A Comprehensive Survey of the Plasmodium Life Cycle by Genomic, Transcriptomic, and Proteomic Analyses

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Plasmodium berghei and Plasmodium chabaudi are widely used model malaria species. Comparison of their genomes, integrated with proteomic and microarray data, with the genomes of Plasmodium falciparum and Plasmodium yoelii revealed a conserved core of 4500 Plasmodium genes in the central regions of the 14 chromosomes and highlighted genes evolving rapidly because of stage-specific selective pressures. Four strategies for gene expression are apparent during the parasites’ life cycle: (i) housekeeping; (ii) host-related; (iii) strategy-specific related to invasion, asexual replication, and sexual development; and (iv) stage-specific. We observed posttranscriptional gene silencing through translational repression of messenger RNA during sexual development, and a 47-base 3’ untranslated region motif is implicated in this process.

Rodent malaria parasite species provide model systems that allow issues to be addressed that are impossible with the human-infectious species Plasmodium falciparum and P. vivax (1). Three closely related species, P. chabaudi, P. yoelii, and P. berghei, are in common use in the laboratory. Comparative sequencing and analysis of the genomes of such model species, in addition to the complete genome sequence of P. falciparum (2), provide insights into the evolution of Plasmodium genes and gene families (3).

The malaria parasite differentiates into a series of morphologically distinct forms in the vertebrate and mosquito hosts. It alternates between morphologically related invasive stages (sporozoite, merozoite, and ookinete) and replicative stages (pre-erythrocytic, erythrocytic-schizont, and oocyst) interposed by a single phase of sexual development that mediates transmission from the human host to the anopheline vector (1). This report integrates genome sequence analyses of P. berghei and P. chabaudi with transcriptome and proteome data for P. berghei, allowing the categorization of protein expression, the analysis of regulation mechanisms for gene expression, and the identification of species-specific gene families and genes under selective pressure.

**Genome sequencing and annotation.** Partial shotgun sequencing (4) of the genomes of P. c. chabaudi (AS clone) and P. berghei (ANKA clone) generated assemblies of ~17 and ~18 Mb, respectively (Table 1 and table S1). Orthologous genes of these two genomes and of P. y. yoelii (3) and P. falciparum (2) were inferred through bidirectional BLAST searches (Table 2). Combining the gene predictions of the three rodent parasites revealed that 4391 genes had orthologs in P. falciparum. These orthologs represent a universal Plasmodium gene set (table S2), which was mainly distributed across the central “core” regions of the 14 P. falciparum chromosomes. For example, in the core region of P. falciparum chromosome 2, 144 of 158 genes had rodent parasite orthologs (Fig. 1), whereas in the subtelomeric regions, only 3 of 65 genes showed (low) homology to rodent parasite genes (figs. S1 to S14). In addition to BLAST analysis, we manually examined the orthology of gene models on the basis of the conservation of gene order between the rodent parasites and P. falciparum, resulting in the identification of an additional 109 orthologs (table S3). There were no orthologs in the rodent parasite genomes for 736 P. falciparum genes, and 161 of these were located in the core regions (table S3). The other 575 are located in the subtelomeric regions, and Markov clustering (5) of these P. falciparum–specific genes revealed that almost half could be assembled into 12 distinct gene families (Fig. 1 and table S4). Only five subtelomeric gene families are obviously shared between all the sequenced Plasmodium species (table S4) (6). Previous studies have shown that a subtelomeric gene family of P. vivax, the P. vivax interspersed repeats (vir) (7), has related gene families in P. berghei (bir), P. chabaudi (cir), and P. yoelii (vir) (8, 9), and we suggest pir (Plasmodium interspersed repeats) to collectively describe the families. The bir and cir families code for highly variable proteins that share ~30% sequence identity at the amino acid level. The copy number appears to be much higher in P. y. yoelii (>800 copies) compared to P. berghei (180 copies) and P. c. chabaudi (138 copies).

