B) Total cellular ATP level assays performed in triplicate, with the assay performed at days 3, 5, 7, and 9 post differentiation. The Tb1-/Tb1- cell line was assayed only up to day 5, as insufficient numbers of viable cells remained after that time point. Error bars indicate standard deviations. The ATP level from day 3 for each cell line was set as 100%.

mitochondria. We then measured "in organello" ATP production via the generation of a proportional luminescent signal (110). Using succinate as a substrate, preparations from L262P/L262Py cells generated significantly less ATP than samples from cells with at least one WT γ allele (Fig. 2A and Table S2). Indeed, inhibition of the F₁F_o-ATP synthase with azide reduced ATP production in L262P/L262Py cell lysates only insignificantly, suggesting a very minor, if any, contribution to ATP production by the mutated enzyme under these conditions. Preparations of WT/L262P γ cells produced higher ATP levels than those of WT γ cells; at present we have no explanation for this observation.



The presence of one or two L262P γ alleles did not affect the level of representative F₁ and F_o ATP synthase subunits (Fig. S3), thus confirming the effect of the mutation is due to functional impairment. Adding back a WT γ allele to L262P/L262P γ restored ATP production via OXPHOS nearly to levels found for WT/WTγ cells (Fig. 2A). Interestingly, adding pyruvate as an additional substrate, which theoretically should allow additional ATP production via the acetate:succinate coenzyme A (CoA) transferase (ASCT) cycle, did not significantly increase ATP production in mitochondria of any of the cell lines (Fig. 2A; Table S2). This suggests the absence or low levels of either important transporters, enzymes, or cofactors for this pathway in these early differentiated cells. Similarly, α -ketoglutarate as the sole substrate, which should allow ATP production by substrate phosphorylation via succinyl-CoA synthetase, with production of succinate and subsequent ATP production via OXPHOS, resulted in only a moderate production of ATP in mitochondria from newly differentiated WT γ cells.

The maintenance of total cellular ATP levels after differentiation was also impacted by the presence of homozygous L262Py. By day 9, the level of cellular ATP in L262P/ L262P γ cells was considerably lower than that of cells expressing at least one WT γ allele (Fig. 2B). This is consistent with an impairment of ATP synthase complexes with an L262P γ mutation in coupling the $\Delta\Psi$ m to ATP synthesis, as suggested above. ATP production assays could not be performed on Tb1-/Tb1- cell lines due to the large number of viable differentiated cells required, but measurement of cellular ATP levels at day 5 showed that ATP levels were much depleted in these dying cells (Fig. 2B).

Motility analysis of in vitro cultured F_1F_0 -ATP synthase mutant cell lines. Microscopic observation of L262P/L262Pγ cells grown in SDM80 medium had suggested a motility phenotype. We decided to investigate this further by recording motility tracks from videos of freshly differentiated cells in SDM80 medium. L262P/L262P γ cells showed an evident motility defect in comparison with cells expressing at least one WT γ allele (Fig. 3A). The average curvilinear velocities measured over the actual point-to-point route followed by the cell, i.e., the mean instant speeds, were calculated from these tracks, which confirmed that L262P/L262Py cells had a substantial progressive velocity defect manifesting between day 6 and day 9 after differentiation (Fig. 3B). The WTy add-back cell line L262P/WTy, however, had a motility similar to that of WT/ WTy and WT/L262Py cells. An overall trend of increasing motility in the other cell lines between days 6 and 9, despite a trend of slightly declining total cellular ATP in at least the parental cell line (Fig. 2B), suggests that the relationship between total cellular ATP concentration and motility is not simply linear but that there is possibly a threshold of ATP availability below which motility cannot be maintained. Consistent with the more rapid drop in cellular ATP in Tb1-/Tb1- cells, these mutants showed a pronounced reduction in mean velocity immediately after differentiation, at a time when the velocity of L262P/L262Py cells was still unaffected (Fig. 3C; Fig. S4). The rapid death of the Tb1⁻/Tb1⁻ cells after differentiation precluded an analysis of later time points.

Ability of F₁F₀-ATP synthase mutants and kDNA° cells to colonize the tsetse midgut. To test whether trypanosomes devoid of kDNA could establish a PCF midgut infection in vivo, we fed teneral tsetse flies with horse blood containing stumpy-form WT/L262Pγ kDNA^o parasites or kDNA⁺ stumpy-form parasites with the parental, WT/ WTγ, WT/L262Pγ, or L262P/L262Pγ genotype at the Liverpool School of Tropical Medicine (LSTM) tsetse fly colony. At day 9 postinfection, we dissected the flies and isolated, disrupted, and inspected their midguts under a microscope to assess the parasite infection prevalence. The parental cell line was able to differentiate to the PCF and established a midgut infection in around 85% of flies dissected (Fig. 4A), with 70% of the flies having high levels of infection. In contrast, kDNA° T. brucei cell lines were unable to establish even a single midgut infection.

Flies infected with WT/L262P γ or L262P/L262P γ cells had a slightly lower midgut infection level than flies infected with parental or WT/WT γ cells; approximately 70% were midgut infected, but the difference in overall infection rates was not statistically significant (Fig. 4A). However, flies infected with L262P/L262Pγ parasites showed a significantly lower proportion of highly infected midguts. Infections conducted at a



