Genome-wide regulatory dynamics of translation in the *Plasmodium falciparum* asexual blood stages.

**Permalink**
https://escholarship.org/uc/item/10k3q7ns

**Authors**
Caro, Florence  
Ahyong, Vida  
Betegon, Miguel  
et al.

**Publication Date**
2014-12-10

**DOI**
10.7554/elife.04106

Peer reviewed
Genome-wide regulatory dynamics of translation in the *Plasmodium falciparum* asexual blood stages

Florence Caro††, Vida Ahyong††, Miguel Betegon1,2, Joseph L DeRisi1,2*

1Department of Biochemistry and Biophysics, University of California, San Francisco, San Francisco, United States; 2Howard Hughes Medical Institute, University of California, San Francisco, San Francisco, United States

**Abstract** The characterization of the transcriptome and proteome of *Plasmodium falciparum* has been a tremendous resource for the understanding of the molecular physiology of this parasite. However, the translational dynamics that link steady-state mRNA with protein levels are not well understood. In this study, we bridge this disconnect by measuring genome-wide translation using ribosome profiling, through five stages of the *P. falciparum* blood phase developmental cycle. Our findings show that transcription and translation are tightly coupled, with overt translational control occurring for less than 10% of the transcriptome. Translationally regulated genes are predominantly associated with merozoite egress functions. We systematically define mRNA 5′ leader sequences, and 3′ UTRs, as well as antisense transcripts, along with ribosome occupancy for each, and establish that accumulation of ribosomes on 5′ leaders is a common transcript feature. This work represents the highest resolution and broadest portrait of gene expression and translation to date for this medically important parasite.

**Introduction**

The transcriptome of the intraerythrocytic developmental cycle (IDC) of *P. falciparum* is characterized by a continuous cascade wherein the expression of the majority of genes is maximally induced once per cycle and their timing correlates well with the timing for the respective protein’s biological function (Bozdech et al., 2003). The apparent lack of dynamic transcriptional regulation suggested that complementary post-transcriptional mechanisms could play an important role in the regulation of parasite gene expression (Hughes et al., 2010). This is a reasonable assumption, given that global or gene-specific translational regulation of gene expression is a mechanism that allows fast adaptations during drastic changes in environmental conditions as well as during rapid transitions in developmental programs. Indeed a few examples of translational control in *Plasmodium* have been reported. In sporozoites present in the mosquito salivary gland, phosphorylation of the eukaryotic translation initiation factor eIF2α by the kinase IK2, inhibits translation and causes accumulation of mRNAs into granules. Translational repression is alleviated by eIF2α phosphatase during the transition into the mammalian host, allowing parasites to transform into the liver stages (Zhang et al., 2010). Similarly, PK4 kinase activity leads to the reduction of global protein synthesis through phosphorylation of eIF2α in schizonts and gametocytes and is essential for the completion of the parasite’s erythrocytic cycle (Zhang et al., 2012). Gene-specific translational regulation has also been observed in *P. falciparum* and is mediated by cis-acting sequences in combination with RNA-binding proteins. For example, dihydrofolate reductase-thymidylate synthase (DHFR-TS) binds within the coding region of its own cognate mRNA to repress translation (Zhang and Rathod, 2002) and antifolate treatment has been shown to relieve this repressive effect without alteration of mRNA levels.
In Plasmodium berghei, storage of translationally repressed mRNAs prior to fertilization is mediated by mRNA binding via the RNA helicase DOZI and the Sm-like factor CITH (Mair et al., 2006, 2010). Upstream open reading frames (uORFs) found on 5′ UTRs of transcripts have been reported to regulate the translation of specific genes (Morris and Geballe, 2000). In P. falciparum, the only uORF described and functionally characterized to date is a 120 codon region upstream of the var2csa (PFL0030c) coding region, a unique variant of the surface antigen PfEMP1 that mediates adhesion to placenta in pregnant women (Amulic et al., 2009). In this case, translation of the uORF modulates repression of var2csa translation. Aside from these examples, the extent to which global and gene-specific translational control operates in P. falciparum during the IDC remains sparse.

Since the P. falciparum genome was fully sequenced (Gardner et al., 2002), several large-scale studies have provided detailed insights into the expression of genes and proteins across the parasite’s life cycle. Parallel mass spectrometry-based proteomics and genome-wide expression profiling revealed differences between mRNA abundance and the accumulation of the corresponding protein, supporting the notion that post-transcriptional regulation of gene expression is at play in this parasite (Le Roch et al., 2004; Nirmalan et al., 2004b; Foth et al., 2011). These methods, however, are limited in their ability to measure low abundance proteins and do not capture the underlying relationship between transcriptional activity and translational efficiency. More recently, polysome profiling was used to monitor discrepancies between polysome-associated and steady-state mRNAs in 30% of the P. falciparum blood stage transcriptome (Bunnik et al., 2013); however, this approach does not reveal...
the precise localization of the ribosomes, and thus can not be used to accurately assess the translational efficiency of a given mRNA (Ingolia, 2014).

Here, we adapted the ribosome profiling technique (Ingolia et al., 2009) to describe the translational dynamics of the P. falciparum asexual blood stage transcriptome. We simultaneously evaluate mRNA abundance, gene structure, ribosome positioning, and translational efficiency for genes expressed through five stages of the IDC. We demonstrate that the data are highly reproducible, and we find that the translational efficiency of the majority of mRNAs expressed follows a narrow distribution, exhibiting a tight coupling between transcription and translation. Only 10% of the genes expressed deviate from this trend and are translationally up- or down-regulated. We found a surprising amount of ribosome density associated with 5′ leaders of transcripts particularly in genes with functions associated with merozoite egress and invasion. Overall, the precision and depth of the dataset presented herein add significantly to our understanding of P. falciparum gene expression by linking transcriptional and translational dynamics throughout the blood stages.

Results

Overview of ribosome profiling in P. falciparum asexual blood stages

To create whole-genome, high-resolution profiles of mRNA abundance and translation during in vivo blood stage development of P. falciparum, we adapted the ribosome profiling technique described by Ingolia et al. (2009). Ribosome profiling is based on the deep sequencing of ribosome protected mRNA fragments obtained by nuclease digestion of polysomes, cycloheximide-arrested ribosomes bound to mRNA. These fragments represent the exact location of the ribosome at the moment the sample was harvested. Five stages representative of the 48-hr IDC of P. falciparum were harvested for both mRNA and polysome isolation; ring, early trophozoite, late trophozoite, schizont stages, and purified merozoites. To assess the reproducibility of the data, we harvested independent biological replicates of each stage. Polysomes were isolated in the presence of the translation elongation inhibitor cycloheximide, then nuclease digested to produce monosomes, and sedimented by centrifugation on a sucrose gradient (Figure 1 and Figure 1—figure supplement 1). To minimize isolation of RNA fragments bound by proteins other than 80S ribosomes, RNA was extracted only from the fractions of the sucrose gradient containing the monosome peak. The resulting ~30 nt fragments of RNA, corresponding to ribosome footprints, were processed into strand-specific deep sequencing libraries in parallel with the mRNA samples, fragmented to ~30 nt for consistency. Despite the unusually high AT content of the P. falciparum genome, over 92% of all 30 nt sequenced reads, derived from coding sequences (CDSs), mapped uniquely to the genome (Figure 1—source data 1 and Figure 1—figure supplement 2).

To quantitatively obtain mRNA abundance and ribosome footprint density measures, we calculated rpkMs (reads per kilobase of exon model per million reads mapped, as in Mortazavi et al. (2008) for each gene. We established the minimum number of mRNA reads sequenced per coding region (rM; reads per million reads mapped) required to confidently include genes in downstream analyses, to be ≥32 rM (Materials and methods, Figure 1—figure supplement 3). Using this conservative threshold, 3605 genes qualified for further analysis. Between biological replicates, Pearson correlation values were consistently high, ranging from r = 0.94 to r = 0.99 (Figure 2A), highlighting the quality and reproducibility of our data. In addition, we compared the RNA-seq transcriptome of the five stages sampled to our previously published transcriptome data set, originally generated using long oligo microarrays (Bozdech et al., 2003). The RNA-Seq transcription profiles of the set of genes shared by the two data sets (n = 1829) were highly correlated (average r = 0.7) to the corresponding 11, 21, 31, 45, and 2 hr post merozoite invasion time points of the microarray data set, despite the use of different methodologies (microarray vs RNA-seq) and the use of different P. falciparum strains (HB3 vs W2, respectively). Because of the higher sensitivity of RNA-seq, we were able to accommodate an additional 743 genes into the cascade-like transcriptome extending it to a total of 3110 genes (Figure 2B, Figure 2—source data 1). The remaining 495 genes in our RNA-seq data set lacked sufficient variation over the five time points for inclusion within the phaseogram. These genes, referred to as non-phanic genes, are nevertheless included in all analyses.