**Selective pressure.** Comparison of orthologous genes of different species through

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models of nucleotide sequence evolution can be used to investigate variable (and positive or negative) selective pressures (10, 11). We determined the relative number of synonymous (dS) versus nonsynonymous (dN) substitutions between orthologs of \textit{P. berghei} and \textit{P. chabaudi}. In general, we found that orthologous gene pairs are under purifying selection pressure (and have dN/dS < 1) and that the observed ratios of median values for genes of rodent parasites (Table 2) were similar to those reported for genes of rodent parasites (Table 2) were similar to those reported for these species. Thus, the gene numbers indicated are for gene predictions where orthologs were identified in other species only. An excessive number of gene models were predicted for \textit{P. berghei} and \textit{P. c. chabaudi} because of the fragmented nature of the genome sequence data for these species. The high number of orthologs inferred between \textit{P. berghei} and \textit{P. c. chabaudi} compared to pairwise comparisons of the other species most likely reflects the method of automated annotation of both genomes, which used identical gene-finding algorithms (4). Median dN/dS value represents the median value of dN/dS for every gene pair and is not calculated from the median dN and dS values for each comparison. The median dN/dS for comparisons with \textit{P. falciparum} are low because of the saturation of synonymous changes in the alignments, resulting in high dS values.

**Table 1.** Genome summary statistics. A more detailed set of statistics is given in table S1.

<table>
<thead>
<tr>
<th>Statistic</th>
<th>\textit{P. berghei}</th>
<th>\textit{P. c. chabaudi}</th>
<th>\textit{P. y. yoellii}</th>
<th>\textit{P. falciparum}</th>
</tr>
</thead>
<tbody>
<tr>
<td>Size (bp)</td>
<td>17,996,878</td>
<td>16,866,661</td>
<td>23,125,449</td>
<td>22,853,764</td>
</tr>
<tr>
<td>No. contigs</td>
<td>7,497</td>
<td>10,679</td>
<td>5,687</td>
<td>93</td>
</tr>
<tr>
<td>Average contig size (bp)</td>
<td>2,400</td>
<td>1,580</td>
<td>4,066</td>
<td>213,586</td>
</tr>
<tr>
<td>Sequence coverage</td>
<td>4x</td>
<td>4x</td>
<td>5x</td>
<td>14.5x</td>
</tr>
<tr>
<td>No. protein coding genes</td>
<td>5,864*</td>
<td>5,698*</td>
<td>5,878</td>
<td>5,268</td>
</tr>
</tbody>
</table>

*An excessive number of gene models were predicted for \textit{P. berghei} and \textit{P. c. chabaudi} because of the fragmented nature of the genome sequence data for these species. Thus, the gene numbers indicated are for gene predictions where orthologs were identified in other \textit{Plasmodium} species only.

**Table 2.** Genome comparisons between the four sequenced \textit{Plasmodium} species. Av., average; \textit{P. berghei}; \textit{P. c. chabaudi}; \textit{P. f. falciparum}; \textit{P. y. yoellii}.

<table>
<thead>
<tr>
<th>Statistic</th>
<th>\textit{P. berghei} vs. \textit{P. y. yoellii}</th>
<th>\textit{P. berghei} vs. \textit{P. c. chabaudi}</th>
<th>\textit{P. c. chabaudi} vs. \textit{P. y. yoellii}</th>
<th>\textit{P. berghei} vs. \textit{P. f. falciparum}</th>
<th>\textit{P. c. chabaudi} vs. \textit{P. f. falciparum}</th>
</tr>
</thead>
<tbody>
<tr>
<td>Av. protein identity (%)</td>
<td>88.2</td>
<td>83.2</td>
<td>84.6</td>
<td>61.2</td>
<td>62.9</td>
</tr>
<tr>
<td>Av. nucleotide identity (%)</td>
<td>91.3</td>
<td>87.1</td>
<td>88.1</td>
<td>69.6</td>
<td>70.3</td>
</tr>
<tr>
<td>Median dN</td>
<td>0.005</td>
<td>0.007</td>
<td>0.006</td>
<td>0.29</td>
<td>0.26</td>
</tr>
<tr>
<td>Median dS</td>
<td>0.026</td>
<td>0.49</td>
<td>0.53</td>
<td>49.4</td>
<td>26.1</td>
</tr>
<tr>
<td>Median dN/dS</td>
<td>0.16</td>
<td>0.13</td>
<td>0.11</td>
<td>0.008</td>
<td>0.009</td>
</tr>
<tr>
<td>No. orthologous gene pairs</td>
<td>3153 (4641*)</td>
<td>3318</td>
<td>3375</td>
<td>3890</td>
<td>3842</td>
</tr>
</tbody>
</table>

*The high number of orthologs inferred between \textit{P. c. chabaudi} and \textit{P. berghei} compared to pairwise comparisons of the other species most likely reflects the method of automated annotation of both genomes, which used identical gene-finding algorithms (4). Median dN/dS value represents the median value of dN/dS for every gene pair and is not calculated from the median dN and dS values for each comparison. The median dN/dS for comparisons with \textit{P. falciparum} are low because of the saturation of synonymous changes in the alignments, resulting in high dS values.

**Fig. 1.** Schematic map of \textit{P. falciparum} chromosome 2. Arrow heads and boxes represent genes and their orientation on the DNA molecule. Thin and thick vertical lines represent 1-kb and 10-kb intervals, respectively. \textit{P. falciparum} genes with orthologs in the rodent malaria parasite genomes are marked in shades of blue according to their degree of similarity, from light blue (indicating 40% identity) through dark blue (indicating 100% identity); white genes show <40% identity to their closest ortholog. Weak orthologs not detected by reciprocal BLAST analyses are indicated in dark gray and in light gray if the gene is absent in all rodent malaria parasite genomes. \textit{P. falciparum} genes with no detectable ortholog are classified as follows: orange, \textit{var}, \textit{njf}, and \textit{stevor} gene families; yellow, centrally located expanded gene families shared with the rodent malaria parasite; red, all other \textit{P. falciparum} orphans. A full list of these genes and their classification can be found in table S3. Shaded areas of the map indicate the boundaries of the conserved chromosome core. Transcriptome and proteome data are marked above each gene where available. Transcripts that are up-regulated in asexual and gametocyte stages are shown as red or green horizontal lines, respectively; yellow lines denote genes that are up-regulated in both stages. Protein expression data are indicated by use of a bar code in which shading of each level indicates the following: top bar, sporozoite; second bar, oocyst; third bar, ookinete; fourth bar, gametocyte; and lowest bar, asexual stages. The identifier for every fifth gene (e.g., PF10025C) is indicated. Schematic maps of all the \textit{P. falciparum} chromosomes are shown in figs. S1 to S14.
selection that measured codon volatility in *P. falciparum* (13). There are 15 *P. berghei* genes with a dN/dS ratio > 1 that have detectable orthologs in *P. falciparum*. Not all of these have scores indicating a high volatility, a result consistent with the facts that selection will be operating at different levels in different species and that volatility and dN/dS values measure selection over different time scales.

**Gene expression.** The asexual blood stage cycle of *P. berghei* takes 24 to 24 hours and gametocyte development 30 hours. Gametocytes are morphologically discernable from the asexual trophozoites only after 18 hours (fig. S18). Transcriptome data were obtained from three time points during the G1 phase (rings and young and mature trophozoites) and from two time points during the S/M phase (immature and mature schizonts), as well as from purified immature (24-hour) and mature (30-hour) gametocytes. The transcription profile of these stages was compared through a series of pairwise hybridizations to a *P. berghei* genome survey sequence (GSS) amplicon DNA microarray (4). Proteome data were collected from mixed asexual blood stages (containing both invasive and replicative stages), gametocytes during blood stage development, ookinetes, oocysts (days 9 to 12 postinfection), and salivary gland sporozoites and analyzed by multidimensional protein identification technology (14). The proteome analysis resulted in the identification of 1836 parasite proteins with high confidence (tables S6 to S8) and >5000 parasite proteins with relaxed filtering (4). By comparing expression data for the different life cycle stages, we could categorize proteins into the following four strategies of gene expression: (i) housekeeping, (ii) host-related expression, (iii) strategy-specific expression, and (iv) stage-specific expression.