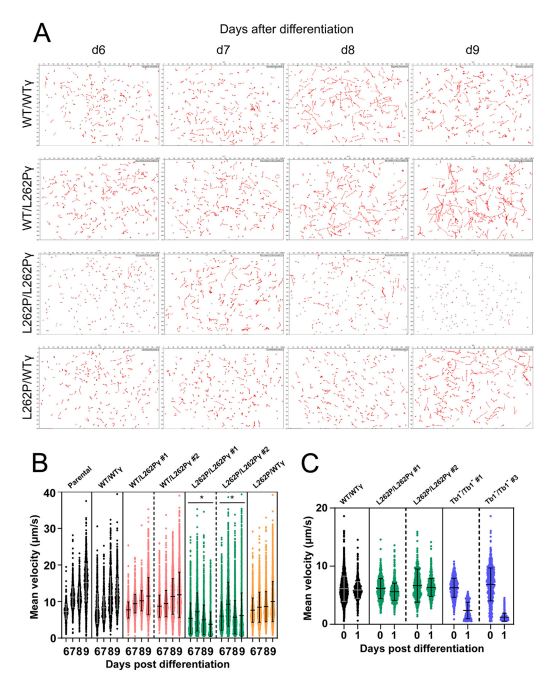


FIG 3 Motility analysis of in vitro cultured parasites. (A) Representative cell movement tracks taken from videos of differentiated PCF cells in vitro. Videos were taken of populations of newly differentiated PCF T. brucei at days 6, 7, 8, and 9 (d6 to d9) after differentiation. (B) Measurement of curvilinear velocity (VCL) from video tracking in panel A, expressed as mean velocity of each tracked cell. VCL was assessed at d6 to d9. At least 965 cells were tracked per cell line and time point. P values were determined using ANOVA between cell lines across the entire d6-d9 data set. *, P < 0.05 (significant reduction in motility in comparison to the other cell lines shown). Error bars indicate standard deviations. (C) VCL assessed as mean velocity of each tracked cell at d0 and d1. At least 750 cells were tracked per cell line and time point. Error bars indicate standard deviations. Representative tracks taken from videos of these cells are shown in Fig. S4.

second tsetse colony at Institut Pasteur, Paris, France, with a subset of the same cell lines showed more reduced levels of midgut infection for L262P/L262Py cells, and this phenotype was rescued in the WTy add-back cell line (Fig. 4B). Flies infected in a separate experiment with Tb1-/Tb1- cells produced infections that were barely detectable in the midgut (Fig. 4A), which is consistent with their rapid death after differentiation in vitro (Fig. 2B). When combined, these results demonstrate that T. brucei requires an



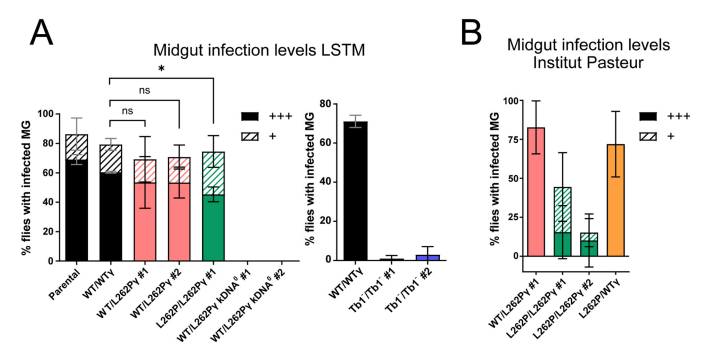


FIG 4 Assessment of fly midgut infectivity of parasites with ATP synthase and kDNA mutations. Midgut infection rates at tsetse fly facilities of (A) Liverpool School of Tropical Medicine (LSTM) and (B) Institut Pasteur Paris. Infected tsetse fly midguts (MG) were harvested at day 9 after the infected blood meal. Density of infection was judged by microscopy: +++, high level of midgut infection at >>10 parasites/field of view; +, low level of midgut infection at ~10 parasites/field of view. For the experiment with Tb1-/Tb1- cells and the corresponding WT/WTy control (A, right), midguts were scored as infected if any trypanosomes were detected. Approximately 25 flies (or 50 flies for the Tb1-/Tb1- cell lines and the corresponding control [A, right]) were infected with one blood cryostock of stumpy-form trypanosomes per cell line and replicate; n = 3. Error bars indicate standard deviations. *, P < 0.05, and ns, not significant, for comparison between the heavy infection levels (unpaired t test).

F₁F₀-ATP synthase complex capable of functioning in OXPHOS to produce an efficient tsetse midgut infection. A reduced capacity for OXPHOS, as demonstrated by the homozygous L262Py parasites, can sustain a midgut infection, but at a reduced rate. Absence of functional F_1F_o ATP synthase, as in $Tb1^-/Tb1^-$ cells, permits minimal midgut infection, whereas absence of all kDNA-encoded products completely prevents midgut infection.

Ability of L262PyF₁F_o-ATP synthase mutants to colonize the tsetse fly PV. We then assessed the ability of the heterozygous and homozygous L262Py mutants to colonize the PV and to continue their development. Whereas a high proportion of midgut infections manifested into PV infections with cells having at least one WT γ allele, infections with L262P/L262Pγ cells produced either very low numbers of (clone 1) or no (clone 2) infected PV (Fig. 5A). As predicted, adding back a WTy allele to an L262P/L262Py cell line restored a normal PV infection rate. Infections performed at the other tsetse colony also produced substantially reduced PV infection rates for L262P/L262Pγ parasites that were also rescued in a WT γ add-back cell line (Fig. 5C). Tracks recorded from cells released from the midgut showed that L262P/L262Py cells displayed significantly less motility than other cell lines within the midgut, although it was notable that there was a large proportion of cells with relatively low motility for all cell lines (Fig. 6). Of note, rescue cell line L262P/WTy showed a somewhat bimodal distribution, with an apparently distinct subpopulation of cells with higher motility. This could reflect heterogenous expression of the transgene in the parasite population. Overall, we conclude that the low PV infection rate is correlated with a decreased motility of these cells.

