The mitochondrial ribosomal protein L13 is critical for the structural and functional integrity of the mitochondrion in Plasmodium falciparum

The phylum Apicomplexa contains a group of protozoa causing diseases in humans and livestock. Plasmodium spp., the causative agent of malaria, contains a mitochondrion that is very divergent from that of their hosts. The malarial mitochondrion is a clinically validated target for the antimalarial drug atovaquone, which specifically blocks the electron transfer activity of the bc1 complex of the mitochondrial electron transport chain (mtETC). Most mtETC proteins are nuclear-encoded and imported from the cytosol, but three key protein subunits are encoded in the Plasmodium mitochondrial genome: cyt b, COXI, and COXIII. They are translated inside the mitochondrion by mitochondrial ribosomes (mitoribosomes). Here, we characterize the function of one large mitoribosomal protein in Plasmodium falciparum, PfmRPL13. We found that PfmRPL13 localizes to the parasite mitochondrion and is refractory to genetic knockout. Ablation of PfmRPL13 using a conditional knockdown system (TetR-DOZI-aptamer) caused a series of adverse events in the parasite, including mtETC deficiency, loss of mitochondrial membrane potential (Δψm), and death. The PfmRPL13 knockdown parasite also became hypersensitive to proguanil, a drug proposed to target an alternative process for PfmRPL13 knockdown parasite also became hypersensitive to proguanil, a drug proposed to target an alternative process for maintaining Δψm. Surprisingly, transmission EM revealed that PfmRPL13 disruption also resulted in an unusually elongated and branched mitochondrion. The growth arrest of the knockdown parasite could be rescued with a second copy of PfmRPL13, but not by supplementation with decylubiquinone or addition of a yeast dihydroorotate dehydrogenase gene. In summary, we provide first and direct evidence that mitoribosomes are essential for malaria parasites to maintain the structural and functional integrity of the mitochondrion.

Malaria remains a severe infectious disease in the tropical and subtropical regions of the world, causing millions of clinical cases and taking the lives of half a million people in 2016 (1).

The causative agents, parasites of Plasmodium spp., have three genomes localized in three distinct cellular compartments, the nucleus, the apicoplast, and the mitochondrion. The nuclear genome is composed of 14 chromosomes (23 mega base pairs) and encodes ~5500 proteins (2). The apicoplast is a relic plastid derived from primary and secondary endosymbiosis, which contains a circular bacterial like DNA (35 kb), encoding ~30 proteins, 2 rRNA genes (16S and 23S), and a complete set of tRNAs required for protein translation within this organelle (3, 4).

The mitochondria of all eu- karyotic organisms are believed to be derived from one single common ancestor, an α-proteobacterium (5). In apicomplexan parasites, the mitochondrial genomes tend to be small, 6–11 kb in length (6, 7). In one genus, Cryptosporidium, the mitochondrial genome has been entirely lost (8). The mitochondrial DNA (mtDNA) of malaria parasites is only 6 kb, encoding three proteins of the mitochondrial electron transport chain (mtETC): cytochrome (cyt) b, cyt c oxidase subunits I and III (COX I and III) (6, 9, 10). These three proteins are translated by the mitochondrial ribosomes (mitoribosomes). The 6 kb mtDNA also encodes mitochondrial rRNA genes; however, these rRNAs are highly fragmented (~30 small pieces of 20–200 bp) and scattered throughout the mitochondrial genome (11–13). It remains entirely unknown how these many small rRNA pieces come together to form a working mitoribosomal complex. Another striking feature of the malarial mtDNA is that it does not encode any tRNA genes, and mitochondrial tRNAs have to be imported from the cytosol (14).

To translate genes encoded on three distinctive genomes localized in three cellular compartments, Plasmodium parasites utilize three types of ribosomes. The cytosolic 80S ribosomes are numerous in the parasite cytoplasm and were easily detected by transmission EM studies carried out more than 50 years ago (15). Their structure has just been resolved by cryo–electron microscopy (cryo-EM) (16, 17). The ribosomes in the apicoplast and mitochondrion of malaria parasites are both prokaryotic type; yet, they possess unique features that are distinct from bacterial ribosomes or organelar ribosomes of other...
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eukaryotes (18). It has been shown that active protein translation occurs in the apicoplast, which renders malaria parasites sensitive to several antibiotics (19–22). In contrast, the mitochondrial protein translation system in Plasmodium has remained an enigma for a long time. On one hand, direct evidence for mitochondrial protein translation in the parasites or in the entire Apicomplexa phylum is lacking, but on the other hand, the essentiality of the mtETC (23, 24) and the appearance of cyt b mutations conferring atovaquone resistance (25) strongly imply that the mitochondrial ribosomes are active.

