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Timothy S. Little

The Francis Crick Institute

Deirdre A. Cunningham

The Francis Crick Institute

Audrey Vandomme

The Francis Crick Institute

Carlos Talavera Lopez

Institute of Computational Biology

Sarah I. Amis

The Francis Crick Institute

Christopher Alder

The Francis Crick Institute

John W.G. Addy

The Francis Crick Institute

Sarah McLaughlin

The Francis Crick Institute

Caroline Hosking

The Francis Crick Institute

George K. Christophides

Imperial College London

Adam J. Reid

Wellcome Genome Campus: Wellcome Sanger Institute

Jean Langhorne (✉ Jean.Langhorne@crick.ac.uk)

Francis Crick Institute <https://orcid.org/0000-0002-2257-9733>

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Analysis of *pir* gene expression across the *Plasmodium* life cycle

Timothy S. Little^{1*}, Deirdre A. Cunningham^{1*}, Audrey Vandomme¹, Carlos Talavera Lopez^{1,4}, Sarah Amis^{1,5}, Christopher Alder¹, John W.G. Addy¹, Sarah McLaughlin¹, Caroline Hosking¹, George Christophides⁶, Adam J. Reid² and Jean Langhorne¹

¹The Francis Crick Institute

²Wellcome Sanger institute, Cambridge, UK, CB10 1SA

⁶ Imperial College London

*shared first authorship

Present address:

⁴Carlos Talavera-López: Institute of Computational Biology; Helmholtz Zentrum für Gesundheit und Umwelt; Munich, Germany.

Corresponding author: Jean.Langhorne@crick.ac.uk

Abstract

Background: *Plasmodium* interspersed repeat (*pir*) is the largest multigene family in the genomes of most *Plasmodium* species. A variety of functions for the PIR proteins which they encode have been proposed, including antigenic variation, immune evasion, sequestration and rosetting. However, direct evidence for these is lacking. The repetitive nature of the family has made it difficult to determine function experimentally. However, there has been some success in using gene expression studies to suggest roles for some members in virulence and chronic infection.

Methods: Here we examined *pir* gene expression across the life cycle of *P. berghei* using publicly available RNAseq data-sets, and at high resolution in the intraerythrocytic development cycle using new data from *P. chabaudi*.

Results: Expression of *pir* genes is greatest in stages of the parasite which invade and reside in red blood cells. The marked exception is that liver merozoites and male gametocytes produce a very large number of *pir* gene transcripts, notably compared to female gametocytes, which produce relatively few. Within the asexual blood stages different subfamilies peak at different times, suggesting further functional distinctions. Representing a subfamily of its own, the highly conserved ancestral *pir* gene warrants further investigation due to its potential tractability for functional investigation. It is highly transcribed in multiple life cycle stages and across most studied *Plasmodium* species and thus is likely to play an important role in parasite biology.

Conclusions: By identifying distinct expression patterns for different *pir* genes and subfamilies we hope to provide a basis for the design of future experiments to uncover their function.

Introduction

The genomes of the malaria parasites (*Plasmodium* spp.) contain a variety of multigene families. These are generally located in the sub-telomeric regions of chromosomes, a feature which is thought to allow regulation of gene expression by heterochromatin and promote diversification of the gene sequences through recombination [1-3]. The most highly studied of these families is *var*, which encodes the PfEMP1 protein. It is present in ~70 copies in the human malaria parasite *Plasmodium falciparum*. The primary role of PfEMP1 appears to be sequestration of the parasite in the vasculature, preventing its destruction in the spleen [4]. Multiple copies of *var* seem to be required to alter the binding specificity and for immune evasion - i.e. switching between antigenically distinct proteins with similar binding properties [5]. This family is however limited to the *Laverania* subgenus, which includes only one of the five species which infect humans. The *Plasmodium* interspersed repeat (*pir*) multigene family has been found in the genomes of rodent malarias, primate malarias and the human-infecting *Plasmodium* species, *P. vivax*, *P. knowlesi*, *P. malariae* and *P. ovale* [6-9]. The number of *pir* genes in the genomes of these species varies considerably, from the lower numbers of 211 members in *P. chabaudi chabaudi* AS and 134 members in *P. berghei*, to more than 1,000 members in *P. yoelii* and up to 1,949 members in *P. ovale curtisi* [9, 10]. The *pir* family is therefore considered to be the largest *Plasmodium* multigene family.

Although it has been suggested that the *pir* family fulfils similar roles in immune evasion and pathogenesis as *P. falciparum var*, there is no direct evidence for this, and the function(s) of *pirs* remains largely unknown. While *P. falciparum* parasites can only express a single *var* gene at a time, individual *P. berghei*, *P. vivax* and *P. yoelii* parasites can express multiple *pir* genes [11-15]. Although some proteomic and immunofluorescence studies show PIR proteins on the surface of infected RBCs (iRBCs), other studies indicate that they are present in the host or parasite cytoplasm, or on the parasitophorous vacuole [6, 16-19] suggesting multiple different functions for PIR proteins during blood-stage infections. In *Plasmodium chabaudi chabaudi* AS infections of mice, different *pir* subfamilies are associated with the acute and chronic phases of the infection, and parasites from the two phases of infection are differently virulent [20]. This association suggests *pir* gene expression may affect virulence of blood-stage *P. chabaudi*, and also that *pir* genes could be involved in evading the initial immune response. Single-cell RNA-seq analysis across the *P. berghei* life cycle has shown that *pir* gene expression is particularly high in blood stages [14]. Male gametocytes were found to express a distinct *pir* repertoire compared to asexual

blood stages, with female gametocytes expressing few *pir* transcripts [13]. Although these data represent only the most highly expressed genes, they suggest there may be different functions for *pir* genes between asexual and sexual blood stages.

This family has been very difficult to study in the laboratory due to there being many genes with varying levels of similarity expressed at a variety of times. Identifying which are expressed at particular life-cycle stages will allow a more targeted approach to determining function. Here we use two rodent-infecting *Plasmodium* species to investigate the stage-specificity of *pir* gene expression. We performed a systematic analysis of the *P. berghei* ANKA *pir* multigene family from published bulk RNA-seq studies, re-processing the raw data using a single bioinformatics pipeline. While single-cell RNA-seq datasets have provided excellent resolution of gene expression in time across the life cycle [14], bulk RNA-seq datasets provide much higher resolution of the transcriptome itself, detecting a larger proportion of transcripts expressed at any one time. Indeed, expression of relatively few *pir* genes was detected in the Malaria Cell Atlas [14]. To examine the dynamics of *pir* expression in asexual blood stages in more detail, we also generated high-resolution transcriptional data from *P. c. chabaudi* AS, across the asexual blood cycle.

Our study demonstrates that the primary function of the *pir* multigene family likely plays out in the blood stages, from merozoite formation in the liver to fertilisation in the mosquito midgut. It is unlikely to be important during the post-fertilisation stages of development in the mosquito through to early liver stages. In both rodent malaria models the *pir* transcriptional repertoire is diverse throughout the intraerythrocytic developmental cycle, however different subfamilies are differently represented over time, suggesting subtle differences in regulation and perhaps function. One *pir* gene, previously described as the putative ancestral *pir* [21], is clearly notable as forming the highest proportion of the *pir* transcriptome of these two species, and of multiple different species of *Plasmodium* for which transcription data is available. This gene may prove a more experimentally tractable target due to its uniqueness and warrants further study.

Methods

Mice

Female C57BL/6 aged 6-8 weeks from the SPF unit at the Francis Crick Institute Mill Hill Laboratory were housed under normal (light 07.00-19.00, dark 19.00-07.00 GMT) or

reverse light conditions (light 19.00-07.00, dark 07.00-19.00 GMT) at 20-22 °C, and had continuous access to mouse breeder diet and water. This study was carried out in accordance with the UK Animals (Scientific Procedures) Act 1986 (Home Office licence 80/2538 and 70/8326) and was approved by The Francis Crick Institute Ethical Committee.

Parasites

A cryopreserved stock of a cloned line of *Plasmodium chabaudi chabaudi* AS, originally obtained from David Walliker, University of Edinburgh, UK, and subsequently passaged through mice by injection of infected red blood cells (iRBC), was used to initiate infections and mosquito transmissions. Transmission of *P. chabaudi* via *Anopheles stephensi* mosquitoes has been described in detail previously [22]. Recently mosquito-transmitted (RMT) *P. chabaudi* derived from a mosquito-initiated infection retain the same phenotype of infection course and gene expression profiles of directly mosquito-transmitted *P. chabaudi* infections [20, 23]. RMT parasites are used here to ensure that each mouse receives a consistent inoculum of blood-stage parasites, rather than the temporarily more variable appearance of parasites in the blood from either a MT- or sporozoite-initiated infection [23]. For RMT-blood stage infections, mice were infected by intraperitoneal injection of 10⁵ iRBC. Blood samples were collected at three-hour intervals over one 24hr asexual cycle. Infections were monitored by light microscopy on Giemsa-stained thin blood smears. Parasitaemia across the 24hrs ranged from 2-10% (Supplementary Information 2).

RNA extraction and sequencing

Mice infected as described above were segregated into eight groups (3 mice/group) according to the scheduled time points 02:00; 05:00, 08:00, 11:00, 14:00, 17:00, 20:00, 23:00 and at each time point, day 7 post-infection, a group was smeared, exsanguinated and blood collected for RNA extraction. Blood samples were depleted of leukocytes by filtration (Plasmodipur, Euro Proxima) and of globin RNA by saponin lysis and centrifugation and RNA extracted as previously described [23]. Briefly, purified parasites were resuspended in Trizol, frozen at -80°C until use. RNA was then extracted and resuspended in water.

P. chabaudi RNA samples were used to make 200-350bp fragment Illumina TruSeq libraries, with 12 cycles of PCR amplification. All 24 samples were tagged, pooled and sequenced on each of two Illumina HiSeq2500 lanes with paired 100bp reads. Recently mosquito-

transmitted RNA samples were used to make 150-250bp fragment Illumina TruSeq libraries with 12 cycles of PCR amplification. All 24 samples were tagged, pooled and sequenced on a single HiSeq2500 lane with paired 100bp reads.

RNAseq analysis

RNAseq datasets were downloaded from SRA using SRA-Tools (SRA_Toolkit_Development_Team), except for the *P. chabaudi* data and the *P. berghei* data from Witmer et al., [24] and Ukegbu & Giorgalli et al., [25] which was transferred manually from the original lab data storage. Technical replicates were concatenated together as fastq.gz files. Both the *P. berghei* and the *P. chabaudi* RNAseq were processed through the rna-seq pipeline v1.4.2 from nf-core repository in the Nextflow pipeline software v20.07.1 [26, 27], using Trim Galore! v0.6.4 to trim sequences, HISAT2 v.2.1.0 as the aligner, and featurecounts v.1.6.4 as the gene count program [28-30]. Quality control was analysed using FastQC v.0.11.8 and collated by MultiQC v.1.7 [31, 32]. Paired-end sequence files and single-end sequence files for *P. berghei* were separated and analysed in different iterations of the same pipeline. PlasmoDB v48 annotations (supported by GeneDB) and genome assemblies were used [10, 33].

For the *P. berghei* datasets the liver stages, and some of the asexual blood stages (rings, trophozoites and schizonts), were defined by time of development in culture. In some published data sets, male and female gametocytes were analysed separately, while in others gametocytes were mixed. The 6h Liver samples from [34] were removed due to low read counts, also removed by the authors in the original publication. Additionally, a 2h Liver sample (SRR11142819) from [35], and two schizont samples (SRR3437888 and SRR3437912) from [36] were removed because hierarchical clustering showed that they were dissimilar to the other samples.

The gene counts were imported into R v4.0.2 where they were normalised by gene length and library size into TPM (R Core Team, 2018). TPM was chosen over RPKM/FPKM (RNA/Fragments Per Kilo) since TPM always totals 1 million in a given sample, while per sample sum of RPKM varies [37].

Pearson correlation (Supplementary Figure 1A) and Principal Components Analysis (PCA; Supplementary Figure 1B) demonstrate that samples from the same life cycle-stage(s) in

different experiments robustly cluster together. PCA was calculated using R's `prcomp` function and the variables were standardised and scaled ('center' and 'scale.' set as TRUE). We concluded that gene expression could be accurately compared between studies, and thus we combined the datasets together to obtain one average sample per life-cycle stage (Supplementary Information 2). Mean expression was calculated, and the median was not used because this leads to the TPM of groups no longer adding up to a constant value. Some life cycle stages from slightly different times or conditions were grouped together as global gene expression was similar, such as liver stage samples at 4h post-infection or earlier (Liver.pre4h), liver stage samples at 48h post infection or later (Liver.post48h), the chronic and acute samples from Brugat et al., 2017 (Asexual.Mixed) [20], the different times of ookinete development from Otto et al., 2014 (Ookinete)[12], the 'activation' state of gametocytes from Ukegbu & Giorgalli et al., 2020 (Gametocytes) [25], 3h post-uptake and earlier blood meal samples from Ukegbu & Giorgalli et al., 2020 (BloodMeal-Pre3h) [25], and the liver stages from different cell lines.

The annotations of *pir* genes in each species were taken from PlasmoDB v48 and can be found on the first sheet of Supplementary Information 2 (*P. berghei*) and the second sheet of Supplementary Information 6 (*P. chabaudi*). Note that this version of PlasmoDB annotated the orthologous genes PBANKA_0524600 and PCHAS_0524800 as members of the *pir* family, but we believe this to be erroneous due to their two-exon structure and lack of similarity with other *pir* genes. Hence, they were excluded from analysis here. The ChAPL/AAPL information for *P. chabaudi* was taken from Brugat et al., 2017 [20].

R packages used in manipulating and analysing the data include `dplyr` [38], `data.table` [39], `reshape2` [40], `stringr` [41], `tibble` [42], `readr` [43], and `readxl` [44][56]. Visualisation was achieved using `ggplot2` [45], `ggpattern` [46], `RColorBrewer` [47], `corrplot` [48], `plotly`, `viridis` [49], `circlize` [50], and `ComplexHeatmaps` [51].

Deconvolution of *P. chabaudi* bulk transcriptomes

To determine the relative proportions of different life stages in the *P. chabaudi* transcriptomes, we used the approach described in Aunin et al. [52]. Briefly, we used pseudobulk samples (excluding mosquito and liver stages) derived from the Malaria Cell Atlas [14] as a reference to deconvolute with CIBERSORT v1.06 using default settings [53].

Statistical analysis

Statistical analyses of gene expression were performed on R v4.0.2. For sets of genes (such as all *pir* genes, sub-families, and Short or Long-form groups) Qusage v.2.22.0 [54] was used with the TPM data, using 2e18 iterations. For individual genes DESeq2 v1.28.1 [55] was used, applying apeglm v1.10.0 [56] log fold change shrinkage to reduce the impact of lowly expressed genes on the differential expression analysis. Individual p values for comparisons were collated and multiple testing correction by the false discovery rate was calculated using R's p.adjust function. In each case the different biological replicates from every experiment were treated as independent samples.

Phylogeny

A 'transformation by orthology' search was conducted on PlasmoDB v48 using *P. berghei* ancestral gene *PBANKA_0100500*, which uses OrthoMCL for this purpose [57]. The *P. c. chabaudi* CB genome was not on PlasmoDB at the time of analysis so this ancestral gene was identified using a BLAST search with the AS strain gene. With nineteen orthologs from the different *Plasmodium* species, MUSCLE (v3.8.31) alignment [58] and PhyML (v3.1) tree-building with LG model 100 bootstrap replicates (otherwise default settings) [59] was conducted on SEAVIEW v4.7 [60]. The tree was imported to R and some additional strains of the same species were removed to simplify the final tree. The R packages used for tree-design were treeio v.1.12.0 [61], ggtree v2.2.1 [62] and ape v5.4 [63].

Ancestral *pir* gene transcriptional analysis

We then investigated whether the orthologs of the ancestral *pir* gene were also highly transcribed in other *Plasmodium* species for which transcription data is available. TPM/RPKM from published studies (Table 2) was used to calculate the proportion of ancestral *pir* expression relative to the rest of the transcriptome.

Data availability

The RNA-seq data relating to *P. c. chabaudi* AS intraerythrocytic developmental cycle are available from the ENA (ERP002273). The relationship between individual samples and ENA accessions is described in Supplementary Table 1.

Results

Asexual blood stages, liver merozoites and male gametocytes are the foci of *pir* gene expression in *Plasmodium berghei*.

The rodent *pir* gene family is divided into two groups named S and L based on sequence similarity and average gene length [12]. These are further classified into subfamilies (L1-L4; S1-S8), with some being unique to a particular rodent species (e.g. S7, *P. chabaudi*; S8, *P. berghei*). In order to determine whether an association exists between transcription of any *pir* subfamily and a particular life cycle stage, we performed a systematic analysis of published transcriptome studies from the rodent *Plasmodium* species, *P. berghei* ANKA (Figure 1, Table 1, and Supplementary Information 1). These included samples from most of the life cycle stages of the parasite in both the mosquito and mouse hosts.

The gene expression values (TPM; transcript-per-million) for the different samples across all genes were used for data normalisation. The individual samples from different experiments separated robustly by stage of development of the parasite (Supplementary Figure 1). As sexual stages may be present in host blood, we used the transcriptional activity of selected sexual stage markers to confirm the relative purity of asexual blood stage samples from across the multiple experiments (Supplementary Figure 2). The genes selected were *p28* (*PBANKA_0514900*) [64] and *nek4* (*PBANKA_0616700*) (female gametocytes and ookinetes) [65], *mapk2* (*PBANKA_0933700*) (male gametocytes) [66], and *hap2* (*PBANKA_1212600*) (gametes) [67]. Their transcriptional activity confirmed that the asexual blood stages had few gametocyte contaminants.

The genome of *P. berghei* ANKA contains 134 *pir* genes. Although multiple *pir* genes are transcribed at all stages of the life cycle (Figure 2A), this begins at very low levels in salivary gland sporozoites and early liver stages, and is then followed by an increase in both the level of transcript, and the total number of *pir* genes transcribed, in the later liver stages. One study analysed RNAseq data from liver merozoites [34], and we find that they display a dramatic increase in the levels of *pir* gene transcripts over earlier liver stages. High levels of transcript are maintained in asexual blood stages declining somewhat by the schizont stage. There are relatively few *pir* transcripts in female gametocytes, whereas the level of *pir* transcription in male gametocytes is of a similar or greater magnitude and breadth to that observed in asexual blood stages. Ookinetes within the mosquito midgut transcribe fewer *pirs* at lower transcription levels, similar to late liver stages, and female gametocytes.

There is one subfamily of *pirs*, distinct from S and L, containing only a single member. It has syntenic orthologues in the genomes of rodent and primate-infecting *Plasmodium* species. This *pir* and its orthologs have been described as the ancestral *pir*, because the other subfamilies may have derived from it [20]. The *P. berghei* ancestral *pir* orthologue (PBANKA_0100500) is consistently highly transcribed from late liver stages, through asexual intraerythrocytic development and in blood stage gametocytes (shown in yellow in Figure 2A and B), dominating the *pir* transcriptional profile whenever *pirs* are highly expressed.

The transcription of L- and S-*pirs* in sporozoite and early liver-stage parasites broadly reflects the genome composition (20% L, 79% S). However, in the later liver stages (post 48h) L-*pirs*, predominantly of the L1 clade, comprise a distinctly higher proportion of the transcripts (61% of TPM; Figure 2B). The increase in expression of the *pirs* between the 24h liver stages and the post 48h liver stages is concentrated in the L-forms (all *pirs* between liver post 48h and 24h: log₂ fold change 0.67, FDR 0.029; L-form *pirs*: log₂ fold change 1.56, FDR < 5x10⁻⁹).

Early intraerythrocytic asexual stages show a lower proportion of L-*pir* transcripts (7-9% in rings and trophozoites), but rising to 33-34% in schizonts – dominated by L2 *pirs*. The stage with the next highest proportion of L2 *pirs* is late liver stages (post 48h), suggesting that the highest proportion of L2 transcripts is associated with development of extracellular invasive forms (merozoites). L1 *pirs* have highest proportions of total *pir* transcripts in the late liver stages. Note that we have previously shown that *P. berghei* L2s have more similar sequence properties to *P. chabaudi* L1s despite their names [20]. The gametocyte stages express slightly lower relative proportions of L-*pir* transcript (7-11%). Members of the S sub-family form a greater proportion of the *pir* transcriptome of rings, trophozoites and gametocytes (Figure 2A, 2B), with the largest subfamily, S1 (Figure 2B), making up the majority of *pir* transcripts.

For those samples which exhibit high levels of *pir* transcriptional activity (male gametocytes, liver merozoites and mixed asexual stages) differential expression analysis of the subfamilies was conducted (Supplementary Information 3). While a number of different subfamilies were found to be differentially expressed to statistical significance in individual pairwise comparisons, no subfamilies were identified as significantly up- or downregulated in the mixed asexual stages compared to both the liver merozoites and the male gametocytes. Thus, for most sub-family divisions all the families tend to increase or

decrease in transcription in tandem with the rest of the *pir* family, however there is an association between L-form *pir* and merozoite production.

Although transcription of individual *pir* genes tended not to be stage-specific, there were *pirs* clearly enriched in certain stages of the life-cycle (Supplementary Information 4). These included the L1 *pirs* PBANKA_0317181 in male gametocytes (compared to mixed asexual stages: log fold change 5.09, FDR < 5×10^{-9} ; compared to liver merozoites: log fold change 10.65, FDR < 5×10^{-7}) and PBANKA_0600031 in liver merozoites (compared to mixed asexual stages: log fold change 5.69, FDR < 5×10^{-7} ; compared to liver merozoites: log fold change 6.82, FDR < 5×10^{-6}). The *pir* gene PBANKA_0400500 was highly transcribed in two of the three experiments performed for the later liver stages (post.48h). This gene is not restricted to liver-stage expression, although notably high in the liver stages, suggesting that it may play a role in late exo-erythrocytic and early asexual stage parasites.

Differential timing of expression of L and S *pir* gene subfamilies during the blood cycle of *P. chabaudi*

We have shown that *pir* gene expression levels in *P. berghei* are low in mosquito stages, but high in the mouse from late liver to asexual blood stages and male gametocytes. We wanted to explore the expression of rodent *pir* genes at higher resolution across the asexual blood stages to determine whether there were more subtle differences in expression of the different subfamilies. To do this, we generated transcriptional data from another rodent malaria parasite, *P. chabaudi chabaudi* AS (Supplementary Information 5). This parasite has a largely overlapping repertoire of *pir* genes. We have a good understanding of *pir* gene expression at 14h post infection and are able to transmit it by mosquito. We have shown that compared to serial blood passage this results in expression of a wide repertoire of *pir* genes which we expect is more representative of the situation in the wild [20, 23].

Microscopic analysis of the proportions of ring and trophozoite forms at each time point shows that development is largely synchronous (Figure 3A; Supplementary Figure 3; Supplementary Information 6). A deconvolution approach using previously published single-cell RNA-seq data [14] confirmed the enrichment of the individual stages at the appropriate time points (Supplementary Figure 4A). Principal Components Analysis of the RNAseq time-points confirmed that most of the different samples form a continuum of transcription over time, with a 'gap' between 14h/17h around when schizonts develop and sequester. This verifies the robust capture of the trophozoite and ring stages of the 24h asexual blood cycle (Supplementary Figure 4B).

Pir genes show a cyclical expression pattern across the developmental cycle, with the majority upregulated at 8-14h (trophozoite stage) (Figure 3B; Supplementary Information 6). This signal suggests that *pir* genes may be required in the schizonts and merozoites, as the transcriptional signal is expected to precede translation of protein by several hours (22). *Pirs* upregulated at ring stages were predominantly of the L1 subfamily, which have a slightly earlier transcriptional peak (8-11h) in the developing trophozoites and are quiescent only in the early ring stages. Transcription of both the S7 and S1 subfamilies peaks sharply at 14h. L4s notably exhibit two distinct temporal *pir* transcription profiles, with a proportion being similar to L1s and the remainder being like the short S *pirs* where (Figure 3B). The ancestral *pir* gene, PCHAS_0101200, is transcribed throughout the intraerythrocytic developmental cycle, but still peaks during the trophozoite stages (Figure 3B and Supplementary Figure 5), as we observed for *P. berghei*. This gene is one of the most highly transcribed *pir* genes, second only to the *pir* PCHAS_0600600.

Brugat et al., 2017 [20] described distinctive clusters of *pir* genes associated with the acute (Acute-Associated *pir* loci or AAPL) and chronic (Chronic-Associated *pir* loci or ChAPL) phases of Mosquito Transmitted (MT) *P. chabaudi* infection. Here, we find that temporal transcription of clusters of *pirs* at ChAPL loci on chromosomes 3 and 6 mirrors that of L1s, and early peaking L4s, as expected, as most ChAPL *pirs* are L1s, while clusters at AAPL loci, mostly S7s and late peaking L4s, peak sharply in late trophozoites (14h) (Figure 3C), suggesting that they perform separate functions during parasite development.

The putative ancestral *pir* gene is a distinctive target for functional studies

We have shown above that the *P. berghei* ancestral *pir* orthologue, *PBANKA_0100500*, is the single most highly transcribed *pir* gene in this species. It contributes a high proportion of the total *pir* steady-state transcriptome in late liver, asexual blood and gametocyte stages. Transcription is only diminished in stages where overall transcription of this multigene family is low i.e. liver stages before 36h of development, ookinetes and sporozoite stages. Similarly, in *P. chabaudi* AS, the ancestral *pir* orthologue (PCHAS_0101200) is highly transcribed across the complete asexual developmental cycle (Supplementary Figure 5). Extending earlier findings, we found that there is a single, syntenic ortholog in all species whose genomes contain canonical *pirs* [10, 20](Figure 4A). The sequences are highly conserved between species, with multiple blocks of 90-100% similarity (Figure 4B). The gene

is within the top 50% of expressed genes in all species for which we could find RNAseq data (Figure 5).

Discussion

We found that high numbers of *pir* gene transcripts were expressed from a variety of family members in stages which produce the invasive blood forms (merozoites) and in male gametocytes. Conversely, only low levels were expressed throughout the mosquito stages, early vertebrate stages and in female gametocytes. This strongly indicates that this multigene family is not required for development within the mosquito beyond sexual reproduction. Instead, it is likely that the bulk of this large family, at least in rodent malaria parasites, is involved in blood stages, leading from red cell invasion by first generation (liver) merozoites to fertilisation and generation of the ookinete form rapidly after mosquito blood feeding. We have previously shown the high levels of *pir* gene expression in *P. chabaudi* parasites in asexual blood stages [22, 23] and the specificity of *pir* gene expression in male *P. berghei* gametocytes versus female gametocytes [13]. This new analysis looks broadly and deeply, confirming the findings based on much lower coverage Malaria Cell Atlas data [14].

We found that L *pir* genes are highly expressed in late liver stages, merozoites and schizonts. These would require some time before being translated into functional proteins and it may be that they function in rings, e.g. soon after red cell invasion. Expression of S-type *pirs* is more steady throughout the development cycle. L-type *pirs* are first transcribed in mature liver schizonts, and upregulated again in asexual blood schizonts, so they are potentially involved in merozoite formation/function. When we look in more detail at this part of the lifecycle in *P. chabaudi* we see dynamic expression of each individual *pir* gene, as has been shown for genes across the *Plasmodium* genome [68]. Again, L and S types behave differently, suggesting differential regulation and function. Underlying this pattern is the relatively early and broad expression of the ChAPL loci, which are rich in L1 and a subset of L4 *pirs*, so they may be required throughout the cycle. The S-rich AAPLs peak later and more sharply, and may only be needed for one stage of the asexual cycle. We previously showed the ChAPLs to be associated with chronic infections. We think that populations of parasites expressing ChAPLs survive the acute immune response, while those expressing AAPL loci are killed [20]. Whether this relates to a function in sequestration, evasion of adaptive immunity or a completely different function is unclear. Molecular mimicry as a means of immune suppression has been described for *P. falciparum* RIFINs, another sequence-variable

multigene family, some of which mimic the natural ligand for LILRB1 [69] and suppress the activity of NK cells. *P. knowlesi pir* sequences have also been shown to have a striking resemblance to 50% of CD99, a T-cell regulating protein, suggesting molecular mimicry and potential immune-modulating activity [8]. Such interactions could promote chronicity, although the mechanisms of the contribution of AAPs and ChAPs in infection outcome have yet to be elucidated.

The other focus of *pir* gene expression is in male gametocytes, with a relative absence of expression in females. A small number of L1 *pir* genes are highly expressed at this stage in *P. berghei*, but S1s and S4s predominate. Could this represent a continuation of function from asexual to sexual parasites? Male gametocytes are found in the bloodstream, whereas immature female gametocytes reside in bone marrow, primarily in the extravascular space. Here, cellular rigidity may be more important than the receptor-mediated interactions of sequestration within the vasculature [70-72].

Our analyses indicate that expression of multiple *pirs* occurs in the parasite developmental stages which are predominantly circulating in the blood, where the parasite-infected cells are targeted by the host's immune system. PIR proteins have been demonstrated to be targeted by antibodies [6] and several studies have shown localisation at or near the parasite surface [16, 19, 73]. Recent structural studies together with sequence analysis and modelling have revealed hydrophobic conserved disulphide bonds forming cysteine residues suggesting that part of the PIR protein may be extracellular [34]. It is hypothesised that hydrophobic domains are exposed on the surface via a flexible linker. The diversity in sequence and length of the flexible loops are consistent with a surface location, and thus could interact directly with the host immune system or be involved in sequestration.

PIRs localise to different cellular compartments depending on the particular stage of asexual development e.g RBC cytoplasm and parasitophorous vacuole [19], [74]. This raises the possibility that PIR proteins are likely to have functions in addition to immune evasion or immune-suppression in the mammalian blood stream. Our transcriptional analyses do not shed light on this. However, we see an association between higher *pir* transcription and the stages involved in or just preceding proliferative steps of development, such as liver schizonts, asexual blood stages (specifically the trophozoites in *P. c. chabaudi*) and male gametocytes, which could be indicative of a role in phase transition rather than overall heightened transcription in these stages.

Although the other major proliferation stage of the parasite, the oocyst in the mosquito, is not covered in the *P. berghei* bulk RNAseq, single cell RNAseq data suggest that *pir* transcription is low in more mature oocysts. However, the earlier oocysts may be high transcribers. Bulk RNAseq analysis of oocysts from rodent *Plasmodium* species at different times post-ookinete differentiation would provide crucial data to investigate this link.

The high level of transcription of one, conserved *pir* gene, across most *Plasmodium* parasites suggests that it may serve an important function in the parasite, again in the blood cycle. Transcription is upregulated just prior to entry into the blood stream and is maintained throughout the erythrocytic developmental stages. The *P. cynomolgi* RNAseq data demonstrated that transcription of its ortholog was much lower than for most of the transcriptome, however this data is from liver stages in which *pir* transcription may be low. Unlike most of the other *pir* family members vector transmission has little impact on the transcription levels of this gene [75][36]. Although its distinctiveness suggests that it may fulfil a different role from other *pirs*, this gene may prove a more tractable target for future studies. We believe that it will provide insights into the molecular function of the whole family.

Conclusion

The *pir* gene family is the largest found in malaria parasites, with potentially important roles in virulence and chronic infection. We have characterised the landscape of *pir* gene expression across the *Plasmodium berghei* lifecycle, highlighting the blood stages as the focus of activity. In depth analysis of the blood stages using the close relative *P. chabaudi* highlighted subtle differences in the timing of expression of different *pir* gene subfamilies. However, the most distinctive expression pattern we found was for the putative ancestral *pir* gene, conserved across much of the *Plasmodium* genus, and very widely and highly expressed. The distinctiveness of this *pir* gene may make discovering its function more tractable while still shedding light on those genes already considered to be involved in host-parasite interactions.

Abbreviations

AAPL: Acute-Associated *pir* loci

ChAPL: Chronic-Associated *pir* loci

iRBC: infected red blood cell

P: *Plasmodium*

pir/PIR: *Plasmodium* interspersed repeat

PCA: Principle Components Analysis

MT: mosquito transmitted

RMT: recently mosquito transmitted

Declarations

Ethics approval and consent to participate

This study was carried out in accordance with the UK Animals (Scientific Procedures) Act 1986 (Home Office licence 80/2538 and 70/8326) and was approved by The Francis Crick Institute Ethical Committee.

Consent for publication

All authors have seen and approved the content.

Availability of data and materials

The RNA-seq data relating to *P. c. chabaudi* AS intraerythrocytic developmental cycle are available from the ENA (ERP002273). The relationship between individual samples and ENA accessions is described in Supplementary Table 1.

Competing interests

The authors declare that they have no competing interests

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Authors' contributions

TSL, GC, AJR and JL conceived and designed the study. TSL performed all the bioinformatics analysis, with substantial input from AJR. TSL, DAC, GC, AJR and JL analysed and interpreted the data. AV led the *P. chabaudi* *in vivo* experiments, RNA-seq sample and raw data generation. AV, SA, CH, SMcL and DAC performed the latter experiments. CA, JWGA and CT-L gave advice and feedback on bioinformatics and

statistics. TSL, DAC, AJR and JL wrote the manuscript. All co-authors have read, reviewed and approved the paper.

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Figure legends

Figure 1. Systematic study of *P. berghei* RNAseq datasets that covers its entire life cycle and demonstrates the dynamic expression of *pir* genes throughout. Schematic of the published experiments used in this study and the *P. berghei* specific developmental stages during which samples were collected. The experiment codes (E01 etc.) correspond to the references in Table 1.

Figure 2. Expression of *pir* genes throughout *P. berghei* life cycle. A: Bar chart of total *pir* gene TPM (y-axis), calculated from the mean expression across all experimental samples of each life cycle stage (x-axis). The colours of the bar chart denote the classification; ancestral (yellow), Long (L) (blue) and Short (S) (orange), with the boxes of the stacks

denoting expression of the individual genes. The background and strip name colours correspond to groups of life cycle stages, including mosquito (green), liver (pink), asexual blood (red), and gametocyte (blue) stages. The genes in each subfamily are ordered by expression levels. Inset of figure shows an enlargement of the sporozoite and liver stages. **B:** Bar chart of the proportion of total *pir* gene TPM (y-axis), contributed by the ancestral *pir* (yellow) and family sub-divisions of the Short/Long *pirs*, for each stage (x-axis). Stages are separated as in (A). Legend shows the number of genes that are members of each grouping, which are collated together for each box in the stacked bars. The ordering is by family sub-division, from ancestral, to Long groups (1-3) and finally Short groups (1-2, 4-6, 8).

Figure 3. Intraerythrocytic development and *pir* gene expression across the 24 hour asexual blood cycle of *P. c. chabaudi* AS parasites transmitted by RMT. **A:** Microscopic counts of ring, trophozoite and schizont stage parasites in Giemsa-stained blood smears, as a proportion of the total parasitaemia (n=3). Graphs show mean (+/-SD). Value 1 would indicate that all parasites counted, are of the specific developmental stage. **B:** Heatmap of *pir* gene expression, $\log(\text{TPM}+1)$, at each timepoint. Genes are grouped according to sub-family (L1, L4, S1 and S7) and ordered by mean expression across the timepoints within each sub-family, with the ancestral *pir* at the very top. Three blood stage-specific genes were included for comparison 'ref': MSP1 (highly transcribed in schizonts/merozoites), HSP70 (highly transcribed in rings/trophozoites) and AMA1 (highly transcribed in schizonts/merozoites). Genes expressed under 1 TPM in every sample were removed. **C:** Heatmap of transcription of the *pir* genes assigned to ChAPL or AAPL loci [20][20], $\log(\text{TPM}+1)$, at each timepoint, grouped according to locus.

Figure 4. Conservation and gene expression of the ancestral *pir* across the *Plasmodium* genus. **A:** Sequence alignment of 15 ancestral *pir* orthologs (i.d.verified 20210527) identified in different *Plasmodium* spp. indicating regions and sequence of high conservation. **B:** Expanded region of the most highly conserved section of 234 aa, illustrated with a sequence logo and percentage match of the consensus.

Figure 5. Phylogenetic analysis of the ancestral *pir* gene. Cladogram of the 15 orthologs, with heatmap of expression of the gene relative to the rest of the genome, where data is available (from multiple transcriptomic datasets as RPKM or TPM - see table 2). With the

exception of *P. cynomolgi* strain M the ancestral ortholog is expressed within at least the top 50% of genes and usually in the top 30% of genes.

Supplementary Figure legends

Supplementary Figure 1

(A) Pearson correlation plot of each transcriptome calculated from genome TPM values (0.04-1.00) for each stage from each experiment. The samples were clustered by hierarchical clustering using Ward's clustering criterion ('ward.d2' setting in corrplot function [85]), and six clusters are highlighted in boxes. Six clusters were chosen as most optimal with the 'elbow method' [86], as implemented through the factoextra package function 'fviz_nbclust' using k-means clustering and within cluster sums of squares. The Experiment codes are listed in Table 1. (B) PCA plot of the genome TPM data coloured by the different stages and shapes determined by the originating experiment. PCA was conducted using the prcomp function in R. (C) PCA plot of the different samples by stage and experiment but using only *pir* gene TPM data, instead of the entire genome as used in (B).

Supplementary Figure 2

Transcription of four gametocyte-specific marker genes across the life cycle stages of *P. berghei* in the different experiments. The genes included are *p28* (*PBANKA_0514900*) and *nek4* (*PBANKA_0616700*), markers for female gametocytes and ookinetes; *mapk2* (*PBANKA_0933700*), marker for male gametocytes [66]; *hap2* (*PBANKA_1212600*), gamete fusion protein [67]. Bar height corresponds to median TPM, with error bars showing the range of TPM values across replicates. Colours correspond to different experiments.

Supplementary Figure 3

Representative Giemsa stained smears of *P. chabaudi* infected iRBCs throughout the 24 hr developmental cycle. Scale bar indicates 10 micrometres.

Supplementary Figure 4

A: Transcriptional deconvolution of each sample from every time point using scRNAseq data [14]. **B:** PCA plot of the genome TPM data coloured by the different timepoints. PCA was conducted using the prcomp function in R.

Supplementary Figure 5

Bar chart of the transcription of the ancestral *pir* gene (PCHAS_0101200) across the *P. c. chabaudi* AS asexual blood cycle. Each point represents one replicate and bars show the median TPM.

Supplementary Table 1.

The relationship between *P. chabaudi* RNA-seq samples used in this study and their entries in the ENA is described.

Figures

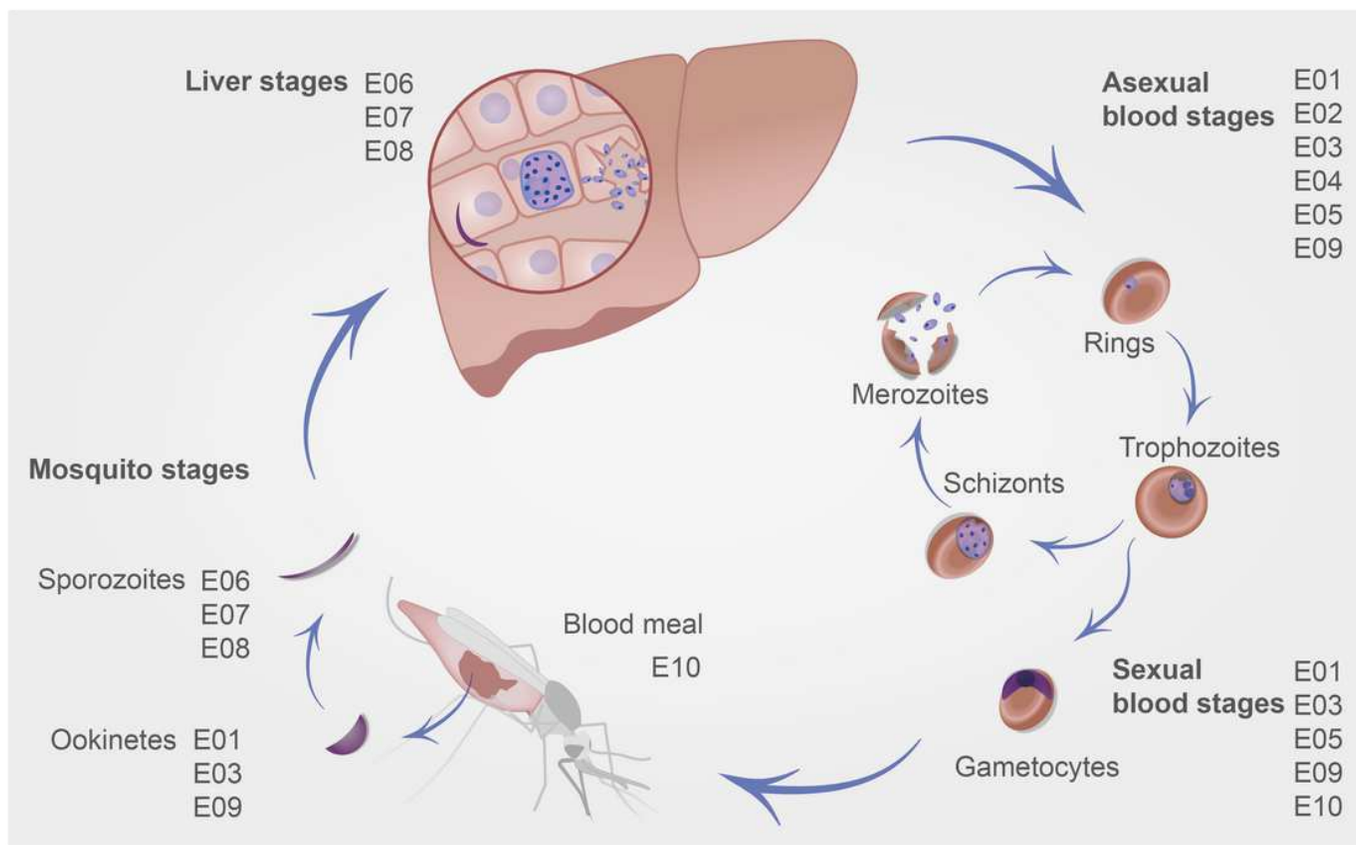


Figure 1

Systematic study of *P. berghei* RNAseq datasets that covers its entire life cycle and demonstrates the dynamic expression of *pir* genes throughout. Schematic of the published experiments used in this study and the *P. berghei* specific developmental stages during which samples were collected. The experiment codes (E01 etc.) correspond to the references in Table 1.

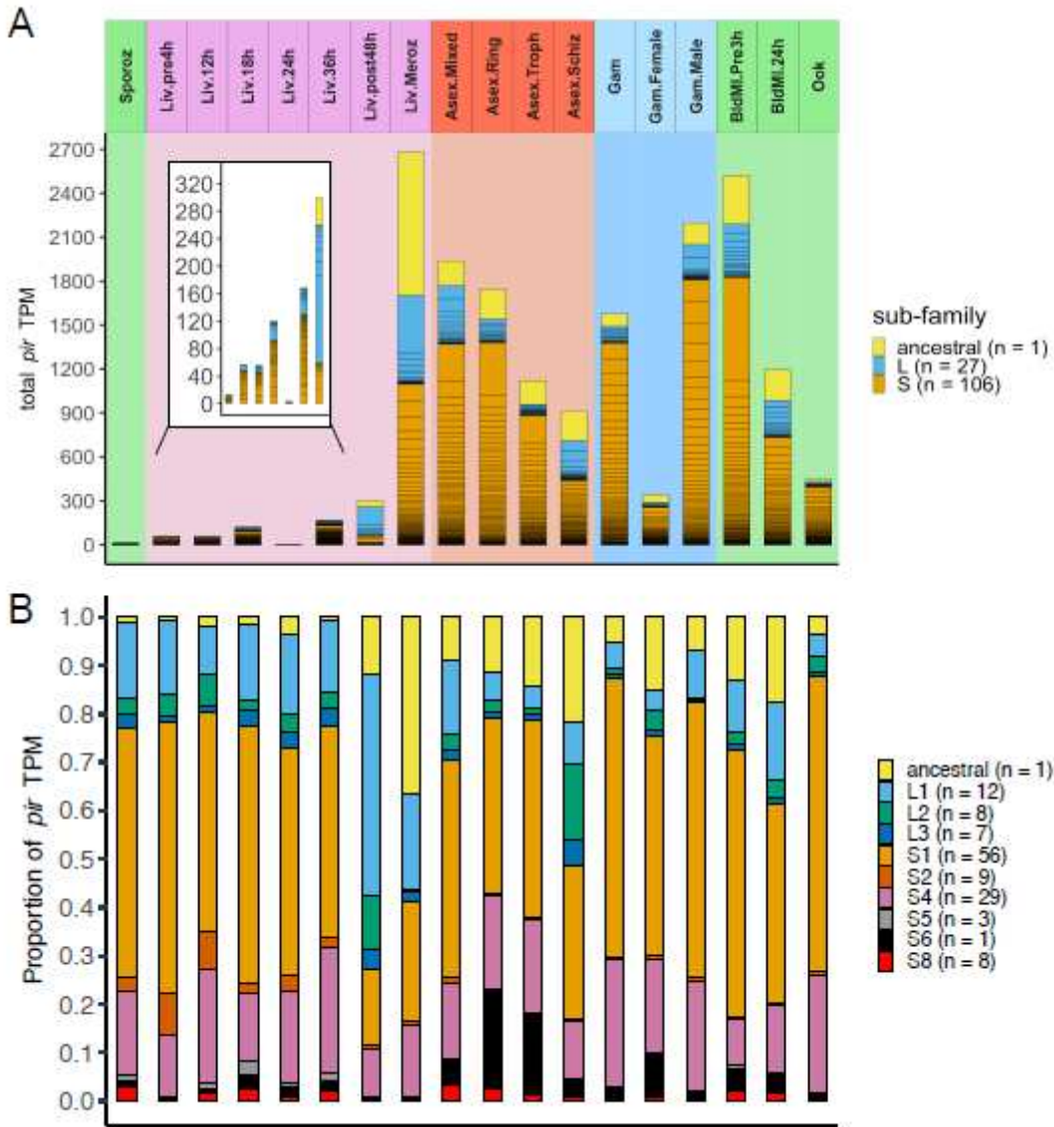


Figure 2

Expression of *pir* genes throughout *P. berghei* life cycle. A: Bar chart of total *pir* gene TPM (y-axis), calculated from the mean expression across all experimental samples of each life cycle stage (x-axis). The colours of the bar chart denote the classification; ancestral (yellow), Long (L) (blue) and Short (S) (orange), with the boxes of the stacks denoting expression of the individual genes. The background and strip name colours correspond to groups of life cycle stages, including mosquito (green), liver (pink), asexual blood (red), and gametocyte (blue) stages. The genes in each subfamily are ordered by expression levels. Inset of figure shows an enlargement of the sporozoite and liver stages. B: Bar chart of the proportion of total *pir* gene TPM (y-axis), contributed by the ancestral *pir* (yellow) and family subdivisions of the Short/Long *pir*s, for each stage (x-axis). Stages are separated as in (A). Legend shows the number of genes that are members of each grouping, which are collated together for each box in the stacked bars. The ordering is by family sub-division, from ancestral, to Long groups (1-3) and finally Short groups (1-2, 4-6, 8).

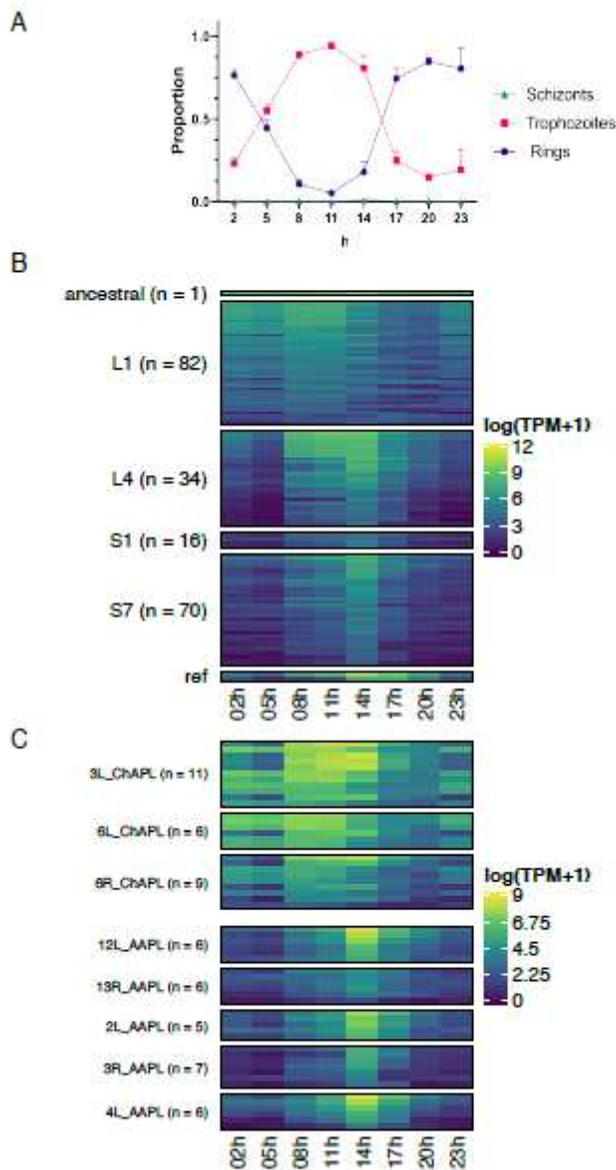


Figure 3

Intraerythrocytic development and *pir* gene expression across the 24 hour asexual blood cycle of *P. c. chabaudi* AS parasites transmitted by RMT. A: Microscopic counts of ring, trophozoite and schizont stage parasites in Giemsa-stained blood smears, as a proportion of the total parasitaemia (n=3). Graphs show mean (+/-SD). Value 1 would indicate that all parasites counted, are of the specific developmental stage. B: Heatmap of *pir* gene expression, $\log(\text{TPM}+1)$, at each timepoint. Genes are grouped according to subfamily (L1, L4, S1 and S7) and ordered by mean expression across the timepoints within each sub-family, with the ancestral *pir* at the very top. Three blood stage-specific genes were included for comparison 'ref': MSP1 (highly transcribed in schizonts/merozoites), HSP70 (highly transcribed in rings/trophozoites) and AMA1 (highly transcribed in schizonts/merozoites). Genes expressed under 1 TPM in every sample were removed. C: Heatmap of transcription of the *pir* genes assigned to ChAPL or AAPL loci [20][20], $\log(\text{TPM}+1)$, at each timepoint, grouped according to locus.