To assess whether capacity for OXPHOS also affected differentiation, we quantified the various parasite stages found in the PV. At the Institut Pasteur facility, we observed that cells with at least one $WT\gamma$ allele could differentiate into long mesocyclic trypomastigote forms as well as into long and short epimastigote forms (Fig. 5B). WT/L262P γ cells showed higher proportions of epimastigote stages than WT/WT γ cells; however,

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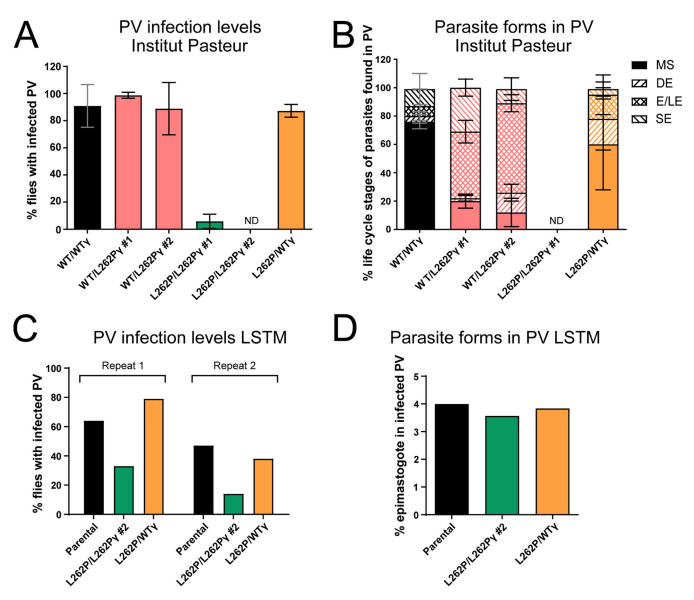


FIG 5 Assessment of fly PV infectivity of parasites with ATP synthase mutations. PV infection and parasite differentiation rates at the tsetse fly facilities of Institut Pasteur and LSTM. At both colonies, infected tsetse fly PV were harvested between 21 and 28 days after infection (A and C), and parasite life cycle stages of cells from these infected PV were blindly quantified from three fixed and DAPI (4',6-diamidino-2-phenylindole)-stained slides per cell line with approximately 100 cells per slide (B and D). (A) A total of 657 flies were dissected at Institut Pasteur (n = 3; error bars indicate standard deviations). (B) L262P/L262Py clone 1 was assessed, but no cells were detected in the PV after fixation due to the very low initial parasite density, and L262P/L262Py clone 2 was not assessed, as no infected PV were detected (see panel A). ND, none detected; MS, mesocyclic; DE, dividing epimastigote; E/LE, epimastigote/long epimastigote; SE, short epimastigote. Error bars indicate standard deviations. (C) One hundred fifty flies per repeat (50 per cell line) were dissected at LSTM.

introduction of a single WTy allele in the L262P/L262Py background (L262P/WTy) resulted in a percentage of epimastigote cells that was more similar to that in the WT/ WT γ cells. The life cycle stages of L262P/L262P γ cells found in the PV could not be assessed for experiments at this fly colony due to the extremely low level of infections found in this organ. The PV infection rate for the L262P/L262Py mutant at the LSTM fly colony allowed us to assess the proportion of epimastigote forms for these cells and for parental and WTy add-back cells as key controls. Interestingly, epimastigotes could be seen at approximately the same proportion in all these cell lines (Fig. 5D), despite the severely reduced levels of PV infection with L262P/L262Py cells. This suggests that, if able to migrate to the PV, L262P/L262P γ cells can physically progress to the epimastigote stage of the life cycle. Thus, whether a reduced capacity of these cells for OXPHOS affects the efficiency of differentiation to the epimastigote stage remains uncertain from these data.



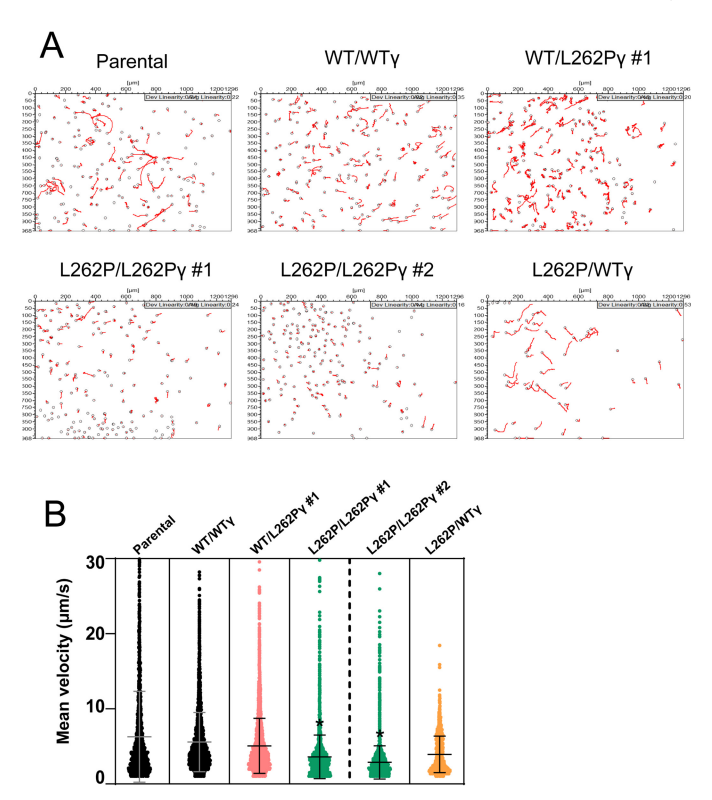


FIG 6 Motility analysis of parasites isolated from the fly midgut (ex vivo). (A) Representative tracks taken from videos of cells from tsetse fly midguts dissected at day 9 postinfection at Institut Pasteur. (B) Measurement of curvilinear velocity (VCL; calculated as mean velocity for each cell) from tracked videos of ex vivo PCF cells. P values were determined using ANOVA between cell lines. *, P < 0.05 (significant reduction in motility in comparison to the other cell lines). Error bars indicate standard deviations. At least 11 movies were taken per fly posterior midgut, with at least 3 flies dissected per strain. At least 1,546 cells were tracked per cell line.



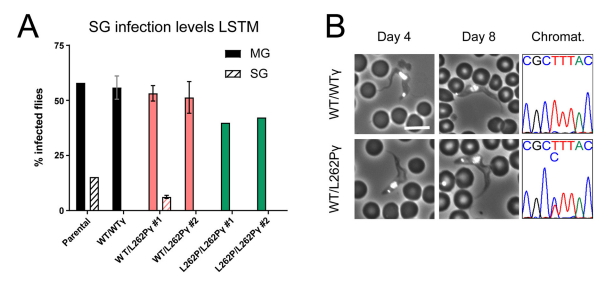


FIG 7 Assessment of fly salivary gland infectivity of parasites with ATP synthase mutations. (A) Infected tsetse fly midguts (MG) and salivary glands (SG) were harvested 4 weeks after infection. For infections with WT/WT₂, WT/L262Py clone 1, and WT/L262Py clone 2, n = 2. For the remaining infections, n = 1. Error bars indicate standard deviations. Around 200 flies were infected with one blood cryostock containing stumpy-form trypanosomes per cell line and replicate. Infections were performed at LSTM. (B) Mice were infected with metacyclic-form trypanosomes isolated from fly salivary glands from panel A. Tail snips were performed daily to assess parasitemia, and DAPI-stained blood smears were collected on days stated. Bar, 10 μ m. The sequencing chromatograms on the right confirm the heterozygous ATP synthase subunit γ genotype (CTT = leucine; CCT = proline).

Ability of L262PyF₁F₀-ATP synthase mutants to complete the life cycle. To investigate whether the life cycle could be completed by L262P γ -expressing cell lines, we dissected infected tsetse flies at 28 days postinfection. Around 12% of tsetse flies fed with parental stumpy cells had detectable parasites in their salivary glands (Fig. 7A), confirming the ability of this parental cell line to complete the *T. brucei* life cycle. WT/ L262Pγ clone 1 produced a lower salivary gland infection rate at an average of 6%. No infected salivary glands were found in flies that had been fed WT/WTy, WT/L262Py clone 2, or L262P/L262Pγ parasites, despite these flies having established midgut infections. The lack of salivary gland infections for the WT/WT γ and one of the WT/L262P γ cell lines suggests that the capacity for this step in the life cycle progression is perhaps unstable and prone to loss in experimental cell lines (although for a given cell line and experiment, salivary gland infection results are reproducible). As a consequence, the significance of the lack of infectivity of L262P/L262Pγ cells is uncertain.

To confirm that the metacyclic form trypanosomes found within the salivary glands in the parental strain and in WT/L262Py clone 1 were indeed animal transmissible, we harvested these forms and used them to infect mice. Slender BSF trypanosomes were visible in both tail blood smears from day 4, with stumpy forms being visible in both strains from day 8 (Fig. 7B). To ensure that cells of the original genotype WT/L262P γ had not reverted to the WT/WTγ genotype, we confirmed by sequencing that both WT and L262Py alleles were detectable at the DNA level. Hence, T. brucei organisms with a WT/L262P γ genotype can complete their parasitic life cycle.

To the best of our knowledge, this is the first published example of a comparison of infection results from two distinct tsetse fly colonies. However, not all cell lines were challenged each time due to fly number restraints and to the extremely laborious nature of the experimental tasks. In total, result trends observed in parallel with the same strains were similar between the tsetse fly colonies at LSTM, Liverpool, and the Institut Pasteur, Paris. They differed only in the degree to which the L262P/L262P γ mutation affected midgut and PV infections. A decrease in midgut infection level was seen for the Institut Pasteur infections (Fig. 4B) compared to those performed at LSTM (Fig. 4A). This correlated with PV infection results, where the L262P/L262Pγ cells barely established a PV infection at Institut Pasteur (Fig. 5A), in contrast to LSTM, where more substantial PV infections were found, but at a lower level than those found in other cell



types (Fig. 5C). These subtle differences could be explained by the different intrinsic vector competences of the two colonies and/or by differences in the infection and maintenance protocols (infection vehicle, blood meal type, and frequency).

DISCUSSION

The kinetoplast (or kDNA) of the sleeping sickness parasite T. brucei encodes essential subunits of respiratory chain complexes, the F₁F₀ ATP synthase and the mitoribosome, which is a property it has in common with mitochondrial DNA of other eukaryotes. Although mitochondrial activity shows dramatic changes during the complex life cycle of the parasite, multiple in vitro studies have confirmed that the major proliferative stages in the mammalian bloodstream (long slender BSF) and in the tsetse fly vector midgut (PCF) depend on kDNA-encoded gene products to manufacture $\Delta\Psi$ m (BSF and PCF) and mitochondrial ATP (PCF). In the BSF, dependence on kDNA for viability can be overcome by compensatory changes in the nucleus-encoded F_1 subunit γ . We utilized a panel of BSF mutants with various defects in kinetoplast-encoded functions (Table 1) combined with a number of multiple experimental challenges to address whether the compensatory mutation that functions in the BSF can compensate for kDNA loss in PCF and to what extent capacity for ATP production by OXPHOS is required for PCF differentiation and viability in vivo, as well as for cyclical development in vivo in the tsetse fly.

Parasites heterozygous for the compensatory subunity mutation can complete the life cycle. A cell line heterozygous for the mutation that enables viability of kDNA^o BSF parasites, WT/L262P γ clone 1, was able to complete the life cycle within the fly and produce animal-infective metacyclic parasites. Growth, capacity for OXPHOS and motility in vitro, and midgut infection prevalence all were unimpaired by the WT/L262Py genotype. Although WT/L262Pγ parasites displayed a higher rate of epimastigote-form infection in the PV than WTy parasites, this observation was probably not linked to the heterozygous genotype itself, as another heterozygous cell line, the WTγ add-back cell line (L262P/WT₇), showed levels of epimastigote forms that were more similar to those in the WT γ cell lines.