In this study, we have characterized one conserved large subunit (LSU) protein of mitoribosomes in Plasmodium falciparum (PfmRPL13, PF3D7_0214200). It has been shown that L13 is a critical component involved in the first step of ribosomal LSU assembly in bacteria (26) and is one of the eight LSU subunits required to form a minimal subribosomal particle maintaining peptidyltransferase activity in Thermus aquaticus (27). Here, for the first time, we provide direct evidence that PfmRPL13 is essential for malaria parasites, highlighting the significance of mitoribosomes for parasite physiology and survival.

Results

PfmRPL13 is essential in asexual blood stages of P. falciparum

PF3D7_0214200 (designated as PfmRPL13) is annotated as the putative mitochondrial ribosomal protein L13 in P. falciparum (www.plasmodb.org) (47). To compare the sequence similarity (or divergence) of mitochondrial ribosomal L13 proteins from various organisms, we performed a multiple sequence alignment and maximum likelihood phylogenetic analysis. As shown in Fig. S1, PfmRPL13 is closely related to its orthologues in other apicomplexan parasites, but shares much less sequence similarity to other mitoribosomal L13 proteins. PfmRPL13 and Toxoplasma mRPL13 share 55% identical residues, but the percent identity between the P. falciparum and human proteins is 23%, as calculated from the alignment (Fig. S1A). As might be expected then, in the phylogenetic results the apicomplexan mRPL13s form a separate, well-supported clade within the overall mitochondrial L13 grouping (Fig. S1B).

To verify the subcellular localization of PfmRPL13 in P. falciparum, we integrated a copy of PfmRPL13 tagged with triple HA (3HA) or GFP into the genome of Dd2attB parasites (28). Interestingly, Dd2attB transfected with the pLN-PfmRPL13-GFP construct grew very slowly and did not express GFP, as determined by fluorescence microscopy and Western blotting (data not shown), suggesting that expression of the PfmRPL13-GFP fusion protein was silenced by some unknown mechanism(s). However, Dd2attB transfected with the pLN-PfmRPL13-3HA construct expressed PfmRPL13-3HA. Immunofluorescence assay showed that PfmRPL13-3HA was localized in the parasite mitochondria almost with no signal in other cellular compartments (Fig. 1). Importantly, Dd2attB-PfmRPL13-3HA grew normally as compared with the parental line (data not shown). Based on its evolutionary sequence conservation (29) and mitochondrial localization, we conclude that PF3D7_0214200 is very likely to be the mitoribosomal large subunit protein L13 in P. falciparum.

To verify whether PfmRPL13 is essential for parasite growth in asexual blood stages, we attempted to knock it out in two WT lines, D10 and NF54, via a CRISPR/Cas9-mediated approach (see “Materials and methods” in the supporting information). Transfections in these lines were performed with circular or linearized template plasmids along with pUF1-Cas9 and selected with either a single drug for the template plasmid or two drugs for both the template and Cas9 plasmids (“Materials and methods”). In cultures transfected with a circular template plasmid, negative selection against maintenance of the plasmid was also performed (“Materials and methods”). However, despite multiple attempts, gene knockout was unsuccessful (data not shown). Because it has been shown that the only essential role of the mtETC in asexual blood stages is to provide an electron sink for the parasite dihydroorotate dehydrogenase (DHODH), which is essential for pyrimidine biosynthesis (23), we hypothesized that PfmRPL13 might be dispensable in the yeast DHODH (yDHODH) transgenic parasites, which do not require a functional mtETC. Knockout approaches as described above were carried out in the NF54attB-yDHODH-GFP line which carries a fusion protein of yDHODH and GFP. Gene knockout in this line was not successful after many attempts, however. In summary, PfmRPL13 is refractory to gene disruption, suggesting that PfmRPL13 is likely to be essential.

Because our knockout studies were unable to definitely determine the essentiality of PfmRPL13 in the parasites, we utilized a recently developed translational knockdown approach, the TetR-DOZI-aptamer system (30). In this system, protein translation is conditionally regulated by addition of anhydrotetracycline (aTc) (ON) or withdrawal of aTc (OFF). As shown in Fig. 2A, the TetR-DOZI-aptamer system was inserted into the genomic locus by double crossover recombination facilitated by CRISPR/Cas9 (“Materials and methods”). We transfected D10 WT parasites with a linearized template vector and two circular guider RNA (gRNA) constructs (“Materials and methods”), selected with blasticidin and aTc, and obtained transgenic parasites 3 weeks post electroporation. As shown in Fig. 2B, the genomic locus of PfmRPL13 was confirmed to be correctly modified in the transgenic parasites by diagnostic PCR analysis. We named this line D10-PfmRPL13-KD (knock-

