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Antibody-independent mechanisms regulate the establishment of chronic *Plasmodium* infection

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Malaria is caused by parasites of the genus *Plasmodium*. All human-infecting *Plasmodium* species can establish long-lasting chronic infections\(^1-^5\), creating an infectious reservoir to sustain transmission\(^1,^6\). It is widely accepted that maintenance of chronic infection involves evasion of adaptive immunity by antigenic variation\(^7\). However, genes involved in this process have been identified in only two of five human-infecting species: *P. falciparum* and *P. knowlesi*. Furthermore, little is understood about the early events in establishment of chronic infection in these species. Using a rodent model we demonstrate that only a minority of parasites from among the infecting population, expressing one of several clusters of virulence-associated *pir* genes, establishes a chronic infection. This process occurs in different species of parasite and in different hosts. Establishment of chronicity is independent of adaptive immunity and therefore different from the mechanism proposed for maintainance of chronic *P. falciparum* infections\(^7-^9\). Furthermore, we show that the proportions of parasites expressing different types of *pir* genes regulate the time taken to establish a chronic infection. Since *pir* genes are common to most, if not all, species of *Plasmodium*\(^10\), this process may be a common way of regulating the establishment of chronic infections.
Long-lasting *Plasmodium falciparum* infections are thought to be maintained by
cytoadherence to avoid splenic clearance and antigenic variation of the adherent proteins
to avoid clearance by antibodies. We wanted to know how chronic infections are
established by malaria parasites lacking genes known to be involved in these processes
(var and sicavar). We used, as an example, the rodent malaria parasite *Plasmodium*
*chabaudi chabaudi* AS that gives rise to chronic infections in laboratory mice (Figure
1A). The acute infection is defined by a peak of parasitaemia (5-10% iRBC)
approximately ten days after mosquito bite, and clearance of the majority of parasites by
day fifteen. A chronic infection is then established, giving rise to several episodes of
patent parasitemia (>0.01%) for up to eighty days. Comparison of the transcriptomes of
parasite populations from the acute and chronic phases of infection showed that among
the multigene families identified in the *P. chabaudi* AS genome, the *pir* family is the
most differentially expressed (Figure 1A, Supplementary Tables 1 and 2). This multigene
family is present in most *Plasmodium* genomes\(^{11}\) and has been associated with
virulence\(^{12}\). More than half of the *pir* genes were differentially expressed between acute
and chronic phases. The majority of these genes had significantly higher expression in the
acute phase of infection, whereas only a few had higher expression in the chronic phase.

A *Plasmodium*-specific antibody response is detectable from day seven of
infection\(^{13}\) and thus the change in *pir* expression might be the result of adaptive
immunity-dependent selection of parasites expressing certain *pir* genes. However, we
observed the same change in mice lacking B cells and antibodies (μMT; Figure 1B) or
lacking T cells able to promote an antibody response (TCRα/-; Figure 1C;
Supplementary Figure 1; Supplementary Table 1). Therefore the transcriptomic changes observed between acute and chronic phases are independent of adaptive immunity and distinct from the mechanism suggested to be involved in maintenance of chronic infection in *P. falciparum*\textsuperscript{9}.

In the genomes of most malaria parasites, *pir* genes are located in subtelomeric regions that are difficult to resolve based on the *de novo* assembly of short sequencing reads. Using Single Molecule Real-Time (SMRT) sequencing, we generated a new genome assembly for *P. chabaudi* AS comprising a complete set of fourteen chromosomes with no gaps (Supplementary Table 3). We observed that some subtelomeres had similar sets of *pir* genes arranged in clusters (Supplementary Figures 2 and 3). In recent work, *pir* genes were classified into several short and long forms (S1-7 and L1-4, respectively)\textsuperscript{14}. We found eight clusters with common structures: three were enriched for the L1 subfamily and five were enriched for the S7 subfamily (Figure 2; Supplementary Table 4). Interestingly, we found that the majority of *pir* expression in our samples came from this small number of clusters, and that genes within clusters were co-expressed during infection (Supplementary Figure 4). While the S7-rich clusters were highly expressed during the acute phase, the chronic phase was dominated by expression of L1-rich loci (Figure 2C; Supplementary Table 5). This pattern was replicated in the αβ T-cell receptor- and B-cell knockout mice (Figure 2C) reinforcing our earlier conclusion that changes in *pir* gene expression are independent of selection by the adaptive immune response. Individual L1- or S7-rich loci are hereafter termed *Chronicity-Associated Pir Locus* (ChAPL), and *Acute-Associated Pir Locus* (AAPL), respectively. We identified an intergenic motif, commonly occurring upstream of most...
ChAPL genes, that might serve as a promoter sequence (Supplementary Figure 5). However, as only a subgroup of ChAPL genes are expressed in each mouse (Figure 2C), it is likely that epigenetic mechanisms regulate access of transcription factors to this motif, as shown for subtelomeric gene families in *P. falciparum*\textsuperscript{15}. Of the three ChAPLs, we observed a preference for clusters on chromosome 6.

We recently showed that increased *P. chabaudi* virulence during serial blood passage (SBP) is associated with a reduced repertoire of *pir* gene transcripts compared with parasites transmitted by mosquito (MT)\textsuperscript{12}. The gene most highly upregulated in SBP parasites was a *pir* from the L1 subfamily (PCHAS\_1100300). Although not found in a ChAPL, this gene is from the same subfamily that is most common in these loci. We therefore hypothesized that chronic phase parasites expressing ChAPLs would have a similar phenotype. To test this, we injected acute and chronic phase parasites into naïve mice. Chronic parasites gave rise to significantly greater parasitaemias and pathology than parasites derived from the acute phase of infection (Figure 3A; Supplementary Figure 6A). This suggests that parasite populations derived from the chronic phase expand better in the face of the response of a naïve mouse than parasites derived from the acute phase. However, as chