Uncovering the essential genome of the human malaria parasite 
*Plasmodium falciparum* by saturation mutagenesis

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Abstract

Severe malaria is caused by the apicomplexan parasite *Plasmodium falciparum*. Despite decades of research the unique biology of these parasites has made it challenging to establish high throughput genetic approaches for identification of therapeutic targets. Using transposon mutagenesis of *P. falciparum* in an approach that exploited its AT-rich genome we generated >38,000 mutants, saturating the genome and defining fitness costs for 95% of genes. Of 5,399 genes we found ~3,000 genes are essential for optimal growth of asexual blood-stages in vitro. Our study defines ~1000 essential genes, including genes associated with drug resistance, vaccine candidates, and conserved proteins of unknown function. We validated this approach by testing proteasome pathways for individual mutants associated with artemisinin sensitivity.
Transposon mutagenesis of *Plasmodium falciparum* was used to generate >38,000 mutants, saturating the genome and defining fitness costs for 95% of genes. We functionally define the relative fitness cost of disruption for 5,399 genes, and find that ~3,000 genes, ~62% of the genome, are essential for optimal asexual blood-stage *in vitro* growth. Our study defines ~1000 essential genes, including genes associated with drug resistance, leading vaccine candidates, and hundreds of *Plasmodium*-conserved proteins of unknown function that are now potential therapeutic intervention targets. We experimentally validated the essentiality of proteasome pathways with drug studies of individual mutants associated with artemisinin sensitivity. This study defines high-priority targets and pathways and points the way for the future of *P. falciparum* high throughput genetics.

**One Sentence Summary:**
Saturation-scale mutagenesis of *Plasmodium falciparum* reveals a core set of genes essential for asexual blood-stage growth *in vitro.*

**Abstract**

**INTRODUCTION:** Malaria remains a devastating global parasitic disease, with the majority of malaria deaths caused by the highly virulent *Plasmodium falciparum*. The extreme AT-bias of the *P. falciparum* genome has hampered genetic studies through targeted approaches such as homologous recombination or CRISPR-Cas9, and only a few hundred *P. falciparum* mutants have been experimentally generated in the past decades. In this study, we have used high throughput *piggyBac* transposon insertional mutagenesis and Quantitative Insertion Site Sequencing (QIseq) to reach saturation-level mutagenesis of this parasite.

**RATIONALE:** Our study exploits the AT-richness of *P. falciparum* genome, which provides numerous *piggyBac* transposon insertion targets within both gene coding and non-coding flanking sequences, to generate over 38,000 *P. falciparum* mutants. At this level of mutagenesis, we could distinguish essential genes as non-mutable and dispensable genes as mutable. Subsequently, we identified 3,357 genes essential for *in vitro* asexual blood-stage growth.
RESULTS: We calculated Mutagenesis Index Scores (MIS) and Mutagenesis Fitness Scores (MFS) to functionally define the relative fitness cost of disruption for 5,399 genes. A competitive growth phenotype screen confirmed that MIS and MFS were predictive of the fitness cost for in vitro asexual growth. Genes predicted to be essential included genes implicated in drug resistance, such as the “K13” Kelch propeller, mdr and dhfr-ts, as well as targets considered to be high-value for drugs development such as pkg, and cdpk5. The screen revealed essential genes that are specific to human Plasmodium parasites but absent from rodent-infective species, such as lipid metabolic genes that may be crucial to transmission commitment in human infections. MIS and MFS profiling provides a clear ranking of the relative essentiality of gene ontology (GO) functions in P. falciparum. GO pathways associated with translation, RNA metabolism, and cell cycle control are more essential, whereas genes associated with protein phosphorylation, virulence factors, and transcription are more likely to be dispensable. Finally, we confirm that the proteasome-degradation pathway is a high-value druggable target based on its high ratio of essential:dispensable genes, and by functionally confirming its link to the mode of action of artemisinin, the current front-line antimalarial.

CONCLUSION: Saturation-scale mutagenesis allows prioritization of intervention targets in the genome of the most important cause of malaria. The identification of the essential genome, consisting of over 3000 genes, will be valuable for antimalarial therapeutic research.

Saturation-scale mutagenesis of Plasmodium falciparum reveals genes essential and dispensable for asexual blood-stage development.

(A) The Mutagenesis Index Score (MIS) rates the potential mutability of P. falciparum genes based on the number of recovered CDS insertions relative to the potential number that could be recovered by large-scale mutagenesis. Genes highlighted are those previously validated to be dispensable (var, eba175, orc-1) or essential (cdpk5, kelch K13, dhfr-ts, pkg). (B) A high-resolution map of a ~50 KB region of chromosome 13 depicts an essential gene cluster, including K13, that lack insertions in the CDS but is flanked by dispensable genes with multiple CDS-disrupting insertions. (C) Conserved essential genes of known (PF3D7_1032900, RNA polymerase II-associated protein 1, putative, PF3D7_1033100, S-adenosylmethionine decarboxylase/ornithine decarboxylase) and unknown function (PF3D7_1033000) were concentrated in syntenic blocks whereas dispensable genes were significantly more common in the syntenic breaks, revealing an evolutionarily-conserved genomic architecture.

Malaria caused by Plasmodium falciparum remains an insidious global health problem, with hundreds of thousands of deaths each year. Recently there have been significant reductions in disease intensity, in part through concerted recent use of artemisinin combination therapies (ACTs), but these gains are now threatened by emerging ACT treatment failures spreading across South East Asia (1, 2). If ACT resistance reaches Africa, a devastating rebound of disease is expected, as occurred with chloroquine resistance in the 1970s. The development of new antimalarial therapies, and identification and prioritization of new targets, is a priority. More than a decade after the completion of the P. falciparum genome, a significant fraction of its genome still lacks functional annotation (3). Although CRISPR-Cas9 and other targeted endonucleases will accelerate functional genomics studies (4–6),

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their usefulness for genome-scale applications is restricted by the absence of non-
homologous end joining in \textit{Plasmodium} parasites, and by the extreme AT-richness of the \textit{P. falciparum} genome which reduces gRNA target site abundance. Therefore, a critical need remains for large-scale genetic analysis to systematically identify essential genes and prioritize parasite metabolic pathways for drug discovery (7).

Large-scale genetic screening methods in model organisms rely on efficient scalable methods for genome engineering. Transposon mutagenesis using the \textit{piggyBac} transposon, which preferentially inserts at the tetranucleotide target sequence TTAA, has been used to carry out whole-genome loss-of-function screens in many organisms (8–11). The highly skewed nucleotide composition of the \textit{P. falciparum} genome, with >81% AT content, is an advantage for the application of \textit{piggyBac} mutagenesis. The skewed composition results in a high density of TTAA sites, averaging one site per 70 bp over both coding and non-coding regions, in theory allowing systematic and saturation-level mutagenesis of the whole genome. While \textit{piggyBac} mutagenesis has been developed and optimized for \textit{P. falciparum}, and previously used for small-scale phenotypic screens and functional characterization of loss-of-function mutants, it has not been used for large-scale screening (12–16). To scale up \textit{piggyBac} mutagenesis in \textit{P. falciparum}, we developed high-throughput transfection mutagenesis methods that mostly create a single insertion per genome, and combined them with an Illumina-based sequencing method for identifying transposon insertion sites (17, 18). This approach, known as Quantitative Insertion-site Sequencing (QIseq), thus allowed whole-genome experimental mutagenesis analysis of \textit{P. falciparum} (fig. S1, A and B).

\textbf{Achieving saturation-level mutagenesis}

In a preliminary study we carried out large-scale transfections followed by short-term \textit{in vitro} growth of mixed pools of drug-selected \textit{P. falciparum} parasites and identified insertion sites by QIseq. This pilot identified a total of 3,651 insertions across the \textit{P. falciparum} genome (table S1)(17, 19). Based on the density and distribution of those insertions modeled by a Negative Binomial Distribution, we predicted that recovery of ≥33,000 insertions would be sufficient to achieve saturation-level mutagenesis in the compact genome of the malaria parasite (fig. S1C-F); this number is sufficient for there to be a high probability that multiple transposon insertions would occur within the CDS of every protein-coding gene larger than 500 bp. We therefore scaled our transfection methods to achieve this number of insertions. Transfected parasite populations were drug-selected briefly to isolate only mutant parasites with integrated \textit{piggyBac} elements carrying the drug resistance gene \textit{hdhfr}, before insertion-site locations were identified by QIseq. Computational analyses were used to verify the sequence reads of the QIseq libraries and the consistency of the raw data (fig. S2A-F); nearly all QIseq-defined insertions occurred at the characteristic TTAA target sequence. All insertions sites that were not flanked by the consensus TTAA (2.49%) insertion site were removed from subsequent analysis (fig. S2A). Previously validated individual \textit{piggyBac} mutant clones were included in each QIseq run to act as a control for the accuracy and sensitivity of each insertion-site identification run (fig. S3A-C and table S2).

The saturation mutagenesis approach identified ~38,000 independent \textit{piggyBac} insertions at distinct TTAA target sites, covering 5,399 nuclear protein-coding genes across all 14
chromosomes of the *P. falciparum* genome (Fig. 1A and table S3). These randomly distributed insertions exceeded those predicted to be required to achieve saturation-level mutagenesis (Fig. 1A), but there were numerous genes and regions encompassing several genes that had significantly fewer insertions than would be expected based purely on the distribution of TTAA-sites across the genome (Fig. 1B, C and D). As well as discrepancies in non-random spatial distribution, coding regions overall lacked insertions compared with intergenic regions (*p* > 2.2e-16, Fisher test) (Fig. 1D). A more detailed analysis of piggyBac insertion density within transcriptional units revealed that insertions in CDS were threefold less common than in flanking intergenic regions, and even within intergenic regions, insertion-site density decreased with proximity to CDS (Fig. 1E). This significant bias towards recovering intergenic insertions in surviving blood-stage parasites, with many CDS having no insertions, are indications that genes lacking insertions are lethal when disrupted by piggyBac insertion.

**Defining gene dispensability using mutagenesis index scores**

While there was a bias against insertions in CDSs, 2,042 genes were disrupted by at least one piggyBac insertion (~38% of genes in the *P. falciparum* genome) (Fig. 1F). The remainder, 3,357 genes (~62% of the genome) had no insertions in their CDSs, and some of these were also completely devoid of insertions in the surrounding intergenic regions (2.9% of genes) (Fig. 1C, fig. S4A and table S4). In the 2,042 mutable genes, insertion sites were distributed uniformly along the gene body CDSs, indicating all disruptions have equivalent consequences for the disrupted gene (Fig. 1E). The uniform distribution of transposon insertions throughout the genome is independent from chromatin structure and gene locations (fig. S4B). We therefore reasoned that the recovery of one or more insertions within the CDS of disrupted genes was an indicator of gene dispensability for *in vitro* asexual blood-stage growth. Therefore, mutable genes are subsequently referred to as dispensable, while the absence of any insertions in the CDS could be considered an indicator that disruptions are lethal, and subsequently these genes are referred to as essential. To quantify the evidence for dispensability or essentiality of each gene we developed a mutagenesis index score (MIS) based on the number of identified piggyBac insertions relative to the number of available TTAA target sites within that gene (Fig. 2A and table S5). Genes with higher MIS (on a scale of 0 to 1) were considered to have a higher possibility of dispensability, while those with lower MIS were considered to have a higher possibility of essentiality. The 2000 genes with highest and lowest MIS represent mutability characterization with the strongest confidence.

**Biological processes identified as dispensable and essential**

Mutants in the lowest MIS quartile were enriched in core metabolic processes shared across eukaryotes, which would be predicted to be essential. Mutants in the highest MIS quartile were enriched in parasite-specific multi-gene families that interact directly with the host, and which are known to be partially functionally overlapping and contain many redundant genes (Fig. 2B). It is important to note that this screen was carried out using *in vitro* cultured parasites, and many of these *in vitro* dispensable genes, such as those for antigenic variation and cytoadherence, have greater importance *in vivo* in human infections (fig. S5). The dispensability of parasite processes involved in host-parasite interactions contrasted with the
essentiality of internal parasite processes, such as those associated with RNA metabolism (Fig. 2C-D, fig. S6 and table S6). RNase II, a gene implicated in a post-transcriptional regulatory mechanism relevant to severe malaria, is an example with a low MIS score, consistent with previous reports of the locus being refractory to disruption (20). Other genes involved in general RNA metabolism that have low MIS include both widely-conserved RNA-binding proteins (such as PABP) and apicomplexan-specific RNA-binding proteins likely to have more specific, mostly unidentified, regulatory targets. The P. falciparum genome has an abundance of RNA-binding proteins that are largely functionally uncharacterized and our analysis suggests that many of these genes are likely to be essential.

MIS therefore correlates with what is known broadly about the importance or redundancy of metabolic pathways. To further validate that MIS is a good predictor of essentiality, we analyzed genes that have been the focus of drug or vaccine development. Genes strongly predicted to be essential based on MIS included the “K13” Kelch propeller implicated in artemisinin resistance, as well as other genes implicated in drug resistance, such as DHFR-TS, MDR and AAC2, involved in pyrimethamine, mefloquine and atovaquone resistance, respectively. Other genes classified as essential based on MIS included high priority blood-stage drug targets such as PKG, and CDPKS. By contrast, most blood-stage vaccine candidates were dispensable, with the notable exception of RH5. Not unexpectedly, sporozoite vaccine candidates CSP and TRAP, which are essential for sporozoite development in mosquitoes but are not required in blood-stage development (21), had high MIS. In contrast, the gene for pore-forming protein Cell Traversal of Ookinotes and Sporozoites (CelTOS) had low MIS. CelTOS is important for these parasite migratory phases and is emerging as a pre-erythrocytic-transmission blocking vaccine candidate; its low MIS indicates it may also have an essential function in blood-stage infections, making it a potential multi-stage vaccine target. MIS analysis also revealed likely essential genes that are specific to human Plasmodium parasites but absent from rodent-infective species, such as the lipid metabolic genes (PCD, PMT) that are crucial in the development of mosquito-transmissible sexual-stages in P. falciparum (22, 23).

**Mutant growth fitness as a measure of gene dispensability**

We previously developed a phenotyping method in which a pool of piggyBac mutants were grown as mixed-mutant pools over multiple generations, and the number of reads for each piggyBac insertion site quantified using next-gen sequencing to measure parasite growth rates (17). We adapted this method to generate a second quantitative measure of gene dispensability independent from MIS, by comparing the normalized number of reads from each insertion site to the total pool of reads across all P. falciparum mutants in the saturation mutagenesis screen (Fig. 2E and table S5). This mutagenesis fitness score (MFS) serves as a proxy for mutant growth fitness. MFS scores strongly correlated with MIS scores, despite the fact that they are independent of each other (Fig. 2F). The MFS of mutable genes was high, in keeping with the MIS prediction of dispensability, while predicted essential non-mutable genes had a low MFS (Fig. 2G). Intermediate MIS and MFS scores were weakly correlated, in keeping with the lower confidence that we can place on the essentiality or dispensability of genes with these intermediate scores.
A competitive in vitro growth screen (17) was used to phenotype four separate pools of mixed-mutant populations to provide additional validation of the correlation between MIS and a gene’s importance for growth fitness. Mutable genes with high MIS had low or minimal fitness cost were competitive growth ‘winners’, whereas growth fitness ‘losers’ had relatively low MIS (Fig. 3A and table S7). Relatively few mutations had little or no fitness cost, as measured by MFS, resulting in a disproportionate number of mutant ‘losers’ (Fig. 3B). Overall both the MIS and MFS were predictive of the fitness cost for in vitro asexual growth (Fig. 3C), and between these metrics we were able to predict separate 5,399 genes in the P. falciparum genome into non-mutable and mutable categories (Fig. 3D and E).

**Association of essentiality with genome structure and transcription patterns**

Dispensable and essential genes rarely occurred as single isolated genes, but rather occurred as multi-gene clusters reminiscent of conserved syntenic blocks (Fig. 1A, C and fig. S4A) (24). We therefore assessed the relationship between evolutionary conservation of genome structure and essentiality/dispensability by using previously defined syntenic relationships between Plasmodium spp. (25). Syntenic genes indeed had lower MIS, suggesting conserved essential functions across Plasmodium spp.; conversely, non-syntenic genes were more likely to be dispensable (Fig. 4A). Mapping chromosomal regions with synteny breaks, which typically harbor gene duplications and paralogs, showed patterns of clustering of essential and dispensable genes in syntenic and non-syntenic chromosomal regions, as seen in examples for chromosomes 13 and 10 (Fig. 4B-E).

Transcription metrics for each gene based on maximum FPKM values (26) were examined for correlations with essentiality and dispensability (Fig. 5A). This analysis showed that essential genes are expressed at significantly higher levels throughout both the asexual and sexual life cycle phases, indicating they may have critical functional roles throughout both the mosquito and human stages and therefore represent potential multi-stage drug targets (Figs. 5B and C). Within the intraerythrocytic cycle, genes with peak expression during the trophozoite stage were more likely to be essential than those expressed during other stages (Fig. 5C). During the trophozoite stage, P. falciparum must acquire nutrients from the host and remodel the infected erythrocyte to avoid host defenses, such as clearance by the spleen. In contrast, dispensable processes were enriched at either end of intraerythrocytic development, reflecting parasite stages with a greater need for functional redundancy, such as those involved in host interaction (e.g., erythrocyte invasion, antigenic variation), and are hence more likely to undergo duplication/paralog evolution. Not unexpectedly, about 41% of the dispensable genes are predominantly expressed in sexual stages (Fig. 5C). These genes may be essential for transmission to or from the mosquito, but do not appear to have an essential function for asexual blood-stage development. The high coverage of sexual-stage genes in this screen emphasizes the potential for piggyBac-based screens for identifying sexual phenotypes.

**Evolutionary conservation of apicomplexan gene essentiality**

We compared the data from our genome-wide saturation screen with data from large-scale genome sequencing studies of P. falciparum, as well as data from recent large-scale mutagenesis studies of apicomplexan genomes (6, 18, 27–29) to evaluate the conservation of
gene essentiality across the organisms’ lineages. Essential *P. falciparum* genes were more highly conserved across *Plasmodium* spp. (Fig. 5D and fig. S8A), are more likely to have a paralog (Fig. 5E and fig. S8B), and encode genes with less genetic variation among *P. falciparum* clinical isolates (Fig. 5F and fig. S8C). Of the *P. falciparum* genes studied here, 2,083 and 1,998 have orthologs in the more distantly-related parasites *Toxoplasma gondii* (6) and *P. berghei* (29), respectively. Overall, there was a strong correlation in gene function between species, particularly with genes predicted as essential (Fig. 5G and H and fig. S8D and E). Genes that are dispensable in *P. falciparum* were also more likely to be dispensable in *P. berghei*. By contrast, the correlation between the dispensable genes of *P. falciparum* and *T. gondii* was weaker, indicating that genus-specific gene functions are enriched for dispensability. Overall, these comparisons show that while phenotype classifications from the large-scale mutagenesis screens in apicomplexan species are also broadly predictive of the *P. falciparum* phenotype classifications, the closer evolutionary relationship of *P. berghei* provides a higher level of sensitivity and specificity for the orthologs that could be disrupted (Fig. 5I and fig. S8F).

Consistent with this assessment that non-mutable genes represented core essential functions in *P. falciparum*, GO enrichment analysis indicated genes with functions associated with translation, RNA metabolism, and cell cycle control were more likely to be essential, whereas genes associated with protein phosphorylation, virulence factors, and transcription were more likely to be dispensable (Fig. 6A). Comparing the relative number of essential, versus dispensable, genes within specific GO biological processes, molecular functions, and cellular components provided a ranking of the essentiality of potential druggable targets and pathways (Figs. 6A-C, figs. S9A-C and table S8). For example, nearly all genes annotated as being involved in ubiquitin-dependent degradation processes were experimentally defined as essential, while individual genes linked to microtubule motility and antigenic variation are experimentally defined as dispensable based on MIS. RNA metabolism and translation-related processes are also highly essential, supporting emerging evidence for the importance of post-transcriptional and translational control (30).

**Essentiality of the proteasome-degradation pathway**

Recent genetic analysis of artemisinin-combination therapy (ACT) resistance has linked drug resistance to cellular stress-response mechanisms involving the *P. falciparum* ubiquitin/proteasome system (31, 32). Genes of the proteasome-degradation pathway were well represented in the *piggyBac* insertion set, and 54 of the 72 genes could be classified as essential genes based on our data (fig. S10A), further strengthening the priority of the proteasome degradation pathway as a high-value druggable target (33). We validated this link between ART sensitivity and proteasome inhibition sensitivity using a set of single-insertion *piggyBac* mutants previously defined by chemogenomic profiling to be part of an ART-sensitivity cluster, including a mutant of the K13 Kelch propeller gene (16, 34). We found ten-fold increased sensitivity to the proteasome inhibitor Bortezomib in mutants of the ART-sensitivity cluster; significantly different relative to wild-type NF54 or mutants with distinct chemogenomic profiles (fig. S10B-E), providing additional support for the association between the ART mechanism of action and proteasome function.
Summary

In conclusion, our genome-wide saturation mutagenesis screen in the major human pathogen, *P. falciparum* has defined genes essential for parasite survival during the blood stage and provided critical functional data for prioritizing high-value drug targets and pathways. The complete characterization of the essential genome with high-throughput saturation mutagenesis will help open new frontiers for antimalarial therapeutic research. The methodology also opens the way for new systematic functional screens for other phenotypes, such as transmission and cytoadherence.

Materials and Methods

Parasite culture

All *P. falciparum* parasite NF54 and piggyBac mutants were cultured in complete RPMI 1640 with 5% hematocrit (medium containing 0.5% Albumax II, 0.25% sodium bicarbonate and 0.01 mg/ml gentamicin). All parasite cultures were maintained by standard methods (35).

piggyBac Transfection (36)

High-efficiency transfection (96-well plate method) was carried out on NF54 schizonts purified by magnetic column (MiltenyiBiotec, CS column) using a transposon plasmid (pLBacII-HDH, containing selection marker human *dhfr*) and a transposase-expressing helper plasmid (pDCBH) (19). RBCs were first loaded with plasmid DNA by electroporation using Gene PulserXCell+CE Module (BioRad). A total of 10 million schizonts, 1200 μg plasmid DNA and 600 μg helper plasmid DNA were used per 96-well plate to achieve maximal transfection efficiency. The drug WR99210 (final concentration 2.5 nM) was added to each plate for selecting transfected parasites, using a robot (Integra VIAFLO 96) to change media with drug every day for 5 days. Transfections were performed in batches over the course of 4 to 6 weeks. Each 96-well plate was cultured in triplicate and cryopreserved in duplicate. One hundred 96-well plates were used to generate the large pools for sequencing (fig. S1A).

For maximum recovery of unique mutants, the transfected culture was distributed into 96-well plates by robot (200μl each well) and grown under two rounds of drug selection. Wells were screened for parasites by PCR and Giemsa-stained thin blood films. >50% of wells became parasite-positive within three weeks. Each positive well with ≥9 unique mutants (one well = one “Mixed Population”, or MP) was cryopreserved in duplicate (two 96 well plates). Additional 96-well plates were used to generate the pools that were then cultured in a T75 flask and harvested after two cycles growth by standard methods for genomic DNA isolation for QiSeq (37). Although mutants grow at different rates, the pools can be expanded for at least 12 asexual life cycle generations (“cycles”) without significant loss in detectable diversity. Importantly, each mutant pool can be regenerated and cloned from their original MPs.
Datasets of reference genomes, transcriptome data and epigenetics data

The NF54 reference genome can be downloaded from (ftp://ftp.sanger.ac.uk/pub/project/pathogens/Plasmodium/falciparum/NF54/Assembly/V1_morphed/). The QIseq sequencing mapping followed previously published methods (37). Ortholog counts, paralog counts, and non-synonymous/synonymous (NonSyn/Syn) ratios can be downloaded from PlasmoDB (http://plasmodb.org/plasmo/) (38). Previously published RNA-seq expression data (26, 39) are used in computational analysis. Formaldehyde-Assisted Isolation of Regulatory Elements (FAIRE-seq) data of seven time points (0, 6, 12, 18, 24, 30 and 36 hour) were used to measure chromatin accessibility (40). The motif information of transcription factor ApiAP2 binding sites is available in previously published work (41). The genome synteny across five Plasmodium species is available in published work (25).