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Figure 2. Genetic ablation of PfmRPL13 through the TetR-DOZI-aptamer system facilitated with CRISPR/Cas9. A, the TetR-DOZI-aptamer system (30) was inserted into the PfmRPL13 genetic locus through a double crossover recombination strategy facilitated by a CRISPR/Cas9 system. TetR-DOZI-RLuc-Bsd is the combined gene for a fusion protein of tetracycline repressor with development of zygote inhibited (DOZI), followed by Renilla luciferase and blasticidin deaminase. 8xapt stands for 8 copies of TetR-binding aptamers. 5’HR and 3’HR are upstream and downstream homologous regions of the PfmRPL13 gene. The green box next to 5’HR indicates shield mutations that keep amino acid sequences intact but change nucleotide sequences to avoid repetitive cutting by Cas9. The dark blue box next to the green box is the 3HA tag. The template plasmid was fully digested with EcoRV before electroporation. Two gRNAs (**) adjacent to the stop codon of PfmRPL13 were cloned into the pAIO plasmid individually. The black bars underneath WT and integrated gene models indicate exons of the gene whereas dashed lines indicate the introns. B, the genotype of the PfmRPL13 knockdown line was verified by diagnostic PCR analysis. D10 genomic DNA served as a control. The sequences of primers P1–P4 are shown in Table S1 and their approximate positions are marked in (A). Lanes 1, 4, and 7 are products from PCR reactions with template DNA from the D10-PfmRPL13-KD line; lanes 2, 5, and 8 with DNA from D10 WT; lanes 3, 6, and 9 are no DNA controls. In lane 7, the fragment to be amplified was too big to work in PCR (>11 kb). C, a representative growth curve of the PfmRPL13 knockdown line grown in the presence or absence of aTc (250 nM). Cultures were split 1:5 on days 2, 4, 6, and 8. Parasitemia was determined under a light microscope by counting more than 1000 red blood cells in each Giemsa-stained thin blood smear. Growth index is the cumulative -fold expansion which is the multiplication of parasitemia and split factors over the time course. A split factor is 5 in a 1:5 split. Data shown are the mean ± S.D. of biological replicates. This experiment has been repeated more than 10 times. aTc, anhydrotetracycline. D, expression of PfmRPL13–3HA in the knockdown parasites maintained with and without aTc for 8 days was examined by Western blotting. The molecular mass of PfmRPL13–3HA is 28 kDa. P. falciparum aldolase (41 kDa) served as a loading control. E, MitoTracker staining patterns of healthy versus parasites with depolarized mitochondria. F, quantification of mitochondrial staining patterns in the PfmRPL13 knockdown line maintained with or without aTc, respectively. Green bars, tubular. Red bars, diffused. Data shown are averaged from n = 3 experiments and analyzed by a Student’s t test. **, p < 0.01; *, p < 0.05.

down). It has been shown that protein knockdown efficiency in the TetR-DOZI-aptamer system is determined by aptamer copy number, but aptamers are not very stable (31). To delin-
The expression of PfmRPL13 was then controlled by addition and removal of aTc in the culture. To rule out any possibility that aTc itself would interfere with parasite growth, the WT D10 line was exposed to media supplemented with aTc (250 nM) and no aTc for 2 weeks but no differences between the two were noticed (data not shown), suggesting that aTc did not alter parasite growth. To determine the effect of knocking down PfmRPL13 on parasite survival, D10-PfmRPL13-KD parasites grown under aTc were enriched by a Percoll gradient, washed thoroughly, and exposed to medium with or without aTc (250 nM) for several intraerythrocyst developmental cycles (ICDs) (“Materials and methods”). As shown in Fig. 2C, D10-PfmRPL13-KD exhibited a severe growth arrest when aTc was removed for three cycles or more. When aTc was removed for more than four cycles, the parasitemia dropped down to a negligible level. To assess parasite growth long term, the aTc-minus culture of D10-PfmRPL13-KD was maintained up to 1 month (split weekly), and growth arrest was maintained for the entire period (data not shown). In the first two cycles post aTc removal, however, the differences between aTc-plus and aTc-minus cultures were not significant under microscopic examination. We then compared their growth rates through [3H]hypoxanthine incorporation assays by titrating aTc in a serial dilution (“Materials and methods”). As shown in Fig. S2, in the first cycle, D10-PfmRPL13-KD incorporated similar levels of [3H]hypoxanthine in low and high concentrations of aTc; however, in the second cycle, the knockdown parasite grown at the lowest aTc concentration (0.24 nM) had a 40% reduction in [3H]hypoxanthine incorporation compared with that of a high aTc (250 nM) culture. In thin blood smears, morphologically deteriorating parasites were observed when aTc was removed for two cycles and the number of morphologically unhealthy parasites increased substantially after removal of aTc for three or more cycles (Fig. S3).

