A first-in-class inhibitor of parasite FtsH disrupts plastid biogenesis in human pathogens

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There is an urgent need for antimalarials with distinct mechanisms-of-action to combat resistance to frontline drugs. The malaria parasite \textit{Plasmodium falciparum} and related apicomplexan pathogens contain an essential, non-photosynthetic plastid organelle, the apicoplast\(^1,2\), which is a key antiparasitic target. Despite its biomedical potential, broadly effective antimalarials targeting the apicoplast have been elusive due to the slow onset-of-action of drugs that inhibit apicoplast translation\(^3,4\) and the apicoplast’s limited metabolic function in the symptomatic stage of \textit{Plasmodium}\(^5\). Apicoplast biogenesis depends on novel, but largely cryptic, mechanisms for protein/lipid import and organelle inheritance during parasite replication\(^6,7\). These critical pathways present untapped opportunities to discover new parasite-specific drug targets. We used an innovative chemical rescue screen\(^5\) to identify the natural product antibiotic, actinonin\(^8\), as a first-in-class antimalarial compound inhibiting apicoplast biogenesis. Both chemical-genetic interaction and resistant mutation indicated that the unexpected target of actinonin in \textit{P. falciparum} and \textit{Toxoplasma gondii} is FtsH1, a homolog of a bacterial membrane AAA metalloprotease. We show that \textit{Pf}FtsH1 is essential for apicoplast biogenesis and parasite replication, making it the first apicomplexan-specific regulator of organelle biogenesis to be identified in a forward screen. Taken together, our findings demonstrate that FtsH1 is a novel and, importantly, druggable antimalarial target. Development of actinonin derivatives as FtsH1 inhibitors will have significant advantages over existing apicoplast-targeting compounds with improved drug kinetics, lower potential for clinical resistance, and multistage efficacy against multiple human parasites.

Because our molecular understanding of apicoplast biogenesis is limited, we performed a chemical screen to identify compounds that disrupt apicoplast biogenesis. Out of >400 antimalarial compounds tested (Extended Data Table 1)\(^9,10\), we identified a single natural product antibiotic, actinonin\(^8\), as a novel inhibitor of apicoplast biogenesis. Previously we demonstrated that biosynthesis of the isoprenoid precursor, isopentenyl pyrophosphate (IPP), is the sole essential function of the apicoplast in blood-stage \textit{P. falciparum}\(^3\). As such, compounds that target the apicoplast cause parasite growth inhibition that is fully rescued by the addition of IPP in the growth media. Actinonin caused \textit{P. falciparum} growth inhibition in a single replication cycle (EC\(_{50}\) = 3.2 \(\mu\)M; 95% CI 2.49-4.13) that was rescued by IPP (EC\(_{50}\) = 60.7 \(\mu\)M; 95% CI 49.6-74.5), demonstrating that it...
specifically inhibits the apicoplast with clinically-relevant kinetics (Fig. 1a; Extended Data Fig. 1a; Extended Data Table 1). In contrast, current drugs that inhibit translation in the apicoplast cause a characteristic “delayed death” after two replication cycles in vitro that limits their antimalarial efficacy and clinical use. When we assessed parasite growth inhibition and apicoplast defects caused by one of these known apicoplast drugs, chloramphenicol, (Extended Data Fig. 1b and 2a,c) actinonin clearly had more rapid antimalarial activity.

Furthermore, actinonin-treated P. falciparum, rescued for growth with IPP, no longer replicated their apicoplast and produced daughter parasites lacking apicoplasts, consistent with a defect in apicoplast biogenesis (Fig. 1b-c, Extended Data Fig. 2a,e). Again, this inhibition phenotype contrasts with that of inhibitors known to disrupt MEP isoprenoid precursor biosynthesis in the apicoplast. When we assessed the inhibition phenotype of fosmidomycin, a MEP inhibitor that is in clinical trials for antimalarial treatment, it also showed single-cycle antimalarial activity rescued by IPP (Extended Data Fig. 1b). Notably, though, fosmidomycin did not cause apicoplast loss (Extended Data Fig 2a,d).

Taken together, actinonin’s inhibition phenotype distinguishes it from known inhibitors that disrupt apicoplast translation and metabolism (Extended Data Fig 1b and 2) and indicates that it has a novel mechanism-of-action.

To further elucidate its mechanism-of-action, we first took a candidate-based approach to identify its target. Actinonin potently inhibits the bacterial and mitochondrial peptide deformylase (PDF), an enzyme that co-translationally removes the formyl group from the initiator methionine. Because the apicoplast translation machinery is prokaryotic in origin and contains a PDF, we tested whether the apicoplast PDF is the target of actinonin. In mitochondria, translation inhibitors suppress the effects of actinonin since translation is upstream of PDF activity. Surprisingly, apicoplast translation inhibition did not suppress the effects of actinonin (Extended Data Fig. 3), suggesting apicoplast PDF is not the target of actinonin.