Approximately 25% of flies with parental strain midgut infections developed salivary gland infections (or \sim 12.5% of the total number of flies fed with parasites), which is consistent with infection levels with this strain reported elsewhere (111). In flies infected with WT/L262Py clone 1 parasites, approximately 10% of flies with a midgut infection developed a salivary gland infection. However, as only the parental line and one WT/L262P γ clone were able to differentiate into the metacyclic stage, and as the ability to invade the salivary glands seems to be a fragile trait that can be lost in laboratory-adapted cell lines, it cannot be ruled out that these differences in infection rate could be due to clone-specific differences unrelated to the genetic manipulation. Nonetheless, our results clearly demonstrate that parasites with heterozygous L262Py mutations enabling kDNA independence can be transmitted by tsetse flies. T. brucei parasites with this or other mutations that obviate kDNA in BSFs are much less sensitive to classes of drugs used in the field to treat animal disease, including isometamidium and homidium, since these drugs, at least in part, target parasite kDNA (112). Exposure to drugs that target kDNA, however, would cause loss of kDNA from WT/L262P γ cells, thus preventing transmission by tsetse flies, as confirmed in the present study (see below). Hence, although a drug-resistant phenotype caused by a heterozygous subunit γ genotype could initially be spread by cyclical transmission, once in a host treated with these drugs, parasites would continue to proliferate, but the opportunity for further tsetse transmission from that host would stop. If such mutations were selected for in an animal-infectious strain of Trypanosoma brucei brucei, there would also be scope for sexually recombining with human-infective parasites to produce parasites with increased drug resistance able to infect humans (113, 114).

Requirement for F_1F_0 -ATP synthase complexes through the life cycle. A comparison of the phenotypes observed for cells with a homozygous L262P F_1 subunit γ mutation versus null mutants for Fo subunit Tb1 provided insight into the requirement for $\rm F_1F_o$ complexes for life cycle completion. First, L262P/L262P γ cells (unlike WT/WT γ and



WT/L262Pγ cells) grow poorly under glucose-depleted conditions, which increase dependence on OXPHOS (28, 37). Tb1-/Tb1- cells, on the other hand, die rapidly under these conditions. Second, early in differentiation to PCF in vitro, L262P/L262P γ cells maintain higher cellular ATP concentrations than Tb1 $^-$ /Tb1 $^-$ cells. Third, L262P/L262P γ cells sustain midgut populations in the glucose-depleted tsetse fly midgut, although these are less dense, and cells are less motile by day 9 than in flies infected with WT/ WT γ or WT/L262P γ cell lines. Tb1 $^-$ /Tb1 $^-$ cells, in contrast, were unable to establish or sustain midgut infections. Together, these results demonstrate that, first, differentiation to PCF and tsetse infectivity depends on functional F₁F_o ATP synthase complexes, which is consistent with the observation that stumpy parasites treated with the F₁F₀ inhibitor oligomycin were unable to reenter the cell cycle during differentiation to PCF cells (68). Second, the L262P γ mutation causes a less profound defect in F_1F_0 -ATP synthase function than the loss of Tb1. This suggests that, whereas the F_1F_o -ATP synthase is nearly completely disrupted in the absence of Tb1 (107, 109), the F_1F_0 complex retains some functionality in L262P/L262Py cells, as these cells can still generate enough ATP mitochondrially to survive under these conditions.

In principle, the effect of the mutations on mitochondrial ATP production could be direct (via F₁F_o-ATP synthase participation in OXPHOS), indirect (via an impact on substrate level phosphorylation), or a combination of both. Crude mitochondrial extracts from L262P/L262Py cells barely produce ATP via OXPHOS, with ATP production levels close to those in the presence of an ATP synthase inhibitor. This is consistent with the expected partial uncoupling between F₁ and F₀ that is caused by the disrupted interaction between the mutated γ subunit and F_1 headpiece (84–87). On the other hand, like stumpy cells (24, 115, 116), newly differentiated L262P/L262Pγ PCF cells are able to metabolize α -ketoglutarate via mitochondrial substrate level phosphorylation to generate ATP. Indeed, we found that L262P/L262P γ clone 2 cells produce ATP from α -ketoglutarate to an extent similar to that of WTy cells. As L262P/L262Py cells can establish infection in the tsetse midgut and progress to the PV, we postulate that the capacity for substrate-level phosphorylation in the absence of glucose may be higher in vivo than in vitro; threonine or glutamine concentrations, for example, may be higher per trypanosome in the midgut (117, 118). The presence of symbiotic bacteria in the tsetse fly midgut and the influence of fly digestion may also impact the metabolism of midgut-associated parasites (35, 119, 120). Nutrients may also be depleted faster within liquid culture than in the fly midgut.

Why would L262P/L262P γ cells and Tb1⁻/Tb1⁻ cells differ in their capacity for mitochondrial ATP production via substrate-level phosphorylation? In L262P/L262Pγ PCF cells, $\Delta \Psi m$ can be generated by electron transport chain components, namely, respiratory complexes III and IV (44, 45). In these cells, the partial uncoupling between the F₁ and F_o parts would be expected to generate a proton leak through F_o into the matrix (84-87). However, the viability of these PCF cells and the fact that they can produce ATP in the presence of α -ketoglutarate suggest that a $\Delta \Psi m$ of sufficient magnitude is maintained to support metabolite transport and mitochondrial protein import for substrate-level phosphorylation. In Tb1-/Tb1- cells, however, the nearly complete loss of Fo from ATP synthase complexes (107, 109) might cause sustained hyperpolarization of the inner mitochondrial membrane, as the $\Delta\Psi m$ generated by complex III and IV cannot be dissipated by proton movement through the F_o moiety (121-123). Hyperpolarization would interfere with metabolite exchange and mitochondrial protein import and cause reactive oxygen species (ROS) generation and enhanced oxidative stress, thus damaging cells in a multitude of ways (124). Indeed, persistent hyperpolarization is an acknowledged cell fate and death checkpoint (125-128). In addition, the profound inability of Tb1-/Tb1- cells to survive as PCF in vitro or in vivo may be influenced by the substantial remodeling of the mitochondrion required during BSF-to-PCF differentiation (9). During this differentiation, the tubular BSF mitochondrion expands in volume and becomes elaborated into a branched network structure with abundant cristae (15, 115, 129), which are important for efficient OXPHOS (130–132). Dimerization of the trypanosome F_1F_0



complex may be critical for folding of the inner mitochondrial membrane and cristae formation and may critically involve F_o, as in other organisms (133–138). The absence of Tb1 and, consequently, Fo in these cells could interfere with the remodeling of the inner mitochondrial membrane during differentiation, preventing the ordered processes required from being completed. The mutants with progressive developmental defects generated here therefore will be a useful resource for future studies aimed at correlating kDNAencoded functions with the changes in mitochondrial physiology and ultrastructure that are a hallmark of trypanosome differentiation.