To determine PfmRPL13 expression levels in the knockdown parasites, a knockdown assay was set up using the same protocol as described above. Protein samples were collected every 2 days over four ICDs and examined by Western blotting (“Materials and methods”). As shown in Fig. 2D, PfmRPL13 expression was substantially diminished after aTc was removed for just one cycle. We then qualitatively monitored \( \Delta \psi_m \) using MitoTracker in the knockdown parasites maintained with or without aTc (“Materials and methods”). As shown in Fig. 2E, a healthy mitochondrion in a late trophozoite has a tubular structure with MitoTracker staining largely confined to the organelle (Tubular, top panel), whereas a sick parasite was unable to constrain MitoTracker staining to the mitochondrion, and it diffused throughout the cytosol (Diffused, bottom panel). We quantified the percentages of Tubular (healthy) versus Diffused (sick) MitoTracker staining patterns in the PfmRPL13 knockdown parasites over four ICDs. As shown in Fig. 2F, D10-PfmRPL13-KD parasites grown with aTc continuously present maintained healthy mitochondria, with very few parasites showing a diffused pattern. However, when aTc was removed, the percentage of parasites with diffused staining in the PfmRPL13 knockdown parasites increased dramatically in the third and fourth cycles. These data suggest that PfmRPL13 and, hence, functional mitoribosomes were critical to maintain mitochondrial membrane potential and parasite health. Taken together, our data strongly imply that PfmRPL13 is essential for parasite growth and survival in the asexual blood stages of P. falciparum. In addition, our results agree with the recent genome-wide gene disruption study carried out in the rodent parasite, which revealed the essentiality of mitochondrial ribosomal protein L13 in P. berghei (32).

**Genetic ablation of PfmRPL13 leads to mtETC deficiency and proguanil hypersensitivity**

Because the mtETC is the recipient of the three proteins translated by mitoribosomes in malaria parasites, it is critical to evaluate the function of the mtETC in the PfmRPL13 knockdown parasites. To do that, we directly measured the \( bc_3 \) complex enzymatic activity in vitro by an assay measuring its ability to reduce oxidized cyt c (a model of Q cycle (33) was depicted in Fig. 3A). We isolated mitochondria from D10-PfmRPL13-KD parasites maintained under aTc continuously and in the absence of aTc for two cycles and four cycles (“Materials and methods”). As shown in Fig. 3B, when aTc was removed for two cycles, there was a ~20% reduction in the \( bc_3 \) complex enzymatic activity (\( p < 0.05 \)); however, after aTc was removed for four cycles, the \( bc_3 \) complex enzymatic activity was diminished by 70% (\( p < 0.001 \)). These data provide direct evidence that knocking down PfmRPL13 results in defects in the \( bc_3 \) complex. It has been shown that whereas yDHODH transgenic parasites are resistant to atovaquone and other \( bc_3 \) complex inhibitors, they become hypersensitive to proguanil when their mtETC is chemically inhibited (23). To explore whether mtETC malfunction in the PfmRPL13 knockdown parasites would also make these parasites hypersensitive to proguanil, we cultured the parasites in the absence of aTc for three ICDs and performed a [3H]hypoxanthine incorporation assay. As shown in Fig. 3C, in the WT D10 line, there was no change in proguanil EC_{50} in the presence or absence of aTc (250 nM), suggesting that aTc did not affect proguanil sensitivity. However, in the knockdown parasite, there was a 55-fold increase in proguanil sensitivity when PfmRPL13 was genetically abolished. The EC_{50} of proguanil in D10-PfmRPL13-KD parasites was 9.56 \( \mu \text{M} \) with aTc present, whereas it was reduced to 0.17 \( \mu \text{M} \) when aTc had been previously removed for three cycles. The yDHODH transgenic line was used as a positive control in the assay (23). As expected, the yDHODH line exhibited proguanil hypersensitivity when the parasite was added with 50 \( \mu \text{M} \) atovaquone (Fig. 3C). Remarkably, the level of proguanil hypersensitivity in our PfmRPL13 knockdown parasites (55-fold) was comparable with that of the yDHODH line under atovaquone treatment (62-fold). These data suggest that PfmRPL13 genetic ablation caused a severe mtETC deficiency in a degree that was similar to that triggered by atovaquone inhibition. To rule out the possibility that hypersensitivity to proguanil in the D10-PfmRPL13-KD line was merely a result of poor parasite growth in the absence of aTc, we tested its sensitivity to other antimalarial compounds including PA21A092 (34), a PfATP4 disruptor, and artemisinin. As shown in Fig. S4, in the presence or absence of aTc, D10-PfmRPL13-KD parasites exhibited the same levels of sensitivity to PA21A092 and artemisinin, respectively. Taken together, these data strongly
suggest that proguanil hypersensitivity in the PfmRPL13 knockdown parasites was specifically induced by ablation of PfmRPL13 (see “Discussion”).