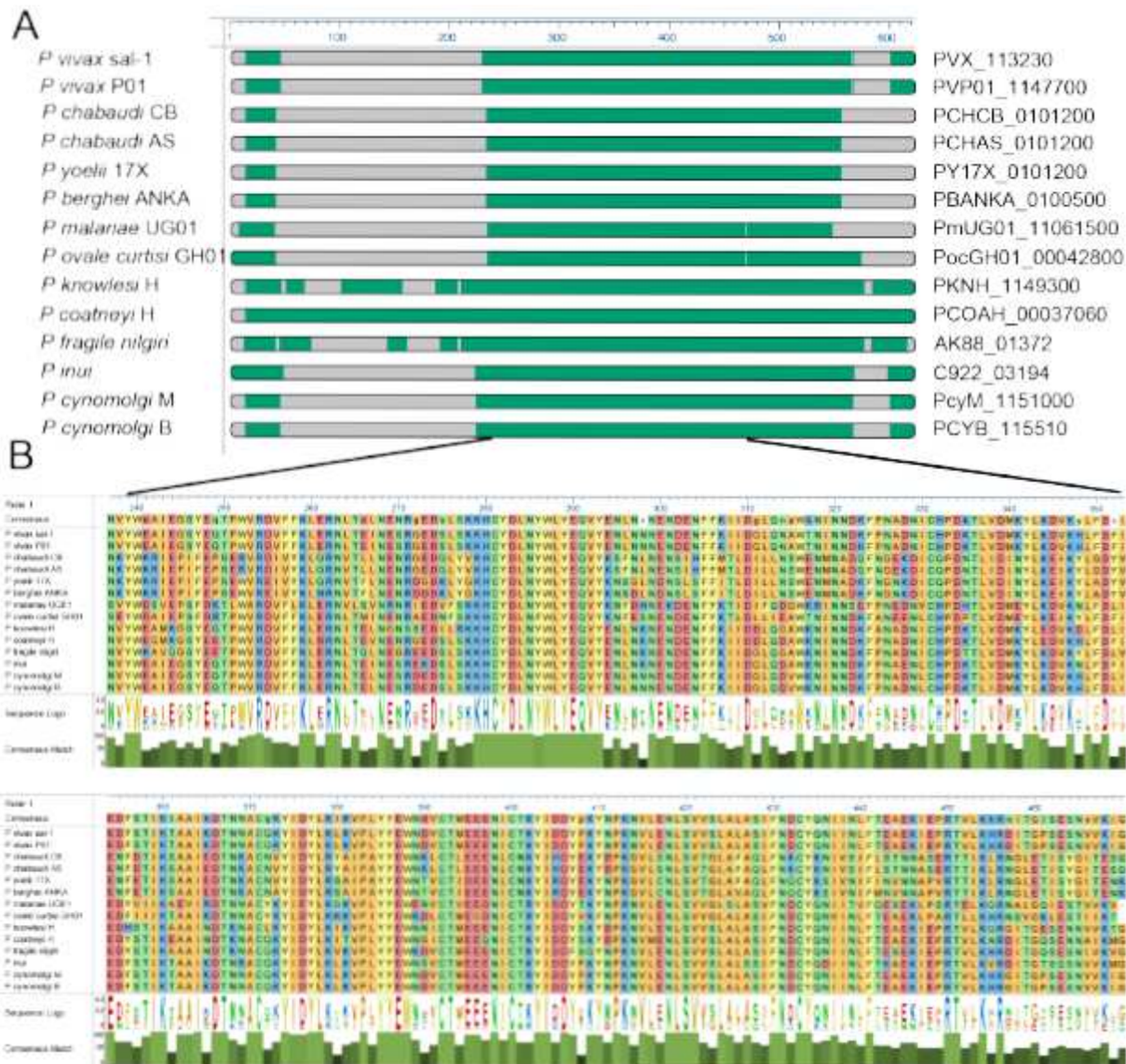


Figure 4

Conservation and gene expression of the ancestral *pir* across the *Plasmodium* genus. A: Sequence alignment of 15 ancestral *pir* orthologs (i.d.verified 20210527) identified in different *Plasmodium* spp. indicating regions and sequence of high conservation. B: Expanded region of the most highly conserved section of 234 aa, illustrated with a sequence logo and percentage match of the consensus.

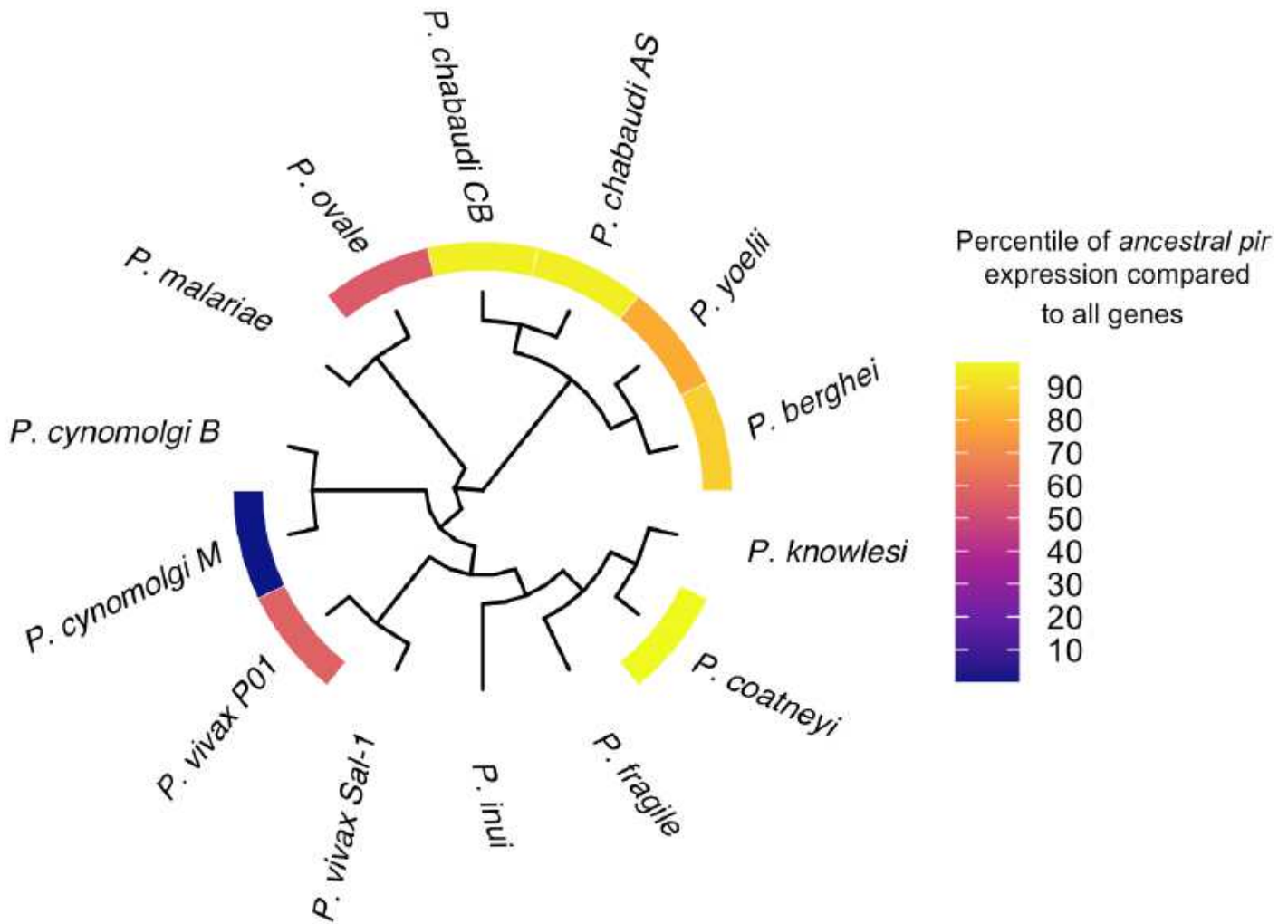


Figure 5

Phylogenetic analysis of the ancestral *pir* gene. Cladogram of the 15 orthologs, with heatmap of expression of the gene relative to the rest of the genome, where data is available (from multiple transcriptomic datasets as RPKM or TPM – see table 2). With the exception of *P. cynomolgi* strain M the ancestral ortholog is expressed within at least the top 50% of genes and usually in the top 30% of genes.

Supplementary Files

This is a list of supplementary files associated with this preprint. Click to download.

- [Table1.pdf](#)
- [Supplementaryalignment.pdf](#)
- [SupplementaryInformation1bergheiCounts.xlsx](#)
- [SupplementaryInformation2.xlsx](#)
- [SupplementaryInformation3.xlsx](#)
- [SupplementaryInformation4.xlsx](#)

- [SupplementaryInformation5.xlsx](#)
- [SupplementaryInformation6.xlsx](#)
- [SupplementaryTable1.xlsx](#)
- [suppinfoDescription.xlsx](#)
- [Table2.pdf](#)