**Housekeeping.** Of the 1836 proteins detected, 136 were expressed in at least four of the five stages analyzed (table S8). Given the lower number of proteins identified in the oocyst (277 proteins) and the sporozoite (134 proteins) compared to the other stages analyzed (733 to 1139 proteins), our analysis will have excluded some of the 301 proteins detected in asexual blood stages, gametocytes, and oocyttes (fig. 3C). Recognizing that these 301 proteins were detected in both vertebrate and mosquito stages, we anticipate that some of these will also be expressed in oocysts and sporozoites.

**Host-related expression.** The proteome and transcriptome data sets revealed that enzymes of the tricarboxylic acid cycle, oxidative phosphorylation, and many other mitochondrial proteins were up-regulated in the gametocyte when compared to the asexual blood stages and were even more abundant in the oocyst (fig. S16 and table S8). These observations suggest that, as in trypanosomes (15), mitochondrial activity increases in the gametocyte as a preadaptation to life in the mosquito vector, and are consistent with the more complex organization of mitochondria in gametocytes (1, 16). Mitochondrial activity apparently continues to increase in the oocyst.

**Strategy-specific expression.** Strategy-specific protein expression is related to invasion, asexual replication, or sexual development. We uniquely detected 966 proteins in invasive zoite (merozoite, oocyst, or sporozoite)—containing preparations, of which 234 were shared between at least two of the three invasive stages but not with the replicative or sexual stages (Fig. 3A). Gliding motility typifies the invasive stages of apicomplexans, and many proteins with a (putative) role in this process were detected. Micronemes and rhoptries are secretory organelles specific to the invasive stages. Although 10 known rhoptry proteins were detected in blood stages and sporozoites, these rhoptry proteins were absent from oocyttes. In contrast, most known micronemal protein families were detected in all zoites but with clear stage-specific expression of different family members. Perforin-like proteins, first described in the micronemes of *P. yoelii* sporozoites (17), contain a membrane attack complex/perforin (MACPF)–like domain and were found both in oocyttes and sporozoites but not in merozoites. We suggest a role for these molecules in parasite entry to and/or egress from target cells, given the role of MACPF-like domains in the formation of pores. Both the oocyst and sporozoite can traverse through several host cells (18, 19), whereas a merozoite enters a target cell only once. Our data therefore support the concept that microneme proteins mediate motility and disruption of the host cell plasma membrane and that the rhoptry proteins are essential to the invasion of the parasitophorous vacuole and host cell survival.

We uniquely detected 472 proteins in replicative stages, i.e., blood stages and oocysts (Fig. 3B). Not unexpectedly and as consistent with findings in *P. falciparum* (20–22), the majority of these genes encode proteins involved in cell growth or division, DNA replication, transcription, translation, and protein metabolism. The more detailed transcriptome analysis of blood-stage gene expression confirmed a cell cycle–related timing of transcription of these genes during the G1 and S/M phases (figs. S18 and S19) and revealed that 215 and 355 were upregulated in the G1 and the S/M phases, respectively.