PfmRPL13 knockdown results in unusual morphological changes in the mitochondria

Next, we used transmission EM (TEM) to assess any morphological alterations in the knockdown parasites. To exclude the possibility that aTc causes changes in parasite morphology, D10 WT parasites were cultured with and without aTc for 2 weeks and their morphologies were examined by TEM (“Materials and methods”). As shown in Fig. S5, with or without aTc, D10 parasites displayed very similar mitochondrial morphologies. Mitochondria in both samples appeared to be circular structures in TEM 2D sections. Because each malaria parasite has just one single tubular mitochondrion (35), it is more likely to obtain cross-sections (circular) than longitudinal sections (tubular) during TEM 2D sample slicing. Mitochondrial morphologies similar to those of D10 were observed in the PfmRPL13 knockdown parasites maintained continuously in medium with aTc; in each TEM section, there were, again, a small number of circular structures present (Fig. 4). In contrast, when aTc was removed for three cycles, it seemed that more mitochondrial pieces were detected on each TEM section of the knockdown parasite (Fig. 4). In addition, many of the mitochondrial pieces appeared as long tubular structures, sometimes exhibiting branching (Fig. 4). To the best of our knowledge, such elongated and branched mitochondria in trophozoite stages of P. falciparum have not been reported previously in TEM studies in the literature. In all, these data revealed interesting morphological changes in parasite mitochondria when PfmRPL13 expression was genetically diminished.

Efforts to rescue PfmRPL13 knockdown parasites grown in the absence of aTc

To gain an understanding of the mechanisms that led to parasite demise when PfmRPL13 was knocked down, we...
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...cantly after five cycles or longer without aTc (long term). These data suggested that, in the long term, yDHODH expression still failed to rescue mtETC defects resulting from PfmRPL13 ablation. We then complemented D10-PfmRPL13-KD with an episomal PfmRPL13 tagged with 3Myc, resulting in another double transgenic line, D10-PfmRPL13-KD-RL13Myc (“Materials and methods”). Expression of PfmRPL13Myc was confirmed by Western blotting in the double transgenic line maintained with or without aTc for four cycles (data not shown). As shown in Fig. 5C, D10-PfmRPL13-KD-RL13Myc grew equally well in the presence or absence of aTc for a long term. To verify aptamer copy number in the endogenous PfmRPL13 locus of the double transgenic line, D10-PfmRPL13-KD-RL13Myc, we checked its 5′ integration site by PCR and sequencing and eight aptamer copies were still intact (data not shown). Therefore, in the absence of aTc, the endogenous PfmRPL13 was not translated but the double transgenic line was sustained by the episomal expression of PfmRPL13 to maintain growth. Clearly, genetic ablation of PfmRPL13 in the parasite was only rescued with provision of complementing episomal PfmRPL13.

Discussion

With the recent decline in the efficacy of artemisinin-based combination therapies (37), there is an urgent need to discover new antimalarial drugs. The mtETC of the parasite is absolutely essential throughout its complex life cycle. The mtETC is critical both to sustain the essential pyrimidine biosynthesis pathway (23) and to maintain the mitochondrial membrane potential (38). The malarial mtETC has been the focus of many endeavors seeking novel antimalarial drugs (39), yet the structure and function of mitochondrial ribosomes (mitoribosomes), which translate critical protein subunits of the mtETC, remain entirely uncharacterized. In this study, we found that one mitochondrial ribosomal large subunit protein L13 (PfmRPL13) was localized to the mitochondrion (Fig. 1) and was essential for growth of asexual blood stage parasites (Fig. 2). Knockdown of PfmRPL13 resulted in a series of adverse events in the parasite, including loss of ΔΨm (Fig. 2F), mtETC deficiency (Fig. 3B), and hypersensitivity to proguanil (Fig. 3C). Remarkably, PfmRPL13 knockdown also led to unusual mitochondrial morphologies (Fig. 4). For the first time, we provided direct evidence that mitoribosomes are essential in asexual blood stages of malaria parasites. Our genetically tagged parasite line will also be a good tool to study structural and functional aspects of mitoribosomes in the future.

The origin of all eukaryotic mitochondria can be traced back to the integration of an α-proteobacterium into the proto-eukaryote ancestor (5, 40). From this single symbiotic event, the common ancestor of all eukaryotic organisms has evolved into the myriad extant cells and organisms with many distinct morphologies and functions that allow survival in drastically different environmental milieus. Needless to say, mitochondria are key players in this ongoing evolutionary adaptation, because they are critical for maintaining cellular bioenergetics, redox balance, signaling, and even life and death decisions (41). However, in apicomplexan parasites, which cause numerous diseases in human and livestock, the mitochondrion has been streamlined by loss of many functions that are important in