We next took an unbiased approach to identify the target of actinonin by attempting to isolate actinonin-resistant P. falciparum but were unsuccessful using multiple selection methods, including chemical mutagenesis of the starting population. Instead we turned to Toxoplasma gondii, a related apicomplexan parasite, because it is easier to grow to large numbers and to genetically modify. As in P. falciparum, actinonin treatment caused growth inhibition (EC₅₀ = 13.7 μM; 95% CI 13.2-14.2; Extended Data Fig 4) and apicoplast loss (Fig. 2a) in T. gondii, suggesting a similar mechanism-of-action. We selected for actinonin-resistant T. gondii and determined the whole-genome sequences for eight independently selected clones (Fig. 2b). Five of these clones harbored a N805S mutation in the metalloprotease domain of the membrane AAA protease TgFtsH1 (TGGT1_259260) (Fig. 2d; Extended Data Table 2). TgFtsH1 was compelling for two reasons. First, TgFtsH1 localizes to the apicoplast. Second, actinonin is a peptide mimic containing a metal-binding hydroxamic acid, a class of molecules that typically binds metalloproteases. Confirming this resistance mutation, replacement of the endogenous TgFtsH1(WT) locus with the allele encoding TgFtsH1(N805S) was sufficient to confer actinonin resistance in T. gondii (Fig. 2c). The increased actinonin EC₅₀ in this
The mutant was similar to that measured for the actinonin-resistant clones that arose following drug selection. Taken together, the known metalloprotease binding of actinonin, the predicted metalloprotease activity of TgFtsH1, and the validated actinonin-resistant mutation in TgFtsH1 support FtsH1 as the target of actinonin in T. gondii, providing a strong candidate for validation in P. falciparum.

The P. falciparum genome contains three FtsH homologs (Pf3D7_1119600, Pf3D7_1239700, and Pf3D7_1464900). One of these, PfFtsH1 (Pf3D7_1239700) is most closely related to TgFtsH1 by phylogenetic analysis. Unlike TgFtsH1, PfFtsH1 was previously reported to localize to the mitochondria, not the apicoplast. However the same study reported that PfFtsH1 undergoes internal cleavage, such that the localization of the N-terminal fragment containing the ATPase and protease domains was unclear.

Setting aside its ambiguous localization, we sought to determine whether PfFtsH1 is required for apicoplast biogenesis and the relevant target of actinonin. We constructed a P. falciparum strain in which the endogenous PfFtsH1 locus was modified with a C-terminal FLAG epitope and 3' UTR tetR-DOZI-binding aptamer sequences to regulate its expression. As expected, PfFtsH1 expression was downregulated when the tetR-DOZI repressor bound the aptamer sequences, and restored when anhydrotetracycline, which disrupts this interaction, was added (Fig. 3a, Extended Data 5).

To test whether PfFtsH1 is essential for apicoplast biogenesis, we downregulated PfFtsH1 and observed a nearly 4-fold decrease in parasitemia after 3 replication cycles, compared to parasites expressing normal levels of PfFtsH1 (Fig 3b, Extended Data Fig. 6a). This growth defect observed after multiple replication cycles is weaker than the single-cycle growth inhibition observed upon actinonin treatment and likely reflects partial knockdown, which has been reported using this tet-regulation system (personal communication, Dan Goldberg). Significantly, growth of PfFtsH1 knockdown parasites was restored by addition of IPP, indicating PfFtsH1 is essential specifically for an apicoplast function (Fig 3b, Extended Data 6a). Finally, we confirmed that loss of PfFtsH1 led to apicoplast loss. Using qPCR for genes in the apicoplast and nuclear genome, we observed a steady decrease in the apicoplast:nuclear genome ratio upon knockdown of PfFtsH1 and growth rescue with IPP compared to control parasites, consistent with loss of the apicoplast (Fig. 3c). These results demonstrate that PfFtsH1 is essential and required for apicoplast biogenesis, making it the first protein with a novel function in apicoplast biogenesis identified in an unbiased screen.

Finally, because knockdown of PfFtsH1 phenocopies the apicoplast biogenesis defect of actinonin, we determined whether knockdown of PfFtsH1 sensitized parasites to actinonin. Indeed, when PfFtsH1 expression was downregulated, the actinonin EC50 decreased >50-fold compared to control parasites (Fig. 3d). Since these growth inhibition assays were performed over a single replication cycle, the downregulation of PfFtsH1 did not significantly affect parasite growth, and the actinonin EC50 could be measured without confounding growth inhibition caused by the decrease in PfFtsH1 levels (Fig. 3b, Extended Data Fig. 6b). Importantly, PfFtsH1 downregulation did not change the EC50 of fosmidomycin, indicating that the hypersensitivity to actinonin after PfFtsH1 downregulation is not due to general apicoplast disruption or parasite death (Extended...
Data Fig. 6c). Combined with the target identification in *T. gondii*, this chemical-genetic interaction strongly implicates FtsH1 as the target of actinonin in apicomplexan parasites.