Although the reduced ATP production levels that we observed for L262P/L262Py cells are sufficient to permit colonization of the midgut, they are correlated with a progressive motility defect that appears to decrease the migration efficiency of the cells in the fly. Only a low level of PV infection was detected in flies infected with L262P/L262Py cells. Swimming ability has been shown to be dictated by the availability of ATP for the dynein motor in numerous eukaryotic flagellate organisms (139-143). In T. brucei, a dynein motor mutant has a motility defect in culture and consequently is unable to migrate to the foregut and PV or infect the salivary glands (144). Furthermore, efficient ATP generation is required for motility of *T. brucei* (145). The ability to efficiently metabolize proline and alternative carbon sources may be particularly important as the parasite tries to migrate to compartments in the tsetse fly that could be less nutrient rich than the midgut, especially at the end of the digestive process (34, 35). The lower infection level found in the L262P/L262Py-infected midguts may also play a role in reduced migration as per the bottleneck model of *T. brucei* differentiation and migration in the tsetse fly (55). In our ex vivo motility analysis, we noticed a pronounced heterogeneity among parasite populations for all cell lines, including the parental line, which could be a contributing factor to the bottleneck effect.

In addition to motility, substantial amounts of ATP are also required for cell cycle progression and differentiation (146). DNA replication complexes and kinesin proteins governing the vast morphological and organelle rearrangements depend on translocations fueled by ATP (147-150). Indeed, the progressive cellular ATP decline over time seen in L262P/L262P γ cells is probably due to the large ATP expenditure necessary for motility and cell cycle progression, and the inefficiency of ATP generation in this cell line. In vivo, L262P/L262Py cells can differentiate into epimastigotes in a limited fashion, agreeing with recent evidence that OXPHOS may not be essential for this life cycle stage (35). The presence of L262P/L262Py epimastigotes was unexpected considering that developmental progression in the insect may be driven by ROS production (63) and that the presence of a L262Py-induced proton leak would be hypothesized to neutralize ROS generation (151).

L262P/L262Pγ cell lines could not infect tsetse salivary glands despite these cell lines being able to sustain a midgut infection for over 4 weeks in these flies. The significance of this observation is uncertain, as it is common to observe a loss of fly infectivity in cultured *T. brucei* strains (152), as observed here for WT/WTy and WT/L262Py clone 2 cell lines. As all cells had been maintained previously in BSF culture, it is possible that there was selection for parasites that had lost genes required for progression from the PCF in the tsetse midaut.

The requirement for kDNA. Stumpy kDNA^o cells were consistently unable to (i) establish a midgut infection and (ii) differentiate into viable PCF cells in vitro after induction with CA, dying within 24 h. The kDNA^o stumpy cells express PAD1 (24) and do have a functional signaling pathway for differentiation, as they respond to CA by expressing EP procyclin at a time point comparable to that seen with kDNA+ cell lines, as reported here and elsewhere (99, 102). However, kDNA^o parasites do not effectively reenter the cell cycle after differentiation from the cell cycle-arrested stumpy BSF. EP procyclin expression appears to be independent of kDNA, but effective reentry into the cell cycle seems to require either the presence of the kinetoplast structure or a kDNA-encoded function, which is in agreement with the work of Timms et al. (68). We suggest that this is due to a complete failure to generate a $\Delta\Psi$ m. In the presence of



functional kDNA, both the slender and stumpy BSFs generate $\Delta \Psi m$ by operating the F_1F_0 -ATP synthase in the reverse direction (17–19, 24, 153). The enzyme switches to the forward direction during stumpy to PCF differentiation (24, 154), and $\Delta\Psi m$ is then generated in a canonical fashion by respiratory complexes III and IV. According to the current model, the kDNA 0 long slender BSF can maintain $\Delta\Psi$ m independently of F_o via the electrogenic action of AAC-mediated ATP⁴⁻/ADP³⁻ exchange across the inner mitochondrial membrane when supported by efficient ATP hydrolysis via an F₁ enzyme bearing a compensatory mutation in subunit γ (17, 24, 66, 82, 83). However, $\Delta\Psi m$ is not generated in kDNA^o stumpy cells, probably due to mitochondrial substrate-level ATP production using α -ketoglutarate preventing the electrogenic import of cytosolic ATP by the AAC (24). In the absence of kDNA-encoded complex III and IV, kDNA⁰ PCF would be unable to generate $\Delta\Psi m$ for the same reason. We can conclude that all naturally occurring kDNA⁻ or kDNA⁰ variants of *T. brucei*, such as *T. b. evansi* and *T. b. equiperdum*, are intrinsically unable to undergo cyclical development in the tsetse fly.