attempted to rescue the knockdown parasites with three different approaches. Previous studies in P. falciparum have shown that mtETC inhibition by atovaquone can be partially rescued with addition of ubiquinone analogs such as decylubiquinone (Qd) (36). As shown in Fig. 5A, however, addition of Qd (50 μM) failed to restore growth of D10-PfmRPL13-KD parasites when aTc had been removed, suggesting that mtETC defects caused by PfmRPL13 knockdown could not be chemically rescued by Qd. Because inhibition of the mtETC by atovaquone can be fully suppressed by expression of a yeast DHODH gene (23), we hypothesized that provision of the yDHODH gene in the knockdown parasites might overcome mtETC defects caused by PfmRPL13 disruption. We then performed a second transfection of the knockdown parasites with an episomal plasmid carrying yDHODH-GFP, resulting in the double transgenic line D10-PfmRPL13-KD-yDHODH-GFP (“Materials and methods”). Expression of the fused yDHODH-GFP protein was confirmed with a fluorescence microscope and Western blotting (data not shown). As shown in Fig. 5B, in the absence of aTc, the double transgenic line displayed moderate growth in the first four cycles (short term) but still succumbed to death significantly after five cycles or longer without aTc (long term). These data suggested that, in the long term, yDHODH expression still failed to rescue mtETC defects resulting from PfmRPL13 ablation. We then complemented D10-PfmRPL13-KD with an episomal PfmRPL13 tagged with 3Myc, resulting in another double transgenic line, D10-PfmRPL13-KD-RL13Myc (“Materials and methods”). Expression of PfmRPL13Myc was confirmed by Western blotting in the double transgenic line maintained with or without aTc for four cycles (data not shown). As shown in Fig. 5C, D10-PfmRPL13-KD-RL13Myc grew equally well in the presence or absence of aTc for a long term. To verify aptamer copy number in the endogenous PfmRPL13 locus of the double transgenic line, D10-PfmRPL13-KD-RL13Myc, we checked its 5′ integration site by PCR and sequencing and eight aptamer copies were still intact (data not shown). Therefore, in the absence of aTc, the endogenous PfmRPL13 was not translated but the double transgenic line was sustained by the episomal expression of PfmRPL13 to maintain growth. Clearly, genetic ablation of PfmRPL13 in the parasite was only rescued with provision of complementing episomal PfmRPL13.

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Figure 4. Morphological changes of mitochondria in the PfmRPL13 knockdown parasites maintained with or without aTc under transmission EM. A–E, TEM images of PfmRPL13 knockdown parasites maintained constantly under aTc. DV, digestive vacuole. AP, apicoplast. N, nucleus. F–J, TEM images of PfmRPL13 knockdown parasites grown without aTc for three cycles. DV, digestive vacuole. Cyt, cytosome. Black arrows indicate mitochondrial sections. Red arrows indicate internal membranes within the mitochondrial matrix. Each panel shows either an entire (A, B, F, G) or a partial (C–E, H–J) image of TEM.
mammalian mitochondria, for example, fatty acid catabolism, amino acid degradation, steroid biosynthesis, and initiation of apoptosis (35). Nevertheless, the mitochondrion is still absolutely essential in every stage of the parasite’s life cycle (39). Even Cryptosporidium, which has lost the mitochondrial genome, has a mitosome (mitochondrion derivative organelle) that harbors at least one essential biochemical pathway—iron-sulfur cluster biogenesis (8, 42). To maintain protein import and critical biochemical pathways, mitochondria and mitochondrial-related organelles require an electrochemical proton gradient (composed of $\text{pH}_{\text{m}}$) across the mitochondrial inner membrane. In malaria parasites, the primary mechanism to sustain $\Delta\psi_{\text{m}}$, the principle component of the proton gradient, is fulfilled by the mtETC (24, 38). Whereas most mtETC proteins are nuclear-encoded and imported into the mitochondrion, three core components are encoded on the 6 kb mtDNA: cyt $b$, COXI, and COXIII. The mitochondrial ribosomes and protein translation system are therefore critical to produce these three mtETC components. In this study, we have shown that PfmrRPL13 ablation in P. falciparum results in mtETC deficiency (Fig. 3B), which leads to the loss of $\Delta\psi_{\text{m}}$ (Fig. 2F) and parasite death (Fig. 2C). For the first time, we have provided direct evidence that mitochondrial ribosomes are essential in malaria parasites. It is likely that a domino effect follows PfmrRPL13 ablation: 1) reduced translation efficiency in mitochondrial ribosomes; 2) shortage of cyt $b$, COXI, and COXIII proteins; 3) failure to assemble functional mtETC complexes; 4) significant reduction of $\Delta\psi_{\text{m}}$; 5) inability to complete pyrimidine biosynthesis, and 6) ultimately parasite death. In the future, we plan to further characterize PfmrRPL13 knockdown parasites to better understand the detailed mechanisms leading to parasite collapse.