Our study represents a rare example of a forward screen to uncover cryptic cellular pathways in *Plasmodium*.[20,21]. Acquired by secondary endosymbiosis of an alga, the apicoplast is evolutionarily distinct. Apicomplexan FtsH1’s role in organelle biogenesis is not conserved in homologs found in mitochondria or primary chloroplasts[22] and likely represents a novel pathway unique to secondary endosymbionts in this parasite lineage[7,15,23,24]. Based on the critical function of FtsH homologs in membrane protein quality control and complex assembly[22], we propose that FtsH1 regulates the proteolysis of key apicoplast membrane protein(s) during parasite replication. FtsH1 offers a rare foothold into a novel apicoplast biogenesis pathway that will yield deeper insight into the molecular mechanisms of eukaryogenesis and uncover additional antiparasitic targets.

Most importantly, our findings present an exciting opportunity for antimalarial drug discovery. FtsH1 inhibitors will have significant advantages over existing antimalarials that target apicoplast metabolism or translation. While metabolic needs vary throughout the parasite lifecycle and even between the same stage of different *Plasmodium* species,[25,26], apicoplast biogenesis is required at every proliferative stage of the parasite lifecycle and highly conserved among apicomplexan parasites. For example, apicoplast translation inhibitors which cause delayed biogenesis defects have broad clinical application as malaria prophylaxis targeting liver-stage *Plasmodium* spp and as a partner drug, in combination with faster-acting compounds, for acute parasitic diseases targeting blood-stage *Plasmodium* spp, *Toxoplasma gondii*, and *Babesia* spp. In fact, the utility of these antibiotics as antiparasitics would be greater if not for their slow activity. Inhibition of FtsH1 retains all the benefits of targeting apicoplast biogenesis with no delay in the onset-of-action. Moreover, our inability to select actinonin-resistant *Plasmodium* contrasts with the ready selection of *in vitro* resistance against antibiotics[27] and MEP inhibitors[9,28,29] and indicates a lower likelihood of clinical resistance to FtsH1 inhibitors.

Finally, several clinical lead candidates based on the actinonin scaffold have advanced into human clinical trials as bacterial PDF inhibitors[30]. We are currently developing an FtsH1-based assay to facilitate high-throughput screening of actinonin derivatives with increased potency and improved drug properties as antimalarials. Overall, FtsH1 inhibitors have potential for rapid onset, minimal clinical resistance, and multi-stage efficacy against multiple parasitic infections, and routes to identifying a clinical lead candidate are readily accessible.

References


### Methods

#### Chemicals

Fosmidomycin was purchased from Santa Cruz Biotechnology and 10mM aliquots were prepared in water. Chloramphenicol was purchased from Sigma Aldrich and 50mM aliquots were prepared in 100% ethanol. Actinonin was purchased from Sigma Aldrich and 25mM aliquots were prepared in 100% ethanol. Anhydrotetracycline was purchased from Sigma and 2.5mM aliquots prepared in 100% ethanol and used at a final concentration of 0.5uM.

Enoxacin, ciprofloxacin, levofloxacain, norfloxacin, novobiocin, coumeramycin, mericitabine, 2’deoxy-2-F-ctydine, gemcitabine, ADEP1a, beta-lactone 4, beta-lactone 7, and rifampin were acquired and solubilized as noted in Supplementary Table 1.

Isopentenyl pyrophosphate (IPP) was purchased from Isoprenoids LC and stored at 2 mg/mL in 70% methanol, 30% 10mM ammonium hydroxide at -80°C. To prevent methanol toxicity, aliquots of IPP were dried in the speed vacuum centrifuge before adding to cultures. All drugs were stored at -20°C and resuspended just prior to use.

#### *Plasmodium falciparum* culture and transfections

*P. falciparum* D10 (MRA-201), and D10 ACP1-GFP (MRA-568) were obtained from MR4. *P. falciparum* NF54<sup>inh</sup> was a gift from David Fidock (Columbia University). NF54
antB strain constitutively expressing Cas9 and T7 Polymerase, generated previously\(^3\), was used in this study. Parasites were maintained in human erythrocytes (2% hematocrit) in RPMI 1640 media supplemented with 0.25% Albumax II (GIBCO Life Technologies), 2 g/L sodium bicarbonate, 0.1 mM hypoxanthine, 25 mM HEPES (pH 7.4), 50 μg/L gentamycin, and 0.4% glucose at 37°C, 5% O₂, and 5% CO₂.