During the first 18 hours of development, gametocytes and asexual trophozoites share the same features of the G1 phase of growth. Subsequently, the gametocytes differentiate into either males that prepare for DNA replication and mitosis or females that prepare for postzygotic growth. Transcriptome analysis demonstrated that 58% of the G1 proteins (125 genes) and 59.4% of the S/M proteins (199 genes) were also upregulated in gametocytes (fig. S19), and the proteome data also emphasized the similarity between protein expression in asexual blood
stages and gametocytes (514 proteins were shared between these stages) (table S8). Despite these similarities, the described unique morphologies indicate that sexual development is a fundamental developmental switch. This is shown by the specific up-regulation of 977 genes (4) (table S10 and fig. S19), including many of the known gametocyte-specific genes, and by the detection of 127 unique proteins in the proteome (Fig. 3C).

Stage-specific expression. Just over half (948) of the proteins detected in the proteome analysis were found in one stage only, suggesting that stage-specific specialization is substantial. However, many of these stage-specific proteins belong to protein families whose expression is strategy-specific, reflecting both conserved mechanisms of parasite development between different stages and subtle molecular adaptations dictated by specific parasite-host interactions. For example, gene families encoding proteins that contain MACPF-like or TSP/ vWA domains are examples of strategy (invasion)–specific expression whose members are stage-specifically expressed. Unexpectedly, the PIR family belongs to this category: Members of the BIR protein family were detected in all stages, but 92% were exclusive to a single stage (fig. S15 and table S8). Peptides were found matching 34 of ~180 predicted Plasmodium genes, and transcription of bir genes was detected in both the asexual blood stage and gametocytes (tables S9 and S10). Although pir are thought to play a role in immune evasion of the blood stages by antigenic variation (7), ~9% of the total BIR repertoire in our analysis was expressed only in the mosquito stages, suggesting that these proteins may have other key functions.

Posttranscriptional gene silencing (PTGS). It has been proposed that transcripts in Plasmodium are essentially produced when needed (22), the so-called “transcripts to go” model (23). However, it has been established that the abundant transcripts for P28 in developing and mature female gametocytes are in a state of translational repression (TR) (24), one mechanism by which PTGS is exercised. In addition, RNA binding proteins (Puf proteins) (25) that play a role in TR are found in Plasmodium and are specifically up-regulated in gametocytes and sporozoites (20, 26). Therefore, we compared the gametocyte transcriptome with the proteomes of both gametocytes and oocokites to determine if additional gametocyte-specific transcripts might be subject to TR. Nine new genes were identified for which transcripts were detected in gametocytes but with protein products specific to the oocokine stage (Fig. 3D and table S11). The analysis of the 3′ untranslated regions (UTRs) of seven of these genes (for two genes, there was insufficient 3′UTR sequence for analysis) and the 3′UTRs of Pbs28 and Pbs25 by the motif identifier program MEME (27) revealed a 47-base motif found in six of the analyzed sequences within 1 kb of the 3′ end of the stop codon (Fig. 3E and fig. S17) (E value = 4.6e-002). Puf proteins bind to a UUGU motif in 3′UTR regions (25, 28), and the 3′UTR regions of all seven candidates and Pbs28 were enriched for this motif (P ≤ 0.001), which was found as a submotif in the 47-base motif. The 47-base motif was used to search the entire P. berghei genome database with MAST (27), and 20 additional genes were identified that had the same motif within 1 kb of their 3′UTR (E < e-05), giving a total of 29 TR candidates. Of these, 22 had orthologs in P. falciparum. Eighteen are up-regulated in gametocytes (16 genes) and/or sporozoites (5 genes), but only two were observed in gametocyte proteomes (table S11). Analysis of 1 kb downstream of the stop codon of 20 of these P. falciparum orthologs, including pfs25 and pfs28, failed to identify a sequence analogous to the P. berghei motif. Nevertheless, visual inspection identified numerous UUGU motifs at analogous positions. This lack of sequence similarity of the predicted 3′UTR binding motif is consistent with the significant sequence diversity in the predicted gene models of the Puf orthologs of P. falciparum and P. y. yoelii (25). The paucity of annotated transcription factors (2, 28) and the phased expression of blood-stage transcripts have led to the proposal that PTGS is a major mechanism of the regulation of gene expression in Plasmodium (28). Our data suggest that, at least in the gametocyte and possibly the sporozoite, TR may be an important component of these regulatory mechanisms.