MATERIALS AND METHODS

Cell line generation. All cell lines used in this study were derived from culture-adapted pleomorphic T. brucei EATRO 1125 AnTat1.1 90:13 BSF parasites (Table S1) (99). Heterozygous cell lines with one F_1F_0 -ATPase subunit γ allele (systematic TriTrypDB ID Tb927.10.180) with the L262P mutation (L262P γ) and akinetoplastic (AK) versions of these cell lines were generated as detailed in reference 24. Homozygous L262Py cell lines were generated by transfecting WT/L262Py T. brucei with plasmid pEnT6-yL262P-BSD (blasticidin resistance marker) to replace the remaining endogenous WTγ allele with an L262Pγ copy. The WTγ add-back cell line was generated by releasing L262P/L262PγT. brucei cells from either puromycin or blasticidin selection before the transfection and subsequently transfecting them with plasmid pEnT6-yWT-PHL (phleomycin resistance marker). Both of these plasmids are based on the pEnT6 backbone (155) and contain either a L262Py gene or a wild-type version (WTy), facilitating the replacement of an ATPase γ subunit allele. The replaced gene is expressed by read-through transcription of the endogenous locus and contains its native 5' untranslated region (UTR) but the aldolase 3' UTR.

To generate a Tb1 (Tb927.10.520)-null (Tb1⁻/Tb1⁻) cell line, pUC57-based plasmids containing either the blasticidin or the phleomycin resistance cassettes were designed and ordered from Biomatik (Fig. S2). Actin 5' and 3' UTRs flank the resistance cassettes, with 350 bp of the Tb1 5' UTR and intergenic region (IGR) positioned upstream of this and 350 bp of the Tb1 3' UTR and IGR positioned downstream of this. The first allele knockout (KO) construct was designed to target sequences distal to those employed by the second allele KO construct (156). These KO constructs were positioned inside HindIII restriction sites to enable digestion before transfection.

For the transfection, the Amaxa Nucleofector II (Lonza) was used with nucleofection solution (90 mM NaH₂PO₄, 5 mM KCl, 0.15 M CaCl₂, 50 mM HEPES [pH 7.3]) and program Z-001. *T. brucei* EATRO 1125 AnTat1.1 90:13 BSF clones were selected after 4 days at 37°C under drug selection as necessary (2.5 $\mu g/$ mL G418, 5 μ g/mL hygromycin, 0.1 μ g/mL puromycin, 5 μ g/mL blasticidin, 5 μ g/mL phleomycin) in HMI-9 medium (157) containing 10% (vol/vol) fetal calf serum (FCS; Gibco). ATPase γ genotypes were verified as detailed by Dewar et al. (24). Tb1 KO cell lines were verified by PCR and Western blotting.

In vitro differentiation and cell culture. Stumpy forms were generated in mice and were either used to make cryostocks containing a parasite population with approximately 90% stumpy bloodstream forms or purified from blood (24), as required. In vitro differentiation was performed by adding 6 mM cisaconitate (CA) to each culture, and cultures were left at 27°C for 24 h. If samples contained blood cells, the flasks were balanced on one corner to allow blood cells to settle out of the medium. Newly differentiated PCF cells were washed and resuspended either in SDM79 containing 10 mM glucose (Invitrogen) (158) or in SDM80 at a density of at least 2×10^6 cells per mL medium. SDM80 was made from SDM79 CGGGPPTA powder (GE Healthcare), an SDM79-based powder that lacks major carbon-containing components (sodium bicarbonate, glucose, glutamine, glutamate, proline, pyruvate, threonine, and sodium acetate). All carbon sources except glucose were added back into the solution at the following concentrations: pyruvate, 100 mg/L; L-proline, 615 mg/L; L-threonine, 394 mg/L; L-glutamine, 320 mg/L; L-glutamate, 24 mg/L; NaHCO₃ 2 g/L; sodium acetate, 10 mg/L. The medium was supplemented with 50 mM Nacetyl D-glucosamine (Sigma) to prevent residual glucose uptake from FCS (159) and maintained under drug selection as necessary (15 μ g/mL G418, 25 μ g/mL hygromycin, 2 μ g/mL puromycin, 10 μ g/mL blasticidin, 2.5 μ g/mL phleomycin). SDM79 and SDM80 were supplemented with 7.5 mg/L hemin and 10% FCS (Invitrogen), and cells were grown at 27°C continuously in these media, maintaining density over 2×10^6 /mL. For time courses, day 0 (d0) was defined as being after 24 h in HMI-9 at 27°C plus CA and then 24 h in SDM80 at 27°C.

Protein expression analysis by flow cytometry and Western blotting. For a quantitative measurement of EP expression, 2×10^6 cells were harvested from +CA and -CA cultures at 0 h, 1 h, 2 h, 3 h, 6 h, 16 h, and 24 h after CA addition. Upon collection, samples were transferred to 5-mL polystyrene round-bottom tubes (BD Flacon 352052), centrifuged at 2,000 \times g for 5 min, and washed in phosphate-buffered saline (PBS; pH 7.4; 137 mM NaCl, 2.7 mM KCl, 10 mM Na_2HPO_4 , 1.8 mM KH_2PO_4). The cell pellet was fixed in 500 μL PBS with 2% formaldehyde and 0.05% (wt/vol) glutaraldehyde overnight. Cells were pelleted, washed three times in PBS, and blocked in PBS with 2% (wt/vol) bovine serum albumin (BSA) for 1 h. After



a PBS wash, cells were stained with PBS containing 2% BSA and 1/500 anti-EP procyclin monoclonal antibody (Cedarlane Laboratories) for 1 h. Cells were washed in PBS, and then PBS containing 2% BSA and 1/1,000 anti-mouse IgG-Alexa 488 secondary antibody was added and left for 1 h. Cells were then washed and resuspended in 500 μ L PBS containing 5 μ g/mL Hoechst DNA staining dye (Life Technologies). After a 30-min incubation at room temperature, cells were analyzed by flow cytometry at an excitation wavelength (λ_{ew}) of 495 nm and an emission wavelength (λ_{ew}) of 519 nm using a Becton Dickinson LSR II machine with BD FACSDiva software, and 2×10^4 events per sample were measured. Results were analyzed with FlowJo software (BD Biosciences).