Parasites were transfected using methods already published\(^2\). Briefly, we used 50 ug of plasmid per 200 μL packed red blood cells (RBCs) adjusted to 50% hematocrit. We used a Bio-Rad Gene Pulser II to pre-load uninfected RBCs using eight square-wave pulses of 365 V for 1 ms, separated by 100 ms. Preloaded RBCs were resealed for 1 hour at 37°C and washed twice in RPMI to remove lysed cells. Schizont stage parasites at 0.5% parasitemia were then allowed to invade half of the preloaded RBCs during two sequential reinvasions. Media was changed daily for the first 12 days and every other day thereafter. Parasites were split 1:1 into fresh blood every 4 days until parasites were visible by Giemsa smear. To select for integration of the pFtsH1 into \(P. falciparum\) NF54antB-pPfCRISPR parasites, transfected parasites were maintained in media containing 5 nM WR99210 and 0.5 μM anhydrotetracycline (Sigma) and then selected with 2.5 mg/l Blasticidin S (Sigma) beginning 4 days after transfection.

**Toxoplasma gondii culture and transfection**

\(T. gondii\) RH and \(T. gondii\) RH \(Δku80Δhxgprt\) strains were a gift from Matthew Bogyo (Stanford University) and maintained by passage through confluent monolayers of human foreskin fibroblasts (HFFs) host cells. HFFs were cultured in DMEM (Invitrogen) supplemented with 10% FBS (Fetal Plex Animal Serum from Gemini), 2mM L-glutamine (Gemini), and 100 ug penicillin and 100 ug streptomycin per mL (Gibco Life Technologies), maintained at 37°C and 5% CO₂. Parasites were harvested for assays by syringe lysis of infected HFF monolayers.

For transfection of \(T. gondii\) Δku80Δhxgprt, 15ug of the pTgCRISPR plasmid was combined with 3 ug of the pFtsH1N805S or pFtsH1WT that had been linearized by NotI digestion. Approximately 10^7 parasites were released from host cells using syringe lysis and washed into 400uL of cytomix containing both plasmids. Parasites were electroporated (BTX Electro Cell Manipulator 600) with 1.2-1.4 kV, 2.5kV/resistance, R2 (24 ohm) and then allowed to recover in the cuvette at room temperature for 10 mins before adding to host cells. After 24 hours, media containing 25ug/mL mycophenolic acid (Sigma) and 50 ug/mL xanthine (Sigma) was added to select for transfectants. After a week, plaques were observed and single clones were isolated using limiting dilution.

**Toxoplasma gondii genome sequencing and SNP identification**

Actinonin resistant and susceptible \(T. gondii\) were grown on 15 cm dishes containing confluent HFF monolayers until spontaneous lysis of the monolayer was achieved. Released parasites were collected and filtered through 5 micrometer syringe filters (Millipore) before isolating DNA (Qiagen DNAeasy Blood & Tissue).

\(T. gondii\) genomic DNA isolated from either the parental \(T. gondii\) RH strain (SRR3666219) or any of its derived mutants (SRR3666219, SRR3666222, SRR3666224,
SRR3666792, SRR3666794, SRR3666796, SRR3666798, SRR3666799, SRR3666801) was sequenced in an Illumina NextSeq apparatus using 2x150bp reads at an average sequencing depth of 35x. Sequencing reads were quality trimmed and remnants of sequencing adaptors removed with trimmomatic (PMID:24695404). Next, reads were mapped to the reference nuclear assembly of the T. gondii GT1 strain (ToxoDB v13.0) and the apicoplast genome assembly from the RH strain (NC_001799) with the program bowtie2 (PMID:25621011). Duplicated aligned reads were removed with picard tools (http://broadinstitute.github.io/picard) and reads spanning InDels were realigned with GATK (PMID:20644199). Afterwards, allelic variants were called with samtools mpileup followed by bcftools call with –p set to 0.05 (PMID:26826718). Finally, classification of mutations was performed with snpEff (PMID:22728672).

**Growth inhibition assays**

For *P. falciparum* EC50 calculations, growth assays were performed in 96 well plates containing serial dilution of drugs in triplicate. Media was supplemented with 200 uM IPP as indicated. Growth was initiated with ring-stage parasites (synchronized with 5% sorbitol treatment 48 hours prior) at 1% parasitemia and 1% hematocrit. To calculate growth, cultures were incubated for 72h and growth was then terminated by incubation with 1% formaldehyde (Electron Microscopy Services) for 30 minutes at room temperature. Parasitized cells were stained with 50 nM YOYO-1 (Invitrogen) overnight at room temperature and the parasitemia was determined by flow cytometry (BD Accuri C6 Sampler). Data were analyzed by BD Accuri C6 Sampler software.

For *T. gondii* EC50 calculations, plaque assays were performed in 24 well plates containing confluent HFF monolayers serial dilutions of drugs in duplicate. Approximately 50 parasites were counted using flow cytometry and added to each well. After incubating for 6 days, infected monolayers were washed, fixed with methanol for 10 minutes, stained with 2% crystal violet (Sigma) for 30 minutes, and then washed again. Plaques were visualized as non-stained areas. The area of each plaque in a given well was measured and summed using ImageJ as a proxy for growth and normalized to the vehicle only control.