The integration and initial analysis of the four data sets presented here has permitted insights concerning genome evolution, expression of multigene families, and mechanisms of posttranscriptional gene regulation in rodent malaria parasites. This initial overview will be developed further and, as demonstrated here, will continue to emphasize the value of model systems for the study of orthologous features of human malaria parasites.

References and Notes
Gigantic Photoresponse in ½-Filled-Band Organic Salt (EDO-TTF)$_2$PF$_6$

Matthieu Chollet,1 Laurent Guerin,1,2 Naoki Uchida,1 Souichi Fukaya,1 Hiroaki Shimoda,1 Tadahiko Ishikawa,1 Kazunari Matsuda,3 Takumi Hasegawa,4 Akira Ota,5 Hideki Yamochi,6,8 Gunzi Saito,5 Ryoko Tazaki,7,8 Shin-ichi Adachi,7,8,9 Shin-ya Koshibara1,7,8*

We report that the organic salt (EDO-TTF)$_2$PF$_6$ with ½-filled-band (½-filled in terms of holes), which forms an organic metal with strong electron and lattice correlation, shows a highly sensitive response to photoexcitation. An ultrafast, photoinduced phase transition from the insulator phase to the metal phase can be induced with very weak excitation intensity at near room temperature. This response makes the material attractive for applications in switching devices with room-temperature operation. The observed photoinduced spectroscopic change shows that this photoinduced phase transition process is caused by the cooperative melting of charge ordering assisted by coherent phonon generation.

Organic charge transfer (CT) A$_2$B salts composed of a cation (or anion) “A” and a counter ionic “B” in the ratio of 2:1 can display a variety of electronic and magnetic properties, such as superconductivity ($I_1$, $I_2$), metal-insulator (M-I) transition ($I_3$–$I_4$), magnetic frustration ($I_5$), ferroelectricity ($I_6$), and even magneto-dielectric coupled behaviors ($I_7$). The appearance of charge ordering (CO) or a Mott transition accompanied with dimeric distortion in a ½-filled or a ¼-filled (½-filled in terms of holes) network of A molecules with a one-dimensional ($I_8$) or 2D structure plays a key role as the basis for these exotic natures ($I_1$, $I_2$). Recent theoretical studies have revealed that the critical balance among physical parameters, such as bandwidth ($H$), onsite Coulomb interaction ($U$), and nearest-neighbor Coulomb interaction ($V$), leads to the appearance of such novel ground states ($I_7$–$I_{10}$).

These previous studies and a report on the highly efficient photocarrier generation in one of the A$_2$B salts (DCNQI)$_2$Cu (I1) have stimulated research on the photo-induced phase transition (PIPT) in A$_2$B salts. In a crystal that shows multi-instability of its free energy due to critical balance among intrinsic cooperative interactions, the macroscopic phase transition accompanied with large changes in electronic, magnetic, and lattice structures may be triggered by weak photoexcitation ($I_2$, $I_3$), because even a low density of photoexcited species can affect and switch the cooperative interaction ($I_{12}–I_{14}$). From the viewpoint of the highly efficient and nonlinear amplification of the response to the weak photoexcitation via a cooperative channel in a condensed system, this exotic photo effect called PIPT is analogous to the “domino effect” on the molecular scale.

Here, we report that a quasi-1D, ½-filled (in terms of holes) A$_2$B salt (EDO-TTF)$_2$PF$_6$ shows highly sensitive and ultrafast PIPT from an insulator (I) phase accompanied with a CO to a metal (M) phase up to about room temperature (~265 K). [EDO-TTF]

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References and Notes

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Materials and Methods

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SuppFig. 1

Supporting Online Material

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