One proposed alternative pathway to maintain $\Delta\psi_{\text{m}}$ in malaria parasites is rotation of the $\text{F}_0\text{F}_1$ ATP synthase complex (Complex V) in the reverse direction, hydrolyzing ATP (−4 charge) to ADP (−3 charge) (23). One ADP (in the matrix) is then exchanged with one cytosolic ATP, moving one net negative charge across the mitochondrial inner membrane. In addition, ATP hydrolysis by intact coupled Complex V results in the extrusion of protons across the membrane in the opposite direction, providing a further contribution to $\Delta\psi_{\text{m}}$. In the yDHODH transgenic parasites, $\Delta\psi_{\text{m}}$ remained substantially intact even after a complete inhibition of the mtETC by atovaquone, but was rapidly disrupted when proguanil was added subsequently (23). The synergistic effect of atovaquone and proguanil in collapsing $\Delta\psi_{\text{m}}$ suggests that proguanil disrupts the alternative $\Delta\psi_{\text{m}}$ pathway that is likely maintained by Complex V. Remarkably, PfmrRPL13 knockdown parasites exhibited proguanil hypersensitivity to a similar degree as yDHODH transgenic parasites under atovaquone inhibition (Fig. 3C). These results suggest that PfmrRPL13 knockdown caused a severe defect in the mtETC, which rendered parasites reliant on the alternative pathway (proguanil-sensitive) to maintain $\Delta\psi_{\text{m}}$.

Ablation of PfmrRPL13 resulted in a significant morphological change in the parasite mitochondria (Fig. 4). It has been well known that the mitochondrion of P. falciparum in the asexual blood stages is a tubular structure with very few or no cristae on the inner membrane (35). However, in our PfmrRPL13 knockdown parasites, many long tubular and even branched mitochondria were observed in the TEM 2D sections, although they
still remained cristae-deficient (Fig. 4). The mechanisms that led to these unusual morphological changes in the mitochondria of knockdown parasites remain unknown at present. Because there is only one mitochondrion per parasite, it appears that the mitochondrion of the knockdown parasite has undergone a tremendous "growth" in mass and volume. We speculate that mtETC deficiency and/or other mitochondrial defects that result from PfmRPL13 ablation may have triggered a mitochondrial biogenesis pathway(s) to compensate for the loss of mitochondrial components or functions. On the other hand, because these are TEM 2D images, the likelihood that the unusual structures were caused by mitochondrial fragmentation cannot be ruled out. Clearly, further investigation is needed to address these hypotheses.

The results of rescue experiments indicate that the mtETC defects caused by PfmRPL13 knockdown cannot be fully released by addition of the yeast DHODH gene (Fig. 5B), even though yeast DHODH completely restored parasite growth under atovaquone inhibition (23). It has been shown that the bc1 complex of the yDHODH transgenic line remained fully functional (23). It is likely that atovaquone inhibits the electron transfer activity of the bc1 complex by binding to the Qo site of cyt b, but leaving the complex physically intact. In our PfmRPL13 knockdown parasites, however, the mtETC defects seem to be more severe. Mitoribosome deficiency caused by PfmRPL13 knockdown would be expected to compromise both the electron transfer activity and the physical integrity of the mtETC. Although the mtETC can be metabolically bypassed by expression of the yDHODH gene (23), the physical components of the mtETC cannot be supplemented by this exogenous gene. In the PfmRPL13 knockdown parasites, it is likely that loss of integrity of mtETC and \( \Delta \psi_m \) causes a wide range of detrimental defects in the mitochondrion. For example, the mitochondrial iron–sulfur cluster biogenesis pathway, dependent on a functional \( \Delta \psi_m \), will be deprived, which in turn is detrimental to the mitochondrion and the rest of the parasite. Indeed, our knockdown line was only able to be rescued by complementation with episomal expression of PfmRPL13 (Fig. 5C). These data further support that in addition to its enzymatic activities, physical integrity of the mtETC is essential for mitochondrial physiology.

In summary, we provide direct evidence that the mitochondrial ribosomes in the human malaria parasites are essential. Disruption of mitoribosomes by ablation of a critical protein subunit causes a chain of adverse events and eventual parasite demise. Our data highlight the significance of the mitochondrial protein translation machinery, which is evidently critical to sustain mitochondrial function, structure, and physiology.

Experimental procedures

**Plasmid construction**

For localization studies, the coding region of *P. falciparum* mitochondrial ribosomal subunit L13 (PfmRPL13, PF3D7_0214200) was cloned into the pLNmRL2 vectors, bearing either a 3HA or GFP tag (43). For CRISPR/Cas9-mediated knockout studies, two homologous regions and one gRNA of PfmRPL13 were cloned into the pL6 vector, which was kindly provided by Dr. Lopez-Rubio (44). The type II *Streptococcus pyogenes* Cas9 was encoded in the pUFl-Cas9 vector (44). For CRISPR/Cas9-mediated knockdown studies, the TetR-DOZI-aptamer system was utilized, which was kindly provided by Dr. Jacquin Niles (30). The original pMG75-ATP4-aptamer vector (30) was reconstructed to remove the attP sites and to replace the single HA tag with a triple HA tag. The pAll-In-One (pAIO) pre-gRNA vector was kindly provided by Dr. Spillman and Dr. Beck (31), which contained Cas9 fused with yDHODH (23) and elements for expressing a gRNA. For complementation studies, a second transfection was performed in the PfmRPL13 knockdown parasites, introducing episomal plasmids of yDHODH-GFP (23) or PfmRPL13–3Myc encoded on a vector that had a human dihydrofolate reductase selectable marker. Details for other cloning procedures are available in the supporting information.