Gene expression and translation are tightly coupled during the P. falciparum IDC

While RNA-Seq reveals the abundance and architecture of individual mRNAs, ribosome profiling provides a complementary and quantitative measure of mRNA translation. Ribosome occupancy along
the CDS results in a profile that indicates the timing and magnitude of translation of a given mRNA, thus quantitatively delineating regions of each mRNA molecule that are actually bound by 80S ribosomes (Ingolia et al., 2009). To inspect translation on a genome-wide scale, ribosome density...
values of each gene expressed in the data set were organized in the same order as the transcriptome. The translational profile of each gene displayed a cascade-like quality strikingly similar to the transcriptome (Figure 2B). Much like mRNA abundance, translation of phasic genes reaches a single maximum and a single minimum during the IDC. To determine the exact level of correlation between transcription and translation, we directly compared mRNA and ribosome footprint density measurements (Figure 2C). In general, translation is tightly correlated with transcription for all phasic and non-phasic genes in rings ($r = 0.85$), early trophozoites ($r = 0.93$), late trophozoites ($r = 0.91$), schizonts ($r = 0.89$), and purified merozoites ($r = 0.86$). This indicates that when an mRNA is detected in one stage it is associated proportionally with ribosomes within the same stage. An example pair of genes is shown in Figure 3A. Here, mRNA abundance profiles of eukaryotic translation initiation factor eIF2 gamma subunit (PF14_0104) and the conserved protein PF14_0105, show that peak mRNA abundance for these two genes occurs at two different stages, early and late, respectively. Examination of ribosome occupancy of both genes reveals a ribosome density accumulation profile within the coding sequence that mirrors their respective mRNA profiles. As for the majority of genes, ribosome footprint density and mRNA abundance for these two genes are highly correlated ($r = 0.98$ and $r = 0.93$ for PF14_0104 and
PF14_0105, respectively), indicating that mRNA translation occurs proportionally during the same stages at which these genes are transcribed (Figure 3B; Supplementary file 1). Globally, 77% of genes expressed in at least three stages of the IDC display high Pearson correlation ($r \geq 0.7$) between transcription and translation. 

**Figure 3.** Transcription and translation are highly correlated. (A) Ribosome footprint (green) and mRNA (blue) coverage profiles of two neighbor genes, the eIF2 gamma subunit (PF14_0104) and the conserved protein PF14_0105 (CDS, white boxes; HMM-defined UTRs, black lines) in rings (R), early trophozoites (ET), late trophozoites (LT), schizonts (S), and merozoites (M). Mappability = mappability score at that position; range 0 (white) to 30 (black). $rM =$ coverage (reads per million reads mapped). (B) mRNA and ribosome footprint density of the genes in (A) correlate during development. $r =$ Pearson correlation coefficient between ribosome footprint density and mRNA abundance of each gene.

DOI: 10.7554/eLife.04106.010

The following figure supplement is available for figure 3:

**Figure supplement 1.** mRNA abundance and ribosome footprint density are highly correlated for the majority of genes expressed during the IDC.

DOI: 10.7554/eLife.04106.011
mRNA abundance and translation (Figure 3—figure supplement 1). Thus, our genome-wide analysis of translation establishes that for the majority of genes expressed during the IDC, transcription and translation occur proportionally.

**Ribosome profiling reveals instances of translational control of gene expression**

Ribosome profiling allows the monitoring of translation rates through the simultaneous quantitative measure of mRNA abundance and ribosome density on mRNAs. The ratio of the footprint rpkM to the mRNA rpkM for any given gene represents its relative translational efficiency (TE) (Ingolia et al., 2009). To assess the dynamics of translational control and detect variations in control within and between developmental stages, we calculated the relative TE of all expressed genes in our data set (Figure 4A). The shape and the range of TE distributions obtained for each stage sampled is comparable to those seen in other eukaryotes (Ingolia et al., 2009; Dunn et al., 2013). Absolute mean translational efficiencies in all stages (log2TE μRings = −0.43, log2TE μETrophi = −0.56, log2TE μLtrophi = −0.31, log2TE μSchizonts = −0.16 and log2TE μMerozoites = −0.68) had a maximum difference of 1.47-fold observed
between early trophozoites and schizonts. Translational efficiencies display a roughly 100-fold range in absolute values in each of the stages with the exception of the ring and merozoite stages, which exhibit more extreme values. In these stages, the distribution of absolute TE values displays an approximately fourfold larger spread than in early trophozoites, late trophozoites, or schizonts (Figure 4A, Figure 2—source data 1). In rings the gene with the largest TE is the merozoite surface protein 9 (PFL1385c, log$_2$TE = 4.1) and the gene with the lowest TE is the FIKKK family serine/threonine protein kinase (PF14_0734, log$_2$TE = −5.1). In merozoites the largest and lowest TE values correspond to the serine repeat antigen 5 (SERAS5, PFB0340c, log$_2$TE = 4.0) and the alpha adenyl cyclase (PF14_0788a-c, log$_2$TE = −4.7), respectively.

To determine the contribution of translational efficiency to the dynamic range of gene expression, we examined the genes lying at the extremes of the TE distribution. For the purpose of this analysis, genes with a translational efficiency of two standard deviations above or below the mean in any of the stages were considered translationally up- or down-regulated, respectively. A total of 301 genes, 8.3% of the transcriptome, are translationally regulated by this metric, with 124 genes translationally down-regulated and 177 genes translationally up-regulated (Figure 4B, Figure 2—source data 1). The timing of maximum mRNA expression does not influence TE for either of these two groups. Translational efficiencies remain high for the translationally up-regulated and low for the translationally down-regulated genes in all the stages at which they are expressed, regardless of the stage of peak mRNA abundance, suggesting that translational efficiency is largely, but not completely, programmed by the mRNA sequence itself, rather than global factors. For example, translational efficiency of the merozoite surface protein 6 (MSP6, PF10_0346) remains high (log$_2$TE ≥ 2) across all stages irrespective of variations in its mRNA abundance. In contrast TE values for the eukaryotic initiation factor 2alpha kinase 1 (IK1, PF14_0423) are among the lowest measured despite high mRNA abundance across all stages (Figure 4C).

An examination of the 124 translationally down-regulated genes yielded some expected, and in some cases, unexpected findings. As would be expected, two pseudogenes, the ring-infected erythrocyte surface antigen 2 (RESA-2, PF11_0512) and reticulocyte binding protein homologue 3 (PfRh3, PFL2520w), represent a clear example of low translational efficiency. The PfRh3 pseudogene ribosome profile shows that translation of the 5′ end of this transcript occurs up until the encounter of several in-frame stop codons, causing the reduction in ribosome density from this point on (Figure 5A, Figure 5—figure supplement 1). This suggests that a truncated version of the PfRh3 protein is being produced in the W2 strain studied here. Evidence for peptides corresponding to the 5′ end of PfRh3 has been found in gametocytes and sporozoites (however not during the asexual stages) using mass spectrometry (Florens et al., 2002; Lasonder et al., 2002). We note that low levels of ribosomes can still be detected along the full length of this transcript in schizonts and merozoites. Whether these footprints derive from a low level of stop-codon read-through or accumulate via another unknown mechanism remains to be determined.

Ring-infected erythrocyte surface antigen 2, RESA2 (PF11_0512) was first described as a pseudogene based on the presence of an internal stop codon (Cappai et al., 1992). Since then, transcription of this gene has been demonstrated both in vivo (Vazeux et al., 1993) and in vitro (Bozdech et al., 2003). RESA-2 is transcribed but poorly translated in rings, early trophozoites and merozoites (log$_2$TE −3.2, −2.7, −2.9, respectively). Accordingly, the ribosome profile of this gene in merozoites shows a general depletion of ribosomes along the CDS (Figure 5—figure supplement 2). In rings, ribosome density diminishes at the second exon. To validate the RESA2 gene model, we used genomic DNA sequencing data derived from the P. falciparum W2 strain used in this study. We found that 69% (n = 151) of reads mapping to this locus support a single base deletion that creates a premature stop codon exactly at the site of ribosome footprint drop-off (Supplementary file 2). These data suggest that RESA2 is transcribed and actually translated into a shorter protein product of 461 amino acids. Whether or not the protein product is functional or undergoes post-translational degradation remains to be determined.

In addition to expected instances of translational regulation, our data permit the discovery of previously uncharacterized translational regulation, especially at the extremes of the TE distributions. One of the most notable examples of translational silencing is the elf2α kinase IK1 (PF14_0423) for which ribosome footprints accumulate at the 5′ leader and 3′ UTR but not on the CDS, resulting in an extremely low translational efficiency (log$_2$TE = −3.6) despite relatively high transcript abundance across all stages (Figure 5B). The mechanism by which this gene is maintained in a translationally
Figure 5. Translationally down-regulated genes have decreased CDS ribosome density. (A) Ribosome footprint (green) and mRNA (blue) profiles of the PfRh3 pseudogene (PFL2520w) in merozoites (M). In the detail the bars above the gene model indicate AUG, stop, and any other codon, in green, red, and gray, respectively. Boxes indicate the mapping location of peptides identified by mass spectrometry in gametocytes and sporozoites (Florens et al., 2002; Lasonder et al., 2002). Reduction of ribosome footprint coverage occurs upon encounter of consecutive stop codons (extended red lines). (B) eIF2α kinase (PF14_0423) gene in rings (R) showing ribosome footprint accumulation on the 5' leader, 3' UTR, and low translational efficiency of the CDS. (CDS, white boxes; Figure 5. Continued on next page.
down-regulated state is unknown. Another example is the erythrocyte vesicle protein 1 (EVP1, PFD0495c) for which abundant transcript levels can be detected across all stages, with peak mRNA abundance occurring in rings and schizonts (Figure 5—figure supplement 2). Protein levels, however, have been shown to be undetectable (Tamez et al., 2008). Here, we find that translational efficiencies of this gene were low across all stages and lowest in rings and early trophozoites (log\(^2\)TE = −2.6 and −2.9, respectively) demonstrating that post-transcriptional regulation at the level of translation is, at least in part, responsible for its scarcity as a protein.

Thus, our ribosome profiling data set highlights instances of translational control of genes that may not be detected by proteomic methods. Indeed a search for mass spectrometric data showed no evidence for ~70% of genes in this category (Aurrecoechea et al., 2009). Including the aforementioned examples, our data set describes a total 124 translationally down-regulated genes (listed in Figure 2—source data 1) for which translational efficiency values lie at the lower extremes of the distribution.

Protein products of translationally up-regulated genes are likely to be abundant and readily detected using mass spectrometry. Previous proteomic studies show protein evidence in the blood stages for almost all (171 of 177) well-translated genes identified here (Aurrecoechea et al., 2009; Pease et al., 2013). Mass spectrometric evidence for the remaining six genes is either absent (PFL0245w, PFL2510w, PF11_0204) or has only been found in sporozoites (PFE1615c, MAL7P1.300, PF13_0069a). Despite the lack of proteomic data, our data indicate that these genes are both transcribed and translated during the blood stages of the parasite. Whether post-translational control points exist for these proteins is unknown.

Among the top ten most highly translated genes are proteins involved in merozoite egress and invasion MSP3, 6, 7, and 9 (merozoite surface proteins PF10_0345, PF10_0346, PF13_0197, and PFL1385c), serine repeat antigen 5 (SERA5, PFB0340c), and RAP1, 2, and 3 (PF14_0102, PFE0080c, and PFE0075c, respectively) (Figure 2—source data 1). Interestingly, 73 (41%) of all translationally up-regulated genes can be assigned to the repertoire of canonical functions for merozoite egress and invasion described to date (Yeoh et al., 2007; Blackman, 2008; Hu et al., 2010; Farrow et al., 2011). Strikingly, for all genes in this set, maximum mRNA abundance is found during the late stages of the IDC (69 schizont and 4 merozoite stage mRNAs) yet for the majority (50, 70%) peak translational efficiency occurs in rings. Consistent with this, peptides for most of these merozoite function proteins (58 of 73) have been detected in rings (Oehring et al., 2012; Pease et al., 2013). This mode of translational regulation whereby late stage transcripts are highly translated in rings was not exclusively limited to genes related to merozoite egress and invasion. We found evidence for an additional 14 genes with this profile, including, aquaglyceroporin (PF11_0338, log\(^2\)TE = 3.8), tubulin beta chain (PF10_0084, log\(^2\)TE = 1.8), and early transcribed membrane protein 2 (PFB0120w, log\(^2\)TE = 2.5).

Taken together these data demonstrate that transcription and translation are tightly correlated for the majority of genes expressed during the asexual life cycle of *P. falciparum* with few exceptions. These apply to a small subset of translationally down- and up-regulated genes for which translational efficiencies appear to be inherent properties of the mRNA, independent of changes in mRNA abundance. Genes in this category, especially those that exhibit high translational efficiencies, are enriched with functions associated with merozoite egress and invasion during the transition from late stages into rings.

**Ribosome occupancy of 5′ leaders is commonly found on genes expressed during the IDC**

Ribosome profiling provides position specific information along each transcript allowing the detection of changes in ribosome distribution on the mRNA and their relationship to translational efficiency.
To look for ribosome occupancy features beyond the CDS of transcripts, we first took advantage of the deep coverage and strand specificity of our RNA-seq data to identify 5′ leaders and 3′ UTRs of the *P. falciparum* transcriptome. We constructed a hidden Markov model (HMM) to automatically delineate the boundaries of both 5′ leaders and 3′ UTRs for known gene models (see ‘Materials and methods’). Within the limits imposed by our data, we were able to describe 5′ mRNA leaders and/or 3′ UTRs for 3569 genes in at least one of the stages (*Figure 6—figure supplement 1*, *Figure 2—source data 1*). 5′ leaders are on average longer than 3′ UTRs in each of the stages and median lengths across stages vary to a larger degree for 5′ leaders (from 607 to 1040 nt) than for 3′ UTRs (518–622 nt). The longest 5′ mRNA leader was measured in late trophozoites (8229 nt) for the Ap2 transcription factor, PF11_0404, and the longest 3′ UTR stretched 4773 nt for 60S ribosomal protein L7-3, PF14_0231, in rings. An example pair of genes with mapped 5′ leaders and 3′ UTRs is shown in *Figure 6*. Here, our HMM predicts a 636 nt and a 781 nt 5′ leader and a 468 nt and 423 nt 3′ UTR for the Myb2 transcription factor (PF10_0327) and the bromodomain protein (PF10_0328), respectively. These genes, encoded on opposite strands, share a 1536-nt intergenic sequence; however, the span between the region delimited by their 5′ leader sequence is only 120 nt and presumably harbors their respective promoters.

Next, using mRNA boundaries derived from our data, we analyzed ribosome distribution along each transcript during life cycle progression. More than 80% of the ribosome footprints in rings, early trophozoites, late trophozoites, and schizonts, were mapped to CDS regions of the genome, except in merozoites, where only 68% were mapped to the CDS (*Figure 7A*). On average less than 1% of all reads obtained were mapped to 3′ UTRs in each stage, and most transcripts had no observed footprints past the stop codon. In contrast, footprints were far more common in 5′ leaders (9.1%, 4.8%, 7.5%, and 4.8% in rings, early trophozoites, late trophozoites, and schizonts, respectively) particularly in merozoites (23%). Footprint enrichment is specific to 5′ leaders and not due to non-specific background since this would result in an increase of footprints mapping evenly along the length of the transcript, including the 3′ UTR, and not just the 5′ leader. Furthermore, these footprints most likely represent ribosomes because they derive from the 80S monosome fraction of the sucrose gradient, and their footprint read length distributions are equal to those of CDS mapping footprints, whereas they are significantly divergent from rRNA or tRNA read length distributions (*Figure 7—figure supplement 1*).

---

**Figure 6.** Example of extended transcript annotations using the HMM. 5′ leaders and 3′ UTRs of the gene pair Myb2 (PF10_0327) and bromodomain protein (PF10_0328) were defined using the HMM designed (see ‘Materials and methods’). The sizes of 5′ leaders and 3′ UTRs of these genes in the schizont stage are indicated. The intergenic region is 1536 nt and the spanning distance separating the 5′ leaders is 120 nt. Mappability = mappability score at that position; range 0 (white) to 30 (black). rM = coverage (reads per million reads mapped).

DOI: 10.7554/eLife.04106.016

The following figure supplement is available for figure 6:

**Figure supplement 1.** HMM-defined 5′ leader and 3′ UTR characteristics.

DOI: 10.7554/eLife.04106.017
Figure 7. Transcripts accumulate ribosome density within the 5' leader. (A) Proportion of mRNA or ribosome footprint reads mapping to CDS, to HMM-defined 5' leaders and 3' UTRs, antisense to annotated coding genes or to other regions of the genome such as mitochondria, plastid, tRNA, rRNA, ncRNA, and 5' leader and 3' UTR regions not defined by the HMM. (B) Proportion of ribosome footprints mapping inside or outside predicted uORFs in the HMM-defined 5' leaders. (C) Ribosome footprint (green) and mRNA (blue) profiles of the EBA-175 (MAL7P1.176) gene in rings (R) showing ribosome footprint accumulation on the 5' leader. In the detail, the bars above the gene model indicate AUG, stop, and any other codon, in green, red, and gray, respectively and in all three possible frames. Gray bars indicate the 9 uORFs present in the 5' leader, starting with an AUG (green line) and ending with a stop codon (red line). Black bar inside CDS indicates a deletion specific to the W2 strain used in this study. CDS, white boxes; HMM-defined UTRs, black lines. Mappability = mappability score at that position; range 0 (white) to 30 (black). rM = coverage (reads per million reads mapped).

DOI: 10.7554/eLife.04106.018

The following source data and figure supplements are available for figure 7:

Source data 1. Predicted uORFs.
DOI: 10.7554/eLife.04106.019

Figure supplement 1. 5' leader footprints are derived from ribosomes.
DOI: 10.7554/eLife.04106.020

Figure supplement 2. Distribution of uORF coverage on 5' leaders of genes expressed during the IDC.
DOI: 10.7554/eLife.04106.021

Figure 7. Continued on next page
Upstream open reading frames are a major source for 5’ leader ribosome density found from yeast to humans (Ingolia et al., 2009; Brar et al., 2012), and these have been shown to play a role in translational regulation of the downstream ORF in a few well-studied examples (Morris and Geballe, 2000). In P. falciparum, ribosomes have been suggested to accumulate on 5’ leaders of genes displaying a delay in translation presumably due to long uORFs (Bunnik et al., 2013).

We defined 36,086 possible uORF regions in the 5’ leaders of genes expressed during the P. falciparum IDC using a liberal definition that includes any stretch of at least two amino acids, starting with an AUG codon (Figure 7—source data 1). Regardless of stage, half of the total ribosome footprint coverage in 5’ leaders, in aggregate, or on a gene-by-gene basis did not overlap with these predicted uORFs (Figure 7B, Figure 7C—figure supplement 2). We could find no significant correlation between the number of uORFs per gene, the uORF lengths, or the degree to which ribosome density was enriched in uORFs with translational efficiency (Figure 7—figure supplement 3). For example, erythrocyte binding antigen-175 (EBA-175, MAL7P1.176) is well translated in rings (log₂TE = 1.4) and displays a large amount of 5’ leader ribosome occupancy. Half (49%) of the reads map within the nine predicted uORFs on the 5’ leader of this gene, the other half maps outside these uORFs (Figure 7C). Using this liberal definition of an uORF, the data do not support an association between ribosome occupancy in these regions, nor does it support an association between the presence of these regions and translational efficiency.

Nonetheless, there exist at least two clear exceptions. First, we were able to validate translation of the reported uORF present in the 5’ leader sequence of the var2csa mRNA which is expressed only in rings (Amulic et al., 2009). The majority of ribosome footprint density localizes to this uORF, and to a second one just upstream, while the var2csa coding sequence is largely devoid of footprints (log₂TE_rings = −4.2, Figure 7—figure supplement 4), consistent with its translational repression during growth in the absence of placental tissue. Second, another striking example of uORF translation was found on the 5’ leader of a few well-studied examples (Ingolia et al., 2009). In fact 5’ leader ribosome density can be found on 5’ leaders devoid of AUGs. Overall, rings and merozoite stage parasites were found to express transcripts with the highest 5’RD (mean log₂5’RD = −0.03, and 0.11, respectively) relative to early trophozoites, late trophozoites, and schizonts (mean log₂5’RD = −1.11, −0.26, −0.83), where the range of 5’RD values is also narrower (Figure 8A). Interestingly, among genes at the extremes of the 5’RD distributions (mean ± 1 SD), we also found many of our identified translationally up- and down-regulated transcripts (66% and 40%, respectively).
Figure 8. 5′ ribosome density is commonly found on genes expressed during the IDC. (A) 5′ RD distributions in each stage. Transcripts in rings and merozoites have on average higher 5′ RD values; ± 2 SD values lie outside gray shade. μ = mean log$_2$5′RD, n = total number of genes. (B) 5′RD values of the translationally up-regulated set of Figure 8. Continued on next page
Figure 8. Continued

On average, 5′RD was enriched on translationally up-regulated transcripts (mean log2 5′RD = 0.83) and depleted for translationally down-regulated transcripts in all stages (mean log2 5′RD = −1.11), suggesting the possibility that 5′RD is a byproduct of translational efficiency itself (Figure 8B).

In order to determine whether a direct relationship between 5′RD and translational efficiency of the downstream ORF exists, we compared these values for each gene. 5′RD positively correlates, albeit moderately, with translational efficiency in all stages, particularly in rings and merozoites (r = 0.51 and 0.49, respectively). We focused on the subset of genes with highest and lowest 5′RD values (mean ± 2 SD) and found that only a fraction of the translationally up- and down-regulated genes overlap with this category of extreme 5′RDs in each stage (Figure 8C). The largest overlap occurred in rings where the highest 5′RD values were found in 43% (31 genes) of the translationally up-regulated genes, including MSP6, AQP, and SERA5. These results indicate that while in general a correspondence between 5′RD and translational efficiency exists, one is not necessarily predictive of the other and exceptions apply. This is the case, for example, of the translationally down-regulated transcript of the pseudogene PfRh3, which in rings has the second highest 5′RD value (log2 5′RD = 5.1).

In summary, our data establish ribosome accumulation on 5′ leaders as a common feature of transcripts expressed during the IDC. Ribosome density is not restricted to predicted uORFs present within these regions and, with few exceptions, the uORF number, length, or coverage level, is not a requirement for 5′ ribosome density and has no measurable effects on the translation of the downstream ORF. Even though 5′RD is more commonly found on 5′ leaders of highly translated transcripts, this is not a universal trend since only a moderate correlation exists between 5′RD and the translational efficiency of the downstream ORF.

3′ UTR ribosome occupancy is rare

While our data showed 3′ UTRs to be relatively depleted of ribosomes, we searched for rare cases of high 3′ UTR ribosome density, possibly arising from stop codon read-through, alternative stop codon usage, or re-initiation of downstream ORFs (Dunn et al., 2013; Guydosh and Green, 2014). We systematically searched for transcripts for which coverage, in a sliding window of 30 nt, was greater in the 3′ UTR than the CDS, and found 19 genes meeting this criterion. These genes could be qualitatively divided into two categories: 14 with putative stop codon read-through and/or alternate stop codon usage and 5 genes for which the origin of the 3′ UTR density is unclear (listed in Figure 9—source data 1). An example of stop codon read-through is the conserved plasmodium protein (PF13_0160), shown in Figure 9A. Ribosomes not only extend beyond the annotated stop codon of this transcript but also skip subsequent in-frame stop codons present on the predicted 644 nt 3′ UTR. Interestingly, ribosome footprints accumulate in a single large peak approximately 130 nt downstream of the annotated stop codon. On the 1290 nt 3′ UTR of the sodium-dependent phosphate transporter (MAL13P1.206), two large peaks of ribosome footprint density, one approximately 560 nt and the other 860 nt from the stop codon, can be observed (Figure 9B). The origin of these footprints is unclear, and it is possible that these are the product of nuclease protection by RNA-binding proteins that co-sediment with the 80S monosome. To confirm that 3′ UTR mapping reads are derived from ribosome footprints, we compared their cumulative read length distributions against a typical CDS footprint read length distribution (Figure 9—figure supplement 1). For the 16 of the 19 genes we observed no significant difference in footprint size distributions localized to the CDS compared with the 3′ UTR. For the remaining three genes, the sodium-dependent phosphate transporter (MAL13P1.206), the acyl-Coa synthetase (PFD0085c), and the conserved plasmodium protein (PF13_0160), 3′ UTR footprint size distributions were divergent from those on the CDS, implying that footprints found on these genes’ 3′ UTRs may be produced by nuclease protection of these regions by factors other than ribosomes that co-sediment with 80S ribosomes.
Antisense detection plays an important role in gene regulation from bacteria to humans, and while its role is increasingly studied in these organisms (Faghihi and Wahlestedt, 2009), less is known about its relevance in *P. falciparum*. Previous serial analysis of gene expression (SAGE) (Patankar et al., 2001), nuclear run-on experiments (Militello et al., 2005), and more recently antisense splicing events detected by RNA-seq (López-Barragán et al., 2011; Sorber et al., 2011), suggest that antisense RNAs are synthesized by RNA pol II and may constitute up to ~12% of the erythrocytic-stage steady-state RNA (Gunasekera et al., 2004), yet their presence and biological role, if any, remains unclear. A more recent study found no correlation between natural antisense transcript levels and protein abundance (Siegel et al., 2014).

The 30 nt fragmentation and RNA-ligase-based library preparation method employed here affords exquisite strand specificity by minimizing artifacts associated with random priming during reverse transcription. As evidence of this specificity, the highest expressed gene in our data set, histone h2a (PFF0860c), yielded a total of 765,510 reads on the sense strand, and only two reads on

![Figure 9](image-url)

**Figure 9.** 3' UTR ribosome density. (A) Late trophozoite stage mRNA (blue) and ribosome footprint (green) profiles of the conserved plasmodium protein, PF13_0160. Ribosomes can be detected up to ~130 nt beyond the stop codon on the 3' UTR and accumulate in a single large peak. Red lines indicate in-frame stop codons on the 3' UTR. (B) Two large peaks of ribosome footprint density can be detected 560 nt and 860 nt downstream from the stop codon in the 3' UTR of the sodium-dependent phosphate transporter, MAL13P1.206.

DOI: 10.7554/eLife.04106.027

The following source data and figure supplements are available for figure 9:

- **Source data 1.** Genes with 3' UTR ribosome occupancy.
  DOI: 10.7554/eLife.04106.028

- **Figure supplement 1.** 3' UTR ribosome footprint size distribution.
  DOI: 10.7554/eLife.04106.029
the antisense strand, corresponding to a sense:antisense ratio greater than $10^6$. Furthermore, our HMM mapping of 5′ leaders and 3′ UTRs facilitates the differentiation between independently transcribed antisense RNA and transcripts that occur by virtue of being part of an adjacent gene. We took advantage of the nature of our data set to identify antisense transcripts and looked for effects on sense mRNA translation.

For this analysis only, we relaxed our stringent coverage threshold from $\geq 32$ rM to $\geq 16$ rM for inclusion of antisense transcripts. We based our threshold on the presence of an antisense transcript to the sexual stage specific gene pf16 (PFD0310w) confirmed by strand-specific RT-PCR (Figure 10—figure supplement 1). This antisense is predicted by the HMM to be $\sim 4$ kb, extending over the complete coding sequence and beyond and with a coverage level of 23 rM over the sense CDS. Using the 16 rM threshold, we detected 84 antisense transcripts to several known ORFs (listed in Figure 10—source data 1), including the nucleoside transporter pfNT4 (PFA0160c) depicted in Figure 10A. The merozoite stage contained the highest number of antisense transcripts (46), and the fewest (13) were found in early trophozoites. Manual inspection revealed that in 63% of these instances, the putative antisense transcript actually emanates from the 5′ leader or 3′ UTR of a neighboring gene (not defined by the HMM). Antisense reads for the para-hydroxybenzoate polypropyltransferase (PFF0370w), for example, are actually derived from the 3′ UTR of the neighboring conserved protein PFF0375c (Figure 10B).

We next interrogated the impact of this set of antisense transcripts. Overall, antisense transcripts showed no effect on mRNA abundance and translational efficiencies of the cognate sense transcript. These observations parallel those described for antisense transcripts in yeast (Brar et al., 2012). Thus, at first approximation antisense transcripts do not appear to play a role in translational regulation. However, these observations could be confounded due to the small number of genes in this set, and we cannot exclude the possibility of sense/anti-sense heterogeneity at the single cell level, obscured here at the population level.

Discussion

Herein we present, for the first time, a comprehensive view of the coupled transcriptional and translational dynamics of the *P. falciparum* IDC by determination of transcript abundance and architecture together with ribosomal density and positioning. The quality of our data relies on several critical features: (1) high temporal specificity and reproducibility of fully independent biological replicas of five strictly staged cultures; (2) purified merozoites to allow discrete measurements in this stage without confounding contributions from schizonts or rings; (3) monosome isolation from sucrose gradients to specifically enrich for ribosome-derived footprints and avoid potential complications that can arise with methods like sucrose cushions which are prone to mRNA contamination; (4) sufficient sequencing depth of biological replicates to set a statistical threshold for minimum read coverage and to demonstrate reproducibility; (5) stringent strand specificity to facilitate an HMM for the description of transcript boundaries and the detection of antisense transcription.

Previous studies of the transcript abundance in the malaria blood stages revealed a periodic cascade of gene expression, whereby the majority of expressed genes exhibit one peak of expression per cell cycle (Bozdech et al., 2003). The global profile of transcriptional expression was subsequently found to be highly stereotypical across strains and appeared to lack dynamic responses to perturbation (Llinás et al., 2006; Ganesan et al., 2008). It has been suggested that translational control of protein expression could compensate for the lack of transcriptional dynamics. Proteomic studies described delays in peak mRNA and corresponding protein abundance implicating translational or post-translational mechanisms in the modulation of gene expression (Le Roch et al., 2004; Foth et al., 2011).

Our ribosome profiling results reveal a tight coupling of transcription and translation for the majority of expressed genes, indicating that most protein products are translated with highly similar timing and in proportion to their corresponding mRNA transcripts. Synthesized proteins are likely to exert their functions immediately upon translation but post-translational regulation, not captured by our data, could still be at play. Direct correlations of translational efficiencies measured in this study along with proteomic data sets are hampered by the reduced sensitivity of the latter and differences in temporal resolution and staging of the parasites between data sets. However, the available proteomic evidence is largely consistent with the results presented here, particularly for highly translated proteins. The simultaneous capture of mRNA abundance and
Figure 10. Strand-specific libraries can distinguish antisense from sense gene transcription. (A) Schizont stage mRNA (blue) and ribosome footprint (green) profiles of the nucleoside transporter pfNT4 (PFA0160c). The antisense transcript covers the full extent of the sense transcript and displays ribosome density. (B) An example of antisense reads originating from a neighboring UTR in the schizont stage. The antisense reads in the para-hydroxybenzoate polyprenyltransferase (PFF0370w) stem from the 3‘ UTR of the neighboring conserved plasmodium protein (PFF0375c).

DOI: 10.7554/eLife.04106.030

The following source data and figure supplements are available for figure 10:

Source data 1. Antisense transcripts.
DOI: 10.7554/eLife.04106.031

Figure supplement 1. Strand-specific RT-PCR detection of the antisense to Pf16.
DOI: 10.7554/eLife.04106.032
translation is expected to be a more accurate proxy for protein levels than measurements of mRNA abundance alone (Ingolia et al., 2009) and provides a critical resource for the identification of instances of post-translational regulation of gene expression. However, we note that this data set only provides a direct measure of relative changes in translational efficiencies rather than changes in bulk transcription and translation.

While no up- or down-regulation of global translation efficiencies were observed in any particular stage, more extreme translational efficiencies were measured in subsets of genes expressed in rings and merozoites. We find 177 translationally up-regulated genes with functions predominantly related to merozoite egress and invasion, with peak mRNA in schizonts and peak translational efficiency in rings. It is likely that the genes with unknown functions, regulated in an analogous way during the merozoite to ring transition, are also associated with this process. Our data support a model whereby the transcripts of proteins necessary for merozoite structure and function are made in the previous stage in large abundance, are translationally up-regulated during the invasion process, and remain highly translated well into the ring stage despite rapid mRNA decay during this stage (Shock et al., 2007). Whether the accumulation of 5′ leader ribosome density is a mechanism that assists in this process or is it merely a byproduct of more efficient ribosomal initiation on these templates remains to be tested. With the emergence of genome editing tools such as CRISPR/Cas9 (Ghorbal et al., 2014), it may be possible to create versions of genes with altered cis-acting sequences to test for modulation of 5′ ribosome density and its effect on translational efficiency.

The global nature of ribosome accumulation within the 5′ leader sequences of many transcripts during the IDC and the lack of an association between 5′RD and the number or length of uORFs suggests that ribosomes accumulate on 5′ leaders through means other than a uORF model. For comparison, in yeast under starvation conditions the fraction of ribosome footprints derived from 5′ leaders is increased by sixfold and in some cases no single uORF can account for the entire distribution of ribosomes on the 5′ leader of a gene (Ingolia et al., 2009), much like P. falciparum. What mechanism could account for global ribosome accumulation in the 5′ leader? The presence of apparent 80S ribosomes within the 5′ leader sequence, regardless of whether they cover uORFs or not, suggests an engagement mode in which the fidelity of start codon recognition is altered or suspended. Current models propose that the 43S pre-initiation complex loads onto the mRNA with the assistance of other initiation factors near the 5′ cap and proceeds to scan down the length of the mRNA until it encounters an AUG codon. This is followed by the assembly of the 48S preinitiation complex and then finally the 80S complex (for review, see Hinnebusch, 2011). The AUG that is ultimately chosen is not always the first one encountered, and its sequence context is important for recognition. The factors eIF1, eIF1A, and eIF5 have been implicated in recognition of the ‘correct’ AUG (Aitken and Lorsch, 2012). In the case of P. falciparum, differential regulation or modification of these factors could plausibly result in altered start codon selection and 80S assembly. Whether prematurely initiated complexes are able to scan without synthesizing a peptide or are required to assemble and reassemble until encountering the right start codon remains an open question. Large 5′ ribosome accumulation on translationally up-regulated genes in the ring stage suggests that premature initiation on these transcripts is not detrimental. The development of an in vitro translation system that recapitulates upstream 80S assembly on P. falciparum 5′ leaders will allow direct testing of premature initiation and its effect on translational efficiency in this parasite.

Our ribosome profiling data add an important component to the rich compendium of genome-wide data, including transcript abundance (Bozdech et al., 2003), mRNA decay (Shock et al., 2007), splicing (Sorber et al., 2011), and proteomics for this parasite (Le Roch et al., 2004; Foth et al., 2011). Features such as 5′ leaders, 3′ UTRs, introns, and antisense transcripts are clearly visible and often well delineated. While experimental validation of transcriptional start sites, terminators, and promoters is required, spanning regions between transcripts, such as the one shown in Figure 6, can be used for the search and identification of such functional sites in a reduced sequence space. The data are available at NCBI GEO (accession #GSE58402) to facilitate future queries and normalized read coverage plots for all 5 timepoints are available packed as a single Mochiview file (Caro et al., 2014). Together our results describe a simplified regulatory architecture of gene translation, albeit one that includes peculiar and potentially unique mechanisms specialized for its highly structured and coordinated lifecycle within erythrocytes. Further biochemical dissection of translational initiation mechanisms and determinants of translational efficiency unique to Plasmodium may reveal weaknesses that could be exploited for possible therapeutic intervention.
Materials and methods

Cell culture
W2 strain cultures were maintained in Hyperflasks (Corning, Corning, NY) in 500 ml RPMIc (RPMI 1640 media supplemented with 0.25% Albumax II (GIBCO, Grand Island, NY), 2 g/L sodium bicarbonate, 0.1 mM hypoxanthine, 25 mM HEPES (pH 7.4), and 50 µg/L gentamycin), at 37°C, 5% O₂, and 5% CO₂, to maximum 10–15% parasitemia at 5% hematocrit (HC) and frequent media changes (at least every 6–8 hr). Cells were synchronized by two consecutive sorbitol treatments for three generations, for a total of six treatments. Maximum invasion, point at which half of the culture is either rings or schizonts, was defined as hour zero and independent time points containing ~10^10 parasites were harvested 11, 21, 31, and 45 hr later.

Polysome isolation and library generation
Cultures were incubated for 5 min in 500 ml 37°C RPMIc, 100 µg/ml cycloheximide (Acros Organics, Bridgewater, NJ) and harvested by centrifugation for 8 min at 3.65×g at room temperature. An aliquot was removed and flash frozen in liquid nitrogen for total RNA purification, followed by poly(A)-purification and chemical fragmentation with Zn^{2+} to ~30 nt for consistency in mRNA-Seq library preparation. The remaining culture was treated with ice-cold 0.1% saponin in 1X PBS, 100 µg/ml cycloheximide, for RBC lysis. Parasites were resuspended in ice-cold parasite lysis buffer (15 mM KOAc, 15 mM MgOAc, 10 mM Tris HCl pH 7.4, 0.5 mM DTT, 0.5% Triton X-100, 100 µg/ml cycloheximide) and dripped into a conical tube filled with, and immersed in, liquid nitrogen. Frozen cells transferred placed in liquid nitrogen pre-chilled chambers and pulverized for 3 min at 15 Hz, on a Retsch MM301 mixer mill. Pulverized cells were thawed on ice, and cell debris was removed by centrifugation at 4°C, 16,000×g for 10 min. The supernatant was treated with 2.88 U/µg Micrococcal nuclease for 30 min at room temperature and immediately loaded onto sucrose gradients for ultracentrifugation at 35,000 rpm for 3 hr at 4°C in a L8-60 M Beckman centrifuge. Monosome fractions only, were collected to generate ribosome footprint libraries for deep sequencing using the HiSeq 2000 (Illumina, San Diego, CA), as described (Ingolia et al., 2009).

Merozoite purification
Late stage schizonts (40–44 hpi) were magnetically purified using MACS LD columns (Miltenyi Biotec, San Diego, CA) and resuspended in RPMIc without blood addition. After reaching maximum invasion (1:1 schizont to ring ratio), cultures were harvested by centrifugation at 1500 rpm at room temperature for 5 min. Pelleted cultures were resuspended into fresh RPMIc and placed at 37°C. Merozoites in the supernatant were treated with 100 µg/ml cycloheximide for 5 min at room temperature, harvested at 4000 rpm at 4°C for 5 min and resuspended in RPMIc and passed again through a MACS LD column. Parasite lysis buffer was added to the merozoite-enriched flow-through and flash frozen in liquid nitrogen. This procedure was repeated three times every 45 min using the original culture. The same procedure described above was used for RNA extraction and library preparation.

SNP-corrected W2 genome
W2 strain genomic DNA was isolated from >90% synchronized ring stage cultures. Paired end libraries were constructed using the Nextera DNA Sample Prep Kit (Epicenter Biotechnologies, Madison, WI) according to the manufacturer’s instructions reducing PCR cycles from nine to six and using 80% A/T dNTPs. Libraries were sequenced using the HiSeq 2000 (Illumina). Reads were aligned to the P. falciparum PlasmoDB 3D7 version 7.1 genome using Bowtie 0.12.1 (Langmead et al., 2009) with parameters –v1 –m 1 (one mismatch allowed, unique mapping). A SNP was called when five or more W2 reads supported, with over 90% agreement, a different base than the one found in the P. falciparum 3D7 7.1 genome. 19401 SNPs (0.08% of total bases) were detected and used to produce the SNP-corrected W2 genome based on the 3D7 genome. Fastq files are available for download at NCBI SRA, accession #SRP042946.

Software pipeline, mappability, and rpkM calculation
Quality-filtered ribosome footprints and mRNA sequencing reads were trimmed to remove library adapter sequence, filtered for P. falciparum rRNA using blast, and aligned uniquely to the W2 SNP-corrected genome using Bowtie 0.12.1 (Langmead et al., 2009) allowing no mismatches. The percentage mappability was calculated using an in silico library of the P. falciparum W2 SNP-corrected genome created using a single nucleotide sliding window of 30 nt. The in silico library was uniquely
aligned to the genome allowing no mismatches. The mappability score is given by the number of 30 nt sequences covering each nt position in the genome, such that any position has a score that ranges from 0 to 30. Both mRNA and ribosome footprint rpkmS were calculated as in Mortazavi et al. (2008), excluding the first 50 bases of each gene to eliminate bias introduced by the observed ribosome accumulation peak near the start codon. Genes with fewer than 80% mappable bases (248 genes) or any overlapping non-CDS feature on the same strand (77 genes) were excluded from this calculation. Data are available for download at NCBI GEO, accession #GSE58402. MochiView genome browser data tracks are available in Supplementary file 1 (Homann et al., 2010).

**Extended phaseogram**

The genes of the RNA-seq transcriptome obtained in this work were listed in the same phaseogram order as the previously published microarray transcriptome (Bozdech et al., 2003). The criteria for inclusion of a gene into the phaseogram was mRNA ≥ 32 rM, >2 peak to trough ratios, and Pearson correlation coefficient >0.8 with the expression profiles of the two neighboring genes.

**Hidden Markov model to describe transcript boundaries**

The HMM was built using RNA-Seq data obtained in this study and two states: transcript (t) or intergenic (i) with three possible emissions: (1) read present, (2) read not present but position is unmappable, (3) read not present but the position is mappable. Both state and emission probabilities were calculated using ~30 kb training set of manually identified transcript and non-transcript regions. The initial probabilities were set to 0.5. Transition probabilities were estimated from the median length of intergenic regions of (1252 nt) and median lengths of CDS regions (2545 nt), where the P<sub>t→i</sub> = (1/2545), P<sub>t→t</sub> = (2544/2545), P<sub>i→t</sub> = (1251/1252), and P<sub>i→i</sub> = (1/1252). We applied the Viterbi algorithm to predict the optimal path of transcript tracks per time point with a 10 nt window resolution. HMM-defined 5’ leader and 3’ UTR coordinates are available for download at NCBI GEO, accession #GSE58402.

**Strand-specific RT-PCR**

Total RNA from late stage parasites was isolated and reverse transcribed using SuperScript III (Invitrogen, Carlsbad, CA) according to manufacturer’s instructions, using either an antisense-specific primer to Pfs16 (PFD0310w) or a random nonamer. cDNA was amplified using the Pfs16 antisense-specific primer as a forward primer in combination with one of five reverse primers (Supplementary file 3). 18S rRNA primers were used in the control reactions with the random nonamer-derived cDNA.

**Acknowledgements**

The authors would like to thank Dr Graham Ruby for technical expertise in devising the HMM for UTR description and Dr. Giselle Knudsen for technical support. We would also like to thank members of the Weissman Lab: Dr Eugene Oh, and Dr Gloria Brar for helpful experimental advice and suggestions for the analysis of data and Dr Jonathan Weissman for the use of a density gradient fractionator and for helpful discussions during article preparation.

**Additional information**

**Funding**

| Funder                        | Grant reference number | Author                        |
|-------------------------------|------------------------|-------------------------------|
| Howard Hughes Medical Institute | U01 AI075517           | Florence Caro, Vida Ahyong, Miguel Betegon, Joseph L DeRisi |

The funders had no role in study design, data collection and interpretation, or the decision to submit the work for publication.

**Author contributions**

FC, VA, Conception and design, Acquisition of data, Analysis and interpretation of data, Drafting or revising the article; MB, Conception and design, Analysis and interpretation of data; JLDR, Conception and design, Analysis and interpretation of data, Drafting or revising the article.
Additional files

**Supplementary files**
- Supplementary file 1. MochiView genome browser data tracks.
  DOI: 10.7554/eLife.04123.033
- Supplementary file 2. RESA2 mapping reads.
  DOI: 10.7554/eLife.04123.034
- Supplementary file 3. Strand-specific RT-PCR primers.
  DOI: 10.7554/eLife.04123.035

**Major datasets**

The following datasets were generated:

| Author(s)               | Year | Dataset title                                                  | Dataset ID and/or URL | Database, license, and accessibility information |
|-------------------------|------|----------------------------------------------------------------|-----------------------|--------------------------------------------------|
| Caro F, Ahyong V, Betegon M, DeRisi JL | 2014 | Ribosome Profiling in P. falciparum asexual blood stages       | http://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc=GSE58402 | Publicly available at NCBI Gene Expression Omnibus |
| Caro F, Ahyong V, Betegon M, DeRisi JL | 2014 | W2 Plasmodium falciparum genome sequences                       | SRP042946             | Publicly available at NCBI SRA                   |
| Caro F, Ahyong V, Betegon M, DeRisi JL | 2014 | Data from: Genome-wide Regulatory Dynamics of Translation in the Plasmodium falciparum Asexual Blood Stages | http://dx.doi.org/10.5061/dryad.vb855 | Available at Dryad Digital Repository under a CC0 Public Domain Dedication |

References

Altken CE, Lorsch JR. 2012. A mechanistic overview of translation initiation in eukaryotes. *Nature Structural & Molecular Biology* 19:568–576. doi: 10.1038/nsmb.2303.

Amulic B, Salanti A, Lavstsen T, Nielsen MA, Deitsch KW. 2009. An upstream open reading frame controls translation of var2csa, a gene implicated in placental malaria. *PLOS Pathogens* 5:e1000256. doi: 10.1371/journal.ppat.1000256.

Aurrecoechea C, Brestelli J, Brunk BP, Dommer J, Fischer S, Gajria B, Gao X, Gingle A, Grant G, Harb OS, Heiges M, Innamorato F, Iodice J, Kissing JC, Kraemer E, Li W, Miller JA, Nayak V, Pennington C, Pinney DF, Roos DS, Ross C, Stoeckert CJ Jr, Treuman C, Wang H. 2009. PlasmoDB: a functional genomic database for malaria parasites. *Nucleic Acids Research* 37:D539–D543. doi: 10.1093/nar/gkn814.

Blackman MJ. 2008. Malarial proteases and host cell egress: an “emerging” cascade. *Cellular Microbiology* 10:1925–1934. doi: 10.1111/j.1462-5822.2008.01176.x.

Bozdech Z, Llinás M, Pulliam BL, Wong ED, Zhu J, DeRisi JL. 2003. The transcriptome of the intraerythrocytic developmental cycle of *Plasmodium falciparum*. *PLOS Biology* 1:ES. doi: 10.1371/journal.pbio.0000005.

Brar GA, Yassour M, Friedman N, Regev A, Ingolia NT, Weissman JS. 2012. High-resolution view of the yeast meiotic program revealed by ribosome profiling. *Science* 335:552–557. doi: 10.1126/science.1215110.

Bunnik EM, Chung DW, Hamilton M, Ponst N, Saraf A, Prudhomme J, Floresn L, Le Roch KG. 2013. Polysome profiling reveals translational control of gene expression in the human malaria parasite *Plasmodium falciparum*. *Genome Biology* 14:R128. doi: 10.1186/gb-2013-14-11-r128.

Cappai R, Kaslow DC, Peterson MG, Cowman AF, Anders RF, Kemp DJ. 1992. Cloning and analysis of the RESA-2 gene: a DNA homologue of the ring-infected erythrocyte surface antigen gene of *Plasmodium falciparum*. *Molecular and Biochemical Parasitology* 54:213–221. doi: 10.1016/0166-6851(92)90113-X.

Caro F, Ahyong V, Betegon M, DeRisi JL. 2014. Data from: genome-wide regulatory dynamics of translation in the plasmodium falciparum asexual blood stages. Dryad Digital Repository. doi: 10.5061/dryad.vb855.

Dunn JG, Foo CK, Belletier NG, Gavis ER, Weissman JS. 2013. Ribosome profiling reveals pervasive and regulated stop codon readthrough in *Drosophila melanogaster*. *eLife* 2:e01179. doi: 10.7554/eLife.01179.

Faghihi MA, Wahlestedt C. 2009. Regulatory roles of natural antisense transcripts. *Nature Reviews Molecular Cell Biology* 10:637–643. doi: 10.1038/nrm2738.

Farrow RE, Green J, Katsimitouza L, Taylor WR, Holder AA, Molloy JE. 2011. The mechanism of erythrocyte invasion by the malarial parasite, *Plasmodium falciparum*. *Seminars in Cell & Developmental Biology* 22:953–960. doi: 10.1016/j.semcdb.2011.09.022.

Florens L, Washburn MP, Raine JD, Anthony RM, Grainger M, Haynes JD, Moch JK, Muster N, Sacci JB, Tabb DL, Witney AA, Wolters D, Wu Y, Gardner MJ, Holder AA, Sinden RE, Yates JR, Carucci DJ. 2002. A proteomic view of the Plasmodium falciparum life cycle. *Nature* 419:520–526. doi: 10.1038/nature01107.

Foth BJ, Zhang N, Chaal BK, Sze SK, Freiser PR, Bozdech Z. 2011. Quantitative time-course profiling of parasite and host cell proteins in the human malaria parasite *Plasmodium falciparum*. *Molecular & Cellular Proteomics* 10:M110.006411. doi: 10.1074/mcp.M110.006411.
Genomics and evolutionary biology | Microbiology and infectious disease

Ganesan K, Ponnsee N, Jiang L, Fowlie JW, White J, Kamchonwongpaisan S, Yuthavong Y, Wilairat P, Rathod PK. 2008. A genetically hard-wired metabolic transcriptome in *Plasmodium falciparum* fails to mount protective responses to lethal antifolates. *PLOS Pathogens* **4**:e1000214. doi: 10.1371/journal.ppat.1000214.

Gardner MJ, Hall N, Fung E, White O, Berriman M, Hyman RW, Carlton JM, Pain A, Nelson KE, Bowman S, Paulsen IT, James K, Eisen JA, Rutherford K, Salzberg SL, Craig A, Kyes S, Chan MS, Nene V, Shollom SJ, Suh B, Peterson J, Angiuoli S, Pertea M, Allen J, Seidman JG, Haft D, Mathew MW, Vaidya AB, Martin DM, Fairlamb AH, Fraunholz MJ, Roos DS, Ralph SA, McFadden GI, Cummings LM, Subramanian GM, Mungall C, Venter JC, Carucci DJ, Hoffman SL, Newbold C, Davis RW, Fraser CM, Barrell B. 2002. Genome sequence of the human malaria parasite *Plasmodium falciparum*. *Nature* **419**:498–511. doi: 10.1038/nature01097.

Gborral M, Gorman M, Macpherson CR, Martins RM, Scherf A, Lopez-Rubio JJ. 2014. Genome editing in the human malaria parasite *Plasmodium falciparum* using the CRISPR-Cas9 system. *Nature Biotechnology* **32**:819–821. doi: 10.1038/nbt.2925.

Gunasekera AM, Patankar S, Schug J, Eisen G, Kissing J, Roos D, Wirth DF. 2004. Widespread distribution of antisense transcripts in the *Plasmodium falciparum* genome. *Molecular and Biochemical Parasitology* **136**:35–42. doi: 10.1016/j.molbiopara.2004.02.007.

Guydosh NR, Green R. 2014. Dom34 rescues ribosomes in 3’ untranslated regions. *Cell* **156**:950–962. doi: 10.1016/j.cell.2014.02.006.

Hinnebusch AG. 2011. Molecular mechanism of scanning and start codon selection in eukaryotes. *Microbiology and Molecular Biology Reviews* **75**:434–467. doi: 10.1128/MMBR.00008-11.

Homann OR, Johnson AD. 2010. MochiView: versatile software for genome browsing and DNA motif analysis. *BMC Biology* **8**:49. doi: 10.1186/1741-7007-8-49.

Hu G, Cabrera A, Kono M, Mok S, Chaal BK, Haase S, Engelberg K, Chee S, S, Spielmann T, Preiser PR, Gilberger TW, Bozdech Z. 2010. Transcriptional profiling of growth perturbations of the human malaria parasite *Plasmodium falciparum*. *Nature Biotechnology* **28**:91–98. doi: 10.1038/nbt.1597.