Identification of > 38,000 mutants carrying piggyBac insertion

The QIseq sequencing generated a total of 3,301,112 raw reads. First, all the raw reads were filtered by the criteria of matching expected transposon target site ‘TTAA’. This filtering step yielded 2,042,147 reads, which is >6-fold genome coverage. The insertion site QIseq signal was calculated as the reads count of the filtered QIseq reads. To compare the insertion signals between different QIseq runs, we normalized for each insertion i as

$$S_i = \frac{C_i}{\sum_{j \in R} C_j} \times 250000 \qquad (1)$$

, where $S_i$ represents the signal of the insertions i with reads counts $C_i$ in run $R$. All the piggyBac insertions with reads in both 5’ and 3’ ends were treated as true insertions by using an extremely high level of accuracy in target site (target motif TTAA ~ 99%). To identify true insertions with reads only in single 5’ or 3’ ends, we used a method that selected the most accurate signal cutoff based on the ratio of the number of input reads/ the number of output disrupted genes. We computed the number of gene CDSs targeted by the insertions upper cutoff (i.e. more accurate). We noticed that there was a single large and sharp increase in the number of CDSs targeted by the insertions with reads number increasing from cutoff 2.0 to cutoff 2.3 (Fig. S2E, p > 0.05 compared with changing another cutoff). The result indicated a large amount of false positive may be included when setting cutoff ≥2.3. Therefore, the normalized reads count = 2.3 was set as the lower bound for identifying true insertions.

Mutagenesis saturation computational validation

To evaluate the levels of whole-genome mutagenesis and eventual saturation, we used a computational method based on sampling. First, a specific number of insertions were randomly extracted and the fraction of genome elements targeted were counted. This procedure was repeated 1000 times for each query insertion number in fig. S1C. A log-based curve was observed by plotting the mutational events and the median number of genome elements targeted by insertions. The plateauing of the curve at about 20,000 mutational events indicated that a very small number of new genome elements could be targeted even with very large amount of insertions. Importantly, the real data-based sampling curve was
very similar to our independent mathematical predictions of Negative Binomial distributions prior to the start of the saturation mutagenesis experiments (fig. S1D). Together, both our mathematical model and real-data computational sampling show that > 33,000 mutants represent saturation-level mutagenesis of the entire set of protein-encoding genes in the *P. falciparum* genome.

**Sampling methods for accounting for fragment length differences and nucleoside composition bias**

For genes in the *P. falciparum* genome, 5’ upstream and 3’ downstream intergenic regions displayed similar TTAA densities (median number of TTAA sites per 100bp were 1.82 and 1.89, respectively). However, they have significantly different region length distributions (median values are 1681bp and 926bp in 5’ and 3’ direction, respectively; p< 0.001 Wilcoxon test). Therefore, there are different numbers of genomic TTAA sites in 5’ and 3’ intergenic regions, as an intrinsic feature of the *P. falciparum* genome. To account for this region-length and total-TTAA bias in our QiSeq calculations, the TTAA sites in 3’ intergenic regions were sampled based on the TTAA number in 5’ intergenic regions, so that our comparison was based on two identical background target-site distributions. Distribution of insertions and TTAA sites (Fig. 1E) were plotted based on these sampling data.

**Mutagenesis index score (MIS) calculation**

To quantify the mutability of every protein-coding gene in the *P. falciparum* genome, we first checked the number of piggyBac insertions located inside the gene CDS regions. To keep the criteria stringent, the insertions located on the very end (distance from insertion to TSS > 99% of the CDS length) of the CDS were not considered, due to a significantly higher number of insertions at the last 1% of the CDS compared with other regions of the CDS (fig. S2F). We calculated the initial score MIS of gene $MS_g$ based on the equation

$$MS_g = \log\left(\frac{N_g + 1}{D_g}\right)$$  \hspace{1cm} (2)$$

where $N_g$ represents the number of insertions on gene $g$. $D_g$ is the TTAA density of the gene $g$, which could be represented as the number of TTAA per kb of the CDS. We found that the MIS could be decomposed into two mixed Gaussian distributions (Fig. S11). We reasoned that dispensable and essential genes exhibit different MIS distribution. Therefore, a binary variable $\pi_g$ was used to model whether the gene $g$ is dispensable or not: $\pi_g = 1$ corresponds to dispensable and vice versa. The probability of observing a $MS_g$ of gene $g$ is mixture of two Gaussian distributions:

$$P(MS_g) = \sum_{\pi_g = 1, 0} P(\pi_g) N\left(MS_g \mid \mu_{\pi_g}, \sigma_{\pi_g}\right) \hspace{1cm} \sum_{\pi_g = 1, 0} P(\pi_g) = 1$$  \hspace{1cm} (3)$$
Subsequently, the EM algorithm was implemented to search for the optimized value of parameters. The posterior distribution after the EM optimization procedures could be calculated as:

\[
P(\pi_g = 1|MS_g, \Theta) = \frac{P(\pi_g = 1)N(\mu_{\pi_g} = 1, \sigma_{\pi_g} = 1)}{\sum_{\pi_g = 1, 0} P(\pi_g)N(\mu_{\pi_g}, \sigma_{\pi_g})} \tag{4}
\]

, where \(\Theta\) is the parameter space. The MIS was defined as the posterior distribution to be dispensable.

**Mutagenesis fitness score (MFS) calculation**

For the mutants that were subjected to competitive growth essay, we calculated MFS to represent the comparative growth fitness of a mutant. The QIseq sequencing reads distribution reflects the fitness of unique piggyBac mutants in the competitive-growth assays. The start and the end of the growth assay could be represented as \(t_0\) and \(t\). At the end of the assay, \(a_{t,v}^{m,v}\) indicated the relative abundance of each mutant \(m\) targeted into the CDS of gene \(g\) in sample \(v\). Therefore, the abundance of the mutant of the gene \(g\) at the end of the assay could be represented as \(\sum_{m,v} a_{t,v}^{m,v}\). The \(\sum_{m,v} a_{t,v}^{m,v}\) was directly proportional to the QIseq reads number \(\sum_{m} r_{t,g}^{m,v}\). Here, \(r_{t,g}^{m,v}\) represents the normalized reads number of mutant \(m\) targeting gene \(g\) on CDS, which is \(\sum_{m,v} a_{t,v}^{m,v} \sim \sum_{m,v} r_{t,g}^{m,v}\). The QIseq reads starting with ‘TTAA’ were used to measure \(r_{t,g}^{m,v}\). To normalize different \(\sum_{m,v} r_{t,g}^{m,v}\) in different sample \(v\), the average value of \(\sum_{m,v} r_{t,g}^{m,v}\) was calculated as

\[
\frac{\sum_{v} \sum_{m} r_{t,g}^{m,v} \sum_{m} I(m \ on \ g)}{\sum_{m} I(m \ on \ g)} \tag{5}
\]

Here, \(I(\cdot)\) is the indicator function. \(\sum_{m} I(m \ on \ g)\) was the number of mutants targeting gene \(g\) in all samples. At the start of the assay \(t_0\), the relative abundance of each mutant equal to the background TTAA density \(D_g\). Finally, the MFS could be calculated as:

\[
MFS_g = \log \left( \frac{\sum_{v} \sum_{m} r_{t,g}^{m,v} \sum_{m} I(m \ on \ g)}{D_g} \right) \tag{6}
\]

Therefore, the MFS calculation based on individual mutant abundance is a proxy for competitive growth fitness in *in vitro* blood stage.
GO analysis

All the Gene Ontology (GO) terms were downloaded from (http://plasmodb.org/plasmo/). For each specific GO term \( g \), the MIS distribution in the term \( g \) could be represented as \( MIS_g \). The number of genes in term \( g \) is \( NUM_g \). We asked whether the genes in the term \( g \) are more prone to be dispensable or essential. To answer this question, we compared the median value of \( MIS_g \) with sampled data distribution as background. The genes with the same number of \( NUM_g \) were sampled out and represented as \( S_g \). This procedure was repeated 1000 times and the median MIS of each \( S_{g,t} \) (\( t \in 1, 2, 3..1000 \)) was calculated as \( MIS_{S_g,t} \). The p-value was calculated as

\[
P_g = \begin{cases} \sum_{t=1}^{1000} I(MIS_g > MIS_{S_{g,t}}) / 1000 & (MIS_g > 0.5) \\ \sum_{t=1}^{1000} I(MIS_g < MIS_{S_{g,t}}) / 1000 & (MIS_g < 0.5) \end{cases} \tag{7}
\]

Here, \( I(\cdot) \) is the indicator function.

Half-maximal inhibitory concentration (IC50) estimation

To calculate IC50, the dose-response data (e.g., drug concentrations \( c_1, c_2, \ldots c_n \) and growth inhibition \( q_1, q_2, \ldots q_n \)) were used to fit a Hill Equation.

\[
q = U + \frac{U - D}{1 + 10^{(c - \log C)B}} \tag{8}
\]

where \( c \) was drug concentrations in logarithmic form. The parameter C was the estimate of IC50. The goodness of fit was calculated from dose-response data and the Hill Equation.

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

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References and Notes:

37. Bronner IF et al., Quantitative Insertion-site Sequencing (QIseq) for high throughput phenotyping of transposon mutants. Genome research, (2016).
Fig. 1. A genome-wide saturation mutagenesis screen for *Plasmodium falciparum*.
(A) Chromosomal map displays 38,173 piggyBac insertion sites from all mutants evenly
distributed throughout the genome. (B) High-resolution map of a ~50 KB region of
chromosome 13 depicts an essential gene cluster, including K13, flanked by dispensable
genes with multiple CDS-disrupting insertions. (C) High-resolution map of a ~20 KB region
without insertions includes three conserved genes of unknown function (PF3D7_1232700,
PF3D7_1232800, PF3D7_1232900) and a putative nucleotidyltransferase (PF3D7_1232600)
(see also fig. S5). (D) A plot of all piggyBac insertions revealed significantly fewer
insertions were recovered from exon-intron regions compared to the proportion of available
TTAA sites (see also fig. S1D) \(p < 2.2\times 10^{-16}\), Fisher’s Exact test). (E) Density of piggyBac insertion-site distribution revealed threefold fewer insertions recovered in transcriptional regions (blue) than intergenic 5’ (yellow) and 3’ (green) regions, depicted as relative distance upstream and downstream to a gene, respectively. (F) This study determined that under ideal culture conditions for asexual blood-stage growth, 38% of genes in the \(P. falciparum\) genome have mutable CDSs, while 62% of genes have non-mutable CDSs.
Fig. 2. Identification of dispensable and essential genes through Mutagenesis Index Score (MIS) and Fitness Score (MFS).

(A) The Mutagenesis Index Score (MIS) rates the potential mutability of *P. falciparum* genes based on the number of recovered CDS insertions relative to the potential number that could be recovered. Genes known as dispensable or essential are highlighted. (B) MIS violin plots of GO processes grouped from lowest to highest dispensability, according to gene functional annotations. (C) MIS plots and (D) high-resolution chromosome maps highlighting important genes of interest for RNA metabolism [−20kb, +20kb] (see also fig. S4 for MIS...
plots of other genes of interest. (E) The Mutagenesis Fitness Score (MFS) estimates the relative growth fitness cost for mutating a gene based on its normalized QIseq sequencing reads distribution. (F) MIS has significant correlation to MFS (Pearson’s R = 0.67, p < 2.2e-16 compared with permutation). (G) The first and second MFS quartiles were comprised primarily of non-mutable genes, the 4th quartile was comprised mostly of mutable genes, and 3rd quartile had nearly equal numbers of both.
Fig. 3. Validation of mutagenesis score through phenotype screen.

(A) Competitive growth assays of asexual blood-stage growth under ideal *in vitro* culture conditions: phenotypes of four independent mixed-population pools grown for three cycles confirmed losers (left, bottom quantile) and winners (right, top quartile) had significantly different MIS. (B) Overall rank-ordered plot of competitive growth phenotypes shows losers and winners. (C) Competitive growth ‘losers’ had significantly lower MIS and MFS, respectively, validating MIS and MFS as predictor of gene essentiality and dispensability. (D) Circos plot from outer to inner shows the distribution of all piggyBac insertions, MIS
(pink indicates MIS < 0.5, while blue is >0.5), CDS insertions, and MFS along each chromosome of *P. falciparum* genome. (E) Violin plots indicate non-mutable genes had significantly lower MIS and MFS (**** represents Wilcoxon p < 2.2e-16).
Fig. 4. Chromosomal syntenic breakpoints are enriched in dispensable genes. 
(A) Genes within conserved syntenic blocks have significantly lower MIS and MFS (Wilcoxon p < 2.2e-16). Syntenic genes or “syntenic block” is defined as at least three genes in the same order on the same chromosome as their orthologs in another species within a 25-kb search window. (B and C) Scatter plots show the insertion site enrichment along two syntenic breakpoints (Ch13:2,110,000 –2,135,000, Chr10: 642000–666000). Each gap in synteny (white area) is enriched for piggyBac insertions while flanked by essential regions (green shading); black boxes represent the location of CDS. (D and E) Circos plots indicate the syntenic blocks of P. falciparum in relation to other Plasmodium spp. (P. berghei, P. chabaudi, P. knowlesi, P. vivax).
Fig. 5. Distinct biological process and evolutionarily conservation segregate the tendency of dispensable and essential genes.

(A) The genes with lowest FPKM expression value (first quantile) among different stages were enriched for dispensable genes (Wilcoxon P < 2.2e-16 compared with other quantiles) (26). The expression level cut off is set at 20 FPKM. (B) Non-mutable essential genes had significantly higher expression value for blood-stage development. (C) The group of trophozoite-stage genes had the highest proportion of essential genes (red) whereas gametocyte genes had the highest proportion of dispensable genes (blue) (Wilcoxon p < 1e-12). (D to F) Characteristics of essential genes significantly different from dispensable genes include: (D) 1:1 ortholog conserved among Plasmodium spp; (E) absence of paralogs; and (F) reduced rate of non-synonymous to synonymous SNPs. Bars indicate the group median (**** indicates Wilcoxon p < 2.2e-16). (G and H) Essential genes reported in (G) Toxoplasma and (H) P. berghei showed significantly lower MIS in this mutagenesis screen of P. falciparum (**** indicates Wilcoxon p < 2.2e-16). (I) Plot of Receiver Operating Characteristics (ROC) indicate the level of retention of essential genes across species. The

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MIS of *P. falciparum* more strongly correlates with the essentiality phenotype of *P. berghei* than *Toxoplasma*.
Fig. 6. Differentiating dispensable and essential genes and discovering high-priority druggable targets and pathways.

(A) Functional annotations of biological processes are represented by the p-value and the X-axis shows the fraction of the genes with MIS > 0.5. Each GO term is assigned a p-value on the Y-axis to represent the tendency to be essential or dispensable. Essentiality is indicated on a spectrum of red (essential) to blue (dispensable) and circle sizes indicate the GO term enrichment. (B and C) Boxplot of (B) molecular processes and (C) cellular components shows the MIS distribution generated by 1000× sampling of the number of genes in the
query GO-term category. Left (red) and right (blue) triangles indicate GO terms with significantly lower or higher MIS (p-value < 0.05 compared to background), respectively; the heatmap represents the essentiality defined as the fraction of genes per GO term with MIS > 0.5.