**Parasite lines, parasite culture, and transfections**

See supporting information for details.

**Immunofluorescence assay**

Parasite samples (50 μl) were prelabeled with 60 nM MitoTracker Red CMXRos (M7512, Life Technologies by Thermo Fisher Scientific), fixed with 4% formaldehyde/0.0075% glutaraldehyde 1 h at 37 °C, permeabilized with 0.1% Triton X-100/ PBS for 10 min, treated with 0.1 mg/ml NaBH4 for 5 min and blocked with 5% BSA/PBS before primary and secondary antibody incubations. The HA probe (sc-7392, Santa Cruz Biotechnology) and an Alexa Fluor 488 conjugated antimouse secondary antibody (A-21141, Life Technologies by Thermo Fisher Scientific) were used at 1:350 dilution for overnight at 4 °C. All other steps followed the published protocol (43). The parasites were visualized with an Olympus epifluorescence microscope and images were processed using SlideBook software.

**Western blotting**

See supporting information for details.

**Parasite growth curves and knockdown experiment**

Parasite cultures were tightly synchronized with several rounds of alanine treatment (0.5 mM alanine/10 mM HEPE, pH 7.6) and trophozoite/schizont stages were isolated using a Percoll gradient (89428–524, GE Healthcare Life Sciences). The highly enriched parasites were thoroughly washed with medium and inoculated into new cultures with each 10 μl parasitized pellet diluted in 10 ml medium containing 1 ml blood. Cultures were maintained in the presence and absence of 250 nM anhydrotriacycline (Millipore Sigma) and were split 1:5 every 2 days. At each split, samples were collected for thin blood smears and protein analysis by Western blotting. Parasitemia was determined by counting at least 1000 RBCs under a Leica light microscope.

**MitoTracker staining and quantification**

The knockdown experiment was set up as described above. In every 2 days, aliquots of parasitized RBCs (~10 μl pellet)
Mitochondrial ribosomal protein L13 in Plasmodium falciparum

were taken from cultures maintained with or without aTc, re-suspended in 200 μl medium containing 10 nM MitoTracker Red CMXRos and incubated for 30 min. The parasites were washed three times with 1 × PBS, re-suspended in a small volume of medium (~15 μl) and visualized under an Olympus epifluorescence microscope. For each condition, a total of 200 or more parasites were examined within 15 min.

Mitochondrion preparation (Mito Prep)

Mitochondria of P. falciparum cultures were isolated according to a published protocol (45). Briefly, parasites were tightly synchronized and expanded to a large volume (~2 liters). Cultures were harvested at late trophozoite stage and lysed with 0.05% saponin. The pellet was washed, re-suspended in a MESH buffer and disrupted using a N2 cavitation bomb (45). The cell debris was removed by a low speed centrifugation and the supernatant was passed through a MACS Cell Separation Column (Miltenyi Biotec) at 4 °C. The eluted material was pelleted by centrifugation, aliquoted, and stored at −80 °C. Details are listed in the supporting information.

bc, complex enzymatic activity measurement

Cytochrome c reductase activity was assayed by a minor modification of published methods (45, 46). The assay was performed at 35 °C in a stirred cuvette with a total volume of 300 μl, which contained 5 μl Mito Prep sample, 100 μM decyubiquinol, 100 μM horse heart cytochrome c (Millipore Sigma), 0.1 mg/ml m-Docetyl β−Maltoside, 60 mM HEPES, pH 7.4, 10 mM sodium malonate, 1 mM EDTA, and 2 mM KCN. The reduction of cyt c was recorded at 550 nm with a CLARITY VF integrating spectrophotometer (OLIS, Bogart, GA). The short chain ubiquinol analog decyubiquinol was prepared by reducing decyubiquinone in DMSO with sodium borohydride and acidifying the mixture with concentrated HCl. Aliquots were stored under argon at −80 °C. Protein concentration of Mito Prep samples was determined by a Bio-Rad colorimetric assay using a spectrometer (Spectronic Genesys S) at the wavelength of 595 nm.

[3H]Hypoxanthine growth inhibition assay

Growth inhibition assays using [3H]hypoxanthine incorporation were performed in 96-well plates as described previously (36), and data were analyzed by GraphPad Prism 6. See the supporting information for details.

Magnetic enrichment of parasites and transmission EM

aTc was removed from PfmrRPL13 knockdown parasites for three intraerythrocytic cycles. Parasites were enriched by a MACS Cell Separation Column in a magnetic apparatus. See the supporting information for other details.

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