Hughes KR, Philip N, Starnes GL, Taylor S, Waters AP. 2010. From cradle to grave: RNA biology in malaria parasites. *Wiley Interdisciplinary Reviews: RNA* **1**:287–303. doi: 10.1002/wrna.30.

Ingolia NT. 2014. Ribosome profiling: new views of translation, from single codons to genome scale. *Nature Reviews Genetics* **15**:205–213. doi: 10.1038/nrg3645.

Ingolia NT, Ghaemmaghami S, Newman JR, Weissman JS. 2009. Genome-wide analysis in vivo of translation with nucleotide resolution using ribosome profiling. *Science* **324**:218–223. doi: 10.1126/science.1168978.

Langmead B, Trapnell C, Pop M, Salzberg SL. 2009. Ultrafast and memory-efficient alignment of short DNA sequences to the human genome. *Genome Biology* **10**:R25. doi: 10.1186/gb-2009-10-3-r25.

Lasonder E, Ishihama Y, Andersen JS, Vermunt AM, Pain A, Sauerwein RW, Eling WM, Hall N, Waters AP, Stunnenberg HG, Mann M. 2002. Analysis of the *Plasmodium falciparum* proteome by high-accuracy mass spectrometry. *Nature* **419**:537–542. doi: 10.1038/nature01111.

Le Roch KG, Johnson RJ, Floresn L, Zhou Y, Santrosyan A, Grainger M, Yan SF, Williamson KC, Holder AA, Carucci DJ, Yates JR III, Winzeler EA. 2004. Global analysis of transcript and protein levels across the *Plasmodium falciparum* life cycle. *Genome Research* **14**:2308–2318. doi: 10.1101/gr.2523904.

Llinàs M, Bozdech Z, Wong ED, Adai AT, DeRisi JL. 2006. Comparative whole genome transcriptome analysis of three *Plasmodium falciparum* strains. *Nucleic Acids Research* **34**:1166–1173. doi: 10.1093/nar/gkj517.

López-Barragán MJ, Lemieux J, Quiñones M, Williamson KC, Molina-Cruz A, Cui K, Barillas-Mury C, Zhao K, Su XZ. 2011. Directional gene expression and antisense transcripts in sexual and asexual stages of *Plasmodium falciparum*. *BMC Genomics* **12**:587. doi: 10.1186/1471-2164-12-587.

Mair GR, Braks JA, Garver LS, Wiegant JC, Hall N, Dirks RW, Khan SM, Dimopoulos G, Janse CJ, Waters AP. 2006. Regulation of sexual development of Plasmodium by translational repression. *Science* **313**:667–669. doi: 10.1126/science.1125129.

Mair GR, Lasonder E, Garver LS, Franke-Fayard BM, Carret CK, Wiegant JC, Dirks RW, Dimopoulos G, Janse CJ, Waters AP. 2010. Universal features of post-transcriptional gene regulation are critical for Plasmodium zygote development. *PLOS Pathogens* **6**:e1000767. doi: 10.1126/science.1125129.

Millitello KT, Patel V, Chessler AD, Fisher JK, Kasper JM, Gunasekera A, Wirth DF. 2005. RNA polymerase II synthesizes antisense RNA in *Plasmodium falciparum*. *RNA* **11**:365–370. doi: 10.1261/rna.7940705.

Morris DR, Geballe AP. 2000. Upstream open reading frames as regulators of mRNA translation. *Molecular and Cellular Biology* **20**:8635–8642. doi: 10.1128/MCB.20.23.8635-8642.2000.

Mortazavi A, Williams BA, McCue K, Schafer L, Wold B. 2008. Mapping and quantifying mammalian transcriptomes by RNA-Seq. *Nature Methods* **5**:621–628. doi: 10.1038/nmeth.1226.

Nirmanal N, Sims PFG, Hyde JE. 2004a. Translational up-regulation of antifolate drug targets in the human malaria parasite *Plasmodium falciparum* upon challenge with inhibitors. *Molecular and Biochemical Parasitology* **136**:63–70. doi: 10.1016/j.molbiopara.2004.02.013.

Nirmanal N, Sims PFG, Hyde JE. 2004b. Quantitative proteomics of the human malaria parasite *Plasmodium falciparum* and its application to studies of development and inhibition. *Molecular Microbiology* **52**:1187–1199. doi: 10.1111/j.1365-2958.2004.04049.x.

Oehring SC, Woodcroft BJ, Moes S, Wetzel J, Dietz O, Pulfer A, Dekiwadia C, Maeser P, Flueck C, Witter M, Brancucci NM, Niederwieser I, Jenoe P, Ralph SA, Voss TS. 2012. Organellar proteomics reveals hundreds of novel nuclear proteins in the malaria parasite *Plasmodium falciparum*. *Genome Biology* **13**:R108. doi: 10.1186/gb-2012-13-11-r108.

Patankar S, Munasinghe A, Shoaii A, Cummings LM, Wirth DF. 2001. Serial analysis of gene expression in *Plasmodium falciparum* reveals the global expression profile of erythrocytic stages and the presence of
anti-sense transcripts in the malarial parasite. Molecular Biology of the Cell 12:3114–3125. doi: 10.1091/mbc.12.10.3114.

Pease BN, Huttlin EL, Jedrychowski MP, Talevich E, Harmon J, Dillman T, Kannan N, Doerig C, Chakrabarti R, Gygi SP, Chakrabarti D. 2013. Global analysis of protein expression and phosphorylation of three stages of Plasmodium falciparum intraerythrocytic development. Journal of Proteome Research 12:4028–4045. doi: 10.1021/pr400394g.

Shock JL, Fischer KF, DeRisi JL. 2007. Whole-genome analysis of mRNA decay in Plasmodium falciparum reveals a global lengthening of mRNA half-life during the intra-erythrocytic development cycle. Genome Biology 8:R134. doi: 10.1186/gb-2007-8-7-r134.

Siegel TN, Hon CC, Zhang Q, Lopez-Rubio JJ, Scheidig-Benatar C, Martins RM, Sismeiro O, Coppée JY, Scherf A. 2014. Strand-specific RNA-Seq reveals widespread and developmentally regulated transcription of natural antisense transcripts in Plasmodium falciparum. BMC Genomics 15:150. doi: 10.1186/1471-2164-15-150.

Sorber K, Dimon MT, DeRisi JL. 2011. RNA-seq analysis of splicing in Plasmodium falciparum uncovers new splice junctions, alternative splicing and splicing of antisense transcripts. Nucleic Acids Research 39:3820–3835. doi: 10.1093/nar/gkq1223.

Tamez PA, Bhattacharjee S, van Ooij C, Hiller NL, Llinás M, Balu B, Adams JH, Haldar K. 2008. An erythrocyte vesicle protein exported by the malaria parasite promotes tubovesicular lipid import from the host cell surface. PLOS Pathogens 4:e1000118. doi: 10.1371/journal.ppat.1000118.

Vazeux G, Scanf CL, Fandeur T. 1993. The RESA-2 gene of Plasmodium falciparum is transcribed in several independent isolates. Infection and Immunity 61:4469–4472.

Yeoh S, O’Donnell RA, Koussis K, Dluzewski AR, Ansell KH, Osborne SA, Hackett F, Withers-Martinez C, Mitchell GH, Bannister LH, Bryans JS, Kettleborough CA, Blackman MJ. 2007. Subcellular Discharge of a serine Protease mediates Release of Invasive malaria parasites from host erythrocytes. Cell 131:1072–1083. doi: 10.1016/j.cell.2007.10.049.

Zhang M, Fennell C, Ranford-Cartwright L, Sakthivel R, Gueirard P, Meister S, Caspi A, Doerig C, Nussenzweig RS, Tuteja R, Sullivan WJ, Roos DS, Fontoura BM, Ménard R, Winzeler EA, Nussenzweig V. 2010. The plasmodium eukaryotic initiation Factor-2α kinase IK2 controls the latency of sporozoites in the mosquito salivary glands. The Journal of Experimental Medicine 207:1465–1474. doi: 10.1084/jem.20091975.

Zhang M, Mishra S, Sakthivel R, Rojas M, Ranjan R, Sullivan WJ, Fontoura BM, Ménard R, Dever TE, Nussenzweig V. 2012. PK4, a eukaryotic initiation factor 2α(eIF2α) kinase, is essential for the development of the erythrocytic cycle of plasmodium. Proceedings of the National Academy of Sciences of USA 109:3956–3961. doi: 10.1073/pnas.1121567109.

Zhang K, Rathod PK. 2002. Divergent regulation of dihydrofolate reductase between malaria parasite and human host. Science 296:545–547. doi: 10.1126/science.1068274.