For measuring the growth inhibition of *P. falciparum* during the time course, 10 uM actinonin, 10 uM fosmidomycin, 30 uM chloramphenicol, 200 uM IPP, and 0.5 uM of anhydrotetracycline was used as necessary. For comparison of growth between different treatment conditions, cultures were carried simultaneously and handled identically with respect to media changes and addition of blood cells. Daily samples were collected and fixed with 1% formaldehyde for 30 minutes at RT. At the end of the time course, all samples were stained with 50 nM YOYO-1 and parasitemia was measured using flow cytometry. All growth curves were plotted using GraphPad Prism.

For measuring the growth inhibition of *T. gondii* during the time course, 6-well plates were set up with no drug, 40 uM actinonin, 25 nM clindamycin, and 4 uM pyrimethamine. *T. gondii* was added at a MOI = 3. Every 12 hours, parasites were released from HFFs using syringe lysis and counted using flow cytometry (BD Accuri C6 Sampler). After 36 hours, spontaneous lysis of the monolayer was observed and parasites
were counted using flow cytometry and then added back to fresh monolayers at MOI = 3
in the absence of drug and parasites were counted every 12 hours as before.

For co-treatments of *P. falciparum* with actinonin and chloramphenicol, 96 well plates
containing ring stage parasites at 1% parasitemia and 1% hematocrit were treated with
serial dilutions of both actinonin and chloramphenicol alone and in combination. To
determine the effect on growth after one lytic cycle, parasites were fixed at 72h and
parasitemia was measured by flow cytometry as above. To determine the effect on
growth after two lytic cycles, 75% of the media was exchanged at 72 hours and plates
were incubated for an additional 48 hours following fixation and flow cytometry as
above. Media was supplemented with 200 uM IPP as a separate control to insure
specificity of the drug at the concentrations used.

**Quantitative Real-Time PCR**
Parasites from 1 mL of *P. falciparum* culture at ring stage were isolated by saponin lysis
followed by two washes with PBS. DNA was purified using DNAeasy Blood & Tissue
(Qiagen). Primers were designed to target genes found on the apicoplast or nuclear
genome: tufA (apicoplast) 5'-GATATTGATTCCAGCTCCAGAAAGA-3' and CHT1
(nuclear) 5'-TGTTTCCTTCAACCCCTTTT-3' /5'-TGTTTCCTTCAACCCCTTTT-3'.
Reactions contained template DNA, 0.15 uM of each primer, and 1x SYBR Green I
Master mix (Roche). qPCR reactions were performed at 56C primer annealing and 65C
template extension for 35 cycles on a Applied Biosystem 7900HT system. Relative
quantification of target genes was determined. For each time point, the
apicoplast:nuclear genome ratio was calculated relative to the appropriate control
collected at the same time.

**Fluorescence Microscopy**
*P. falciparum* D10 ACP(L)-GFP parasites diluted to 0.05% hematocrit were settled on a
Lab-Tek II Chambered Coverglass (Thermo Fisher) and incubated in 2 ug/mL Hoechst
33342 stain for 15 minutes at 37C. Widefield epifluorescence live cell images were
acquired with an Olympus IX70 microscope. The microscope was outfitted with a
Deltavision Core system (Applied Precision) using an Olympus x60 1.4NA Plan Apo
Lens, a Sedat Quad filter set (Semrock) and a CoolSnap HQ CCD Camera
(Photometrics). The microscope was controlled and images were deconvolved via
softWoRx 4.1.0 software. ImageJ software was used to analyze resulting images.

Live microscopy of *T. gondii* RH FNR-RFP parasites was performing using Lab-Tek II
Chambered Coverglasses containing confluent HFF monolayers. Parasites were added at
an MOI = 1. After 36 hours of incubation, parasites were incubated with 2 ug/mL of
Hoescht for 15 minutes. Widefield epifluorescence live cell images were acquired with a
Nikon Eclipse Ti inverted fluorescence microscope with a NA 1.40 oil-immersion
objective (Nikon Instruments) and controlled using MicroManager v1.4. An iXon3 888
EMCCD camera (Andor) was used for fluorescence imaging and an a Zyla 5.5 sCMOS
camera (Andor) was used for phase contrast imaging. ImageJ software was used to
analyze the resulting images.
Immunoblot
Parasites from 9 mL of *P. falciparum* culture were isolated by saponin lysis, washed with PBS and resuspended in 1 x NuPAGE LDS sample buffer (Invitrogen). Proteins were separated by electrophoresis on 4-12% Bis-Tris gel (Invitrogen) and transferred to a nitrocellulose membrane. After blocking, membranes were probed with 1:2000 monoclonal mouse anti-FLAG M2 (Sigma) and 1:10,000 IRDye 680RD goat anti-mouse IgG (LiCor Bioscience). Fluorescence antibody-bound proteins were detected with Odyssey Imager at 700 nm (LiCor Biosciences).