Western blotting was performed as per Dewar et al. (24). Antibodies used were anti-EP procyclin (1:500), anti-ATP synthase Tb1 (1:2,000) (160), anti-ATP synthase Tb2 (1:2,000) (160), anti-ATP synthase β (1:2,000) (160), anti-COXIV (1:1,000) (41), anti-COXVI (1:500) (44), anti-EF1 α (1:7,000) (Millipore), and anti-COXVI (1:00) mtHSP70 (1:2,000) (161).

Tsetse fly handling and infections. To make the infective parasite stabilates, mouse blood containing stumpy-form trypanosomes was harvested, mixed with 2% (wt/vol) sodium citrate to act as an anticoagulant, and mixed 1:1 with HMI-9 containing 30% (vol/vol) FCS and 7.5% (vol/vol) glycerol for freezing at -80° C. Blood meals for tsetse fly infections were prepared as follows: one 200- μ L aliquot per cell line was thawed at room temperature and mixed with ~5 mL room-temperature, sterile, defibrinated horse blood at the LSTM or with $\sim\!2$ mL room temperature heat-inactivated fetal calf serum at the Institut Pasteur.

At the LSTM, the Glossina morsitans morsitans (Westwood; origin, Kenya) colony was maintained at $26\pm2^{\circ}$ C and 68 to 78% relative humidity, with a 12-h light/12-h dark cycle, and was fed triweekly on sterile, defibrinated horse blood in the dark using a sterile silicon membrane and a heated mat set to 37°C. All flies used in this study were newly emerged, teneral (unfed) male adults. Each experimental group was fed one infected blood meal containing predominantly stumpy-form parasites when flies were at 0 to 24 h after emergence. To obtain midgut infections, flies were offered a parasite-infected blood meal for 10 min, and then 24 h after feeding, flies were chilled to 4°C to remove unfed flies (identified by a nonscarlet abdomen). Flies with a visible blood meal were maintained at 27°C until dissection on day 9 postinfection and fed uninfected blood meals as described above every 2 to 3 days. For salivary gland infections, flies were allowed to feed on an infected blood meal for 15 min. Flies were chilled to 4°C at 24 h after the blood meal to remove unfed flies. Flies were maintained as described above and dissected 4 weeks postinfection.

At the Institut Pasteur, G. m. morsitans tsetse flies (Westwood; origins, Zimbabwe and Burkina Faso) were maintained in Roubaud cages at 27 \pm 1°C and 70% \pm 5% relative humidity, with a 12-h light/12-h dark cycle, and fed twice a week through a silicone membrane with sterile, mechanically defibrinated sheep blood at 37°C. Teneral males (between 24 h and 72 h postemergence) were allowed to ingest BSF parasites in heat-inactivated fetal calf serum at 37°C through a silicone membrane for 10 min during their first meal, and unfed flies were removed after feeding. When possible, flies were starved for at least 24 h before being dissected. For dissection, midguts and PVs were rapidly isolated and placed in distinct drops of PBS with 0.1% glucose as previously described (54, 61). Isolated tissues were assessed by microscopy $(40\times)$ and imaged or allowed to infuse for 5 min in a wet chamber before removal and collection of the remaining PBS drop containing trypanosomes for video recording (tracking analyses; see below) (162).

To infect mice with metacyclic parasites harvested from tsetse flies, infected salivary glands were collected and pooled in ice-cold SDM79. Frozen stocks of these salivarian parasites were prepared by adding glycerol to a final concentration of 7.5% (vol/vol) before freezing at -80° C. Mouse infections, blood smears, and harvesting were performed as detailed in reference 24.

Motility tracking and analysis (in vitro and ex vivo). In silico 2D tracking was performed as previously described (144). Briefly, PCF trypanosomes were sampled from cultures or freshly isolated from tsetse fly tissues, suspended on noncoated glass slides in a drop of PBS with 0.1% (wt/vol) glucose, and maintained at 27°C. Parasites were observed under the 10× objective of an inverted DMI-4000B microscope (Leica) coupled to an ORCA-03G (Hamamatsu) or a PRIM95B (Photometrics) camera. Video recording was performed within 15 min after isolation, and the focal plane was set up in the middle of the PBS drop containing parasites in order to prevent the recording of attached cells at the base of the drop or drifting cells at the top of the drop. For each cell line, 10 to 20 movies were recorded for 20 s (50 ms of exposure). Movies were converted with the MPEG Streamclip V.1.9b3 software (Squared 5) and analyzed with the medeaLAB CASA Tracking V.5.5 software (medea AV GmbH) for quantifying the mean velocity of 200 individual trypanosomes per movie over 149 successive frames. Statistical analyses and plots were performed with XLSTAT 2019.2.01 (Addinsoft) in Excel 2016 (Microsoft) or Prism V8.2.1 (GraphPad). Statistical analyses include two-sided analyses of variance (ANOVA) with Tukey's ad hoc posttests for intergroup comparison at 95% confidence.

In organello ATP production and whole-cell ATP quantification assays. The ATP quantification assay was performed as per the protocol for the CellTiter-Glo 3D cell viability assay kit (Promega), using 1×10^4 cells per sample. In organello ATP production assays were performed using the CLSII ATP bioluminescence assay kit (Roche) and digitonin extractions of 1×10^7 cells per sample (110). Statistical analyses involved one-sided ANOVA with Tukey's ad hoc posttests for intergroup comparison at 95% confidence.

SUPPLEMENTAL MATERIAL

Supplemental material is available online only. MOVIE \$1, AVI file, 0.6 MB.



MOVIE S2, AVI file, 1 MB.

MOVIE S3, AVI file, 0.6 MB.

MOVIE S4, AVI file, 1 MB.

FIG S1, PDF file, 0.7 MB.

FIG S2, PDF file, 0.9 MB.

FIG S3, PDF file, 0.8 MB.

FIG S4, PDF file, 1.2 MB.

TABLE \$1, PDF file, 0.4 MB.

TABLE S2, PDF file, 0.04 MB.

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