Toxoplasma gondii resistance selection
Approximately 2 x 10^6 *T. gondii* RH parasites were added to T25s containing a confluent HFF monolayer and allowed to grow for 24 hours. To mutagenize, between 500 uM – 2 mM N-ethyl N-nitrosourea (ENU) diluted in DMSO was added to flasks and incubated for 2 hours at 37 C. Cultures were then washed twice with 10mLs of cold PBS and then released from host cells using syringe lysis. A quarter of the resulting parasites were passaged to T25s containing a fresh monolayer of HFFs. After two passages, parasites were treated with 40uM actinonin. After one passage under actinonin selection, a severe bottleneck was observed. Plaques of resistant parasites could be observed after one week of constant actinonin pressure with periodic media changes. Finally, single clones were isolated using limiting dilution.

*Plasmodium falciparum* construct generation
The primers used for generating different fragments are listed in the Extended Data Table 3. A construct for regulating expression of FtsH1 (PF3D7_1239700) in *P. falciparum* (pFtsH1) was generated from the parental pSN054, a modified pJazz linear plasmid (Extended Data Table 3). The left homology region was amplified from parasite genomic DNA using primers SMG476 and SMG 477 and was cloned using FseI and AsisI restriction sites. FtsH1 protein coding nucleotides 2348- 2643 were recoded using gene block (IDT) to remove the PAM site. The right homology region was amplified from parasite genomic DNA using primers SMG501 and SMG502. These fragments were cloned using the I-SceI restriction site. Targeting guide RNA was generated by klenow reaction using primers SMG514 and SMG515 and was inserted using the AflII site. All sequences were ligated into the parent plasmid using Gibson assembly. Constructs for regulation FtsH3 (PF3D7_1464900) were generated similarly (SMG505 and SMG506 for the right homology region; SMG495 and SMG496 for the left homology region; SMG518 and SMG519 for the gRNA klenow reaction). While this construct could be generated, no parasites emerged after two transfections. Constructs for regulating FtsH2 (PF3D7_1119600) using primers SMG503 and SMG504 for the right homology region and SMG481 and SMG507 for the left homology region were unable to be generated because of unsuccessful PCR.

Toxoplasma gondii construct generation
A construct for knocking in the FtsH1N805S allele into the endogenous loci of FtsH1 (pFtsHN805S) was generated from the parental pTKO2 vector. Briefly, a ~800 bp sequence upstream of the TGGT1_259260 start codon was amplified as the left
homology region (using primers KAJ1 and KAJ2). The FtsH_{N805S} sequence was then amplified from actinonin resistant cDNA (using primers KAJ3 and KAJ4). The HXGPRT resistance cassette was amplified off of the pTKO2 cassette (using primers KAJ5 and KAJ6). A ~800 bp sequence downstream of the TGTT1_{259260} stop codon was amplified as the right homology region (using primers KAJ7 and KAJ8). To insert the right homology region, the pTKO2 plasmid was cut with HindIII and HpaI and the 800 bp sequence was inserted using infusion (Clontech). To insert the left homology region and the FtsH1_{N805S} allele, the pTKO2 plasmid containing the downstream homology region was cut with NotI and EcoRI and the two PCR products were inserted also using infusion (Clontech). The resulting colonies were tested using a diagnostic HpaI digest and correct clones were subjected to Sanger sequencing of the inserts. To revert the pFtsHN805S construct to pFtsHWT, we used Q5 mutagenesis (NEB) and primers KAJ9 and KAJ10.

To increase the transfection efficiency and specificity, CRISPR-Cas9 was used to insert a double stranded break at the site of insertion (the endogenous TGTT1_{259260} allele). Briefly, pSAG1::Cas9-U6::sgUPRT^{33} was modified to contain a guide sequence specific to a FtsH1 intron using the Q5 mutagenesis kit (NEB) and primers KAJ11 and KAJ12. Sanger sequencing of the guide was used to verify the resulting plasmid.


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K.A-J. and E.Y. conceived and designed experiments. K.A-J. performed the majority of the experiments. S.M.G. and J.C.N. designed and generated the FtsH1 knockdown construct. H.A.L. performed the whole-genome sequencing and variant analysis. K.A-J. and E.Y. analyzed the data and wrote the manuscript. All authors discussed and edited the manuscript.

Competing financial interests
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Fig 1: Actinonin inhibits apicoplast biogenesis in *P. falciparum*

(a) Time course of parasite growth during actinonin treatment with or without IPP, normalized to control cultures with or without IPP as appropriate. Error bars represent the SEM of two biological replicates.

(b) Time course of the apicoplast:nuclear genome ratio measured by quantitative PCR (qPCR) using primers for the apicoplast and nuclear genomes during treatment with actinonin and IPP. Genome ratios were normalized to control parasites grown with IPP only. Error bars as in a.

(c) Representative images of the apicoplast of IPP-rescued control and actinonin treated parasites 24 hours after treatment during the schizont stage. The apicoplast is visualized using the *P. falciparum* reporter strain D10 ACP-GFP in which GFP is targeted to the apicoplast and the nucleus is stained with Hoescht 33342. During *Plasmodium* replication, the apicoplast starts as a single small spherical organelle (ring stage) which branches and divides into multiple apicoplasts (schizont stage). A punctate apicoplast that does not branch indicates a defect in apicoplast biogenesis.
Fig 2: A mutation in the protease domain of *TgFtsH1* is sufficient to confer resistance to actinonin in *T. gondii*

(a) Representative images of the apicoplast of control and actinonin treated parasites 36 hours after infection. Representative parasites missing their apicoplast are denoted with white arrows. The apicoplast is visualized using the *T. gondii* reporter strain RH FNR-RFP in which RFP is targeted to the apicoplast and the nucleus is stained with Hoechst 33342. Each untreated parasite contains one apicoplast, except during cell division when there may be two.

(b) Dose-dependent parasite growth inhibition upon treatment with actinonin for the actinonin-sensitive parent strain (RH) compared with 3 independent clones following selection for actinonin resistance (resistant 1, resistant 2, resistant 3). These three resistant clones are representative of the eight clones submitted for whole genome sequencing. Growth was measured via summed areas of the plaques formed during plaque assays and normalized to untreated controls. Error bars represent the SEM of two biological replicates.

(c) Dose-dependent parasite growth inhibition upon treatment with actinonin for *TgFtsH1*(WT) compared with *TgFtsH1*(N805S) parasites in RH Δ*KU80* strain. Data was measured and analyzed as in 2b.

(d) Schematic of *TgFtsH1*. This protein contains a *N*-unique region containing a putative transmembrane domain, an AAA ATPase domain used for unfolding proteins, a peptidase domain with a zinc co-factor in the catalytic site, and a *C*-unique region. The resistance-conferring variant *TgFtsH*(N805S) is found in the peptidase domain near the catalytic site.
Fig 3: Knockdown of FtsH1 in *P. falciparum* leads to apicoplast loss and hypersensitivity to actinonin

(a) Schematic of the endogenous knockdown strategy. When anhydrotetracycline (aTC) is present in the media, the tet-repressor binds aTC and does not bind the 10x-aptamer sequence, which relieves translational repression, allowing FtsH1 to be expressed. When aTC is washed out of the media, the tet-repressor binds the 10x-aptamer and prevents expression of FtsH1.

(b) Time course of parasite growth without aTC and in the presence or absence of IPP in the media, normalized to the untreated or IPP-rescued parental strain as appropriate. Error bars represent the SEM of two biological replicates.

(c) Time course of the apicoplast:nuclear genome ratio measured by quantitative PCR (qPCR) using primers for the apicoplast and nuclear genomes during treatment with or without aTC. All samples contained IPP to rescue parasite growth. Genome ratios were normalized to respective parental cultures also grown with IPP. Error bars as in c.

(d) Dose-dependent parasite growth inhibition by actinonin in the absence or presence of aTC. Error bars as in c.
Extended Data 1: Actinonin specifically inhibits the apicoplast of *P. falciparum* leading to parasite death after a single replication cycle

(a) Dose-dependent parasite growth inhibition by actinonin in the absence or presence of IPP. Error bars represent the standard error of the mean (SEM) of two biological replicates.

(b) Time course of parasite growth during treatment with actinonin, chloramphenicol, or fosmidomycin in the absence or presence of IPP. Growth is normalized to untreated or IPP-rescued controls as appropriate. Error bars as in a. In contrast to other antimalarials, which cause growth inhibition in a single replication cycle, “delayed death” in *P. falciparum* is associated with inhibitors of apicoplast gene expression and has previously been described in detail. It is characterized by a normal apicoplast and cell division during drug treatment for one replication cycle, followed by halted apicoplast and cell division in daughter parasites of drug-treated parasites.
Extended Data Fig 2: Actinonin has a distinct inhibition phenotype compared to inhibitors of apicoplast metabolism and translation

(a) Time course of the apicoplast:nuclear genome ratio measured by quantitative PCR (qPCR) for targets in the apicoplast and nuclear genome during treatment with actinonin (black), chloramphenicol (blue), or fosmidomycin (green). While actinonin treatment blocks apicoplast genome replication in a single replication cycle, this effect is not observed until the second replication cycle of chloramphenicol treatment and not at all during fosmidomycin treatment. This is consistent with actinonin blocking apicoplast biogenesis, chloramphenicol blocking apicoplast translation leading to a delayed apicoplast biogenesis defect, and fosmidomycin blocking apicoplast metabolic function but not biogenesis. All samples were grown in IPP and genome ratios were normalized to the untreated control cultures also containing IPP. Error bars represent the SEM of at least 2 biological replicates.

(b-e) Representative images of the apicoplast during schizont stage of three successive replication cycles in untreated (b), chloramphenicol (c), fosmidomycin (d) and actinonin (e) treated cultures all grown with IPP. The apicoplast is visualized using the P. falciparum reporter strain D10 ACP-GFP in which GFP is targeted to the apicoplast and the nucleus is stained with Hoescht 33342. In this case, a branched apicoplast indicates successful apicoplast development while punctate apicoplasts (observed in replication cycle 1 for actinonin treatment (e) and replication cycle 2 for chloramphenicol treatment (c)) represents an apicoplast that has failed to develop. The apicoplast is no longer present after replication cycle 2 of actinonin treatment and upon replication cycle 3 of chloramphenicol treatment, which leads to complete mislocalization of the GFP.
Extended Data Fig 3: Growth inhibition in the first replication cycle by actinonin is not suppressed by apicoplast translation inhibition.

Parasite growth after one or two replication cycles after treatment with actinonin, chloramphenicol, or both actinonin and chloramphenicol normalized to growth of an untreated control. Treatment with actinonin alone inhibited growth after the first replication cycle, whereas treatment with chloramphenicol alone inhibited growth after the second replication cycle. Co-treatment with chloramphenicol, which targets apicoplast translation, did not suppress effects of actinonin treatment, which was inconsistent with actinonin targeting the peptide deformylase (PDF) of the apicoplast.
Extended Data Fig 4: Actinonin treatment causes “delayed death” in *T. gondii* associated with apicoplast loss

Time course of parasite growth of untreated (blue), clindamycin (green), pyrimethamine (black), or actinonin (red) treated parasites over the course of two lytic cycles. Unlike *P. falciparum*, *T. gondii* undergoes multiple replication cycles in the host cell before lysis, thus each *T. gondii* lytic cycle represents multiple parasite replication cycles. Pyrimethamine inhibits parasite dihydrofolate reductase and was used as a control for a non-apicoplast targeting drug that inhibits growth in a single lytic cycle. As previously reported, clindamycin, an apicoplast translation inhibitor, gave a “delayed death” phenotype in *T. gondii* characterized by completed cell divisions and host re-invasion during drug treatment in the first lytic cycle, followed by halted cell division in the second lytic cycle. Actinonin also led to growth inhibition in the second lytic cycle, suggesting that it also targets the apicoplast. Error bars represent the SEM of two biological replicates. It is important to distinguish between apicoplast-associated “delayed death” in *P. falciparum* and that in *T. gondii*. In *T. gondii*, apicoplast loss occurs in the first lytic cycle and is temporally separate from defects in parasite cell division and growth inhibition observed in the second lytic cycle; whereas in *P. falciparum* defects in apicoplast biogenesis, parasite cell division, and growth inhibition all occur in the second replication cycle. “Delayed death” in *T. gondii* therefore appears more broadly associated with disruption of apicoplast biogenesis, whereas “delayed death” in *P. falciparum* appears more specific to disruption of apicoplast gene expression that leads to delayed biogenesis defects.
Extended Data Fig 5: Removal of anhydrotetracycline (aTC) from the media results in efficient knockdown of PfFtsH1.

Consistent with the previous report of PfFtsH1 C-terminal processing, we were unable to detect full-length PfFtsH1-FLAG using anti-FLAG in parasites with intact apicoplasts. However, upon loss of the apicoplast, full-length PfFtsH1-FLAG was detectable, suggesting that PfFtsH1 processing occurs in the apicoplast. Therefore, to assess the knockdown efficiency of PfFtsH1, we used a western blot comparing PfFtsH1-FLAG levels in the presence (lanes 1-4) or absence of aTC (lanes 5-8) in IPP-rescued parasites missing their apicoplast. Each sample was taken at the trophozoite stage and cycle 0 indicates 24 hours after the removal of aTC. Lanes 9 and 10 are samples from the parental strain that do not contain the FLAG-tag or the aptamer sequence in the 3' UTR of PfFtsH1. In each case, Cas9-FLAG was used as a loading control. PfFtsH1-FLAG levels were reduced to undetectable levels at 24 hours after aTC removal, validating our knockdown strategy.
Extended Data Fig 6: Knockdown of PfFtsH1 specifically disrupts the apicoplast and leads to specific hypersensitivity to actinonin

(a) Time course of parasite growth with or without anhydrotetracycline (aTC) and with or without IPP in the media. IPP rescues the growth defect observed in upon PfFtsH1 downregulation, indicating that PfFtsH1 is essential for an apicoplast-specific function. Growth is shown normalized to the untreated or IPP-rescued parental strain as appropriate. Error bars represent the SEM of two biological replicates.

(b) Dose-dependent parasite growth inhibition by actinonin with or without aTC for the parental (red) and PfFtsH strain. The EC50 of the parental strain is unchanged by the removal of aTC. Error bars as in a.

(c) Dose-dependent parasite growth inhibition by fosmidomycin with or without aTC in and with or without IPP. The fosmidomycin EC50 is unchanged by regulating levels of PfFtsH1, indicating that the observed hypersensitivity to actinonin upon knockdown of PfFtsH1 is specific to actinonin and does not occur for all apicoplast drug. Error bars represent the SEM of three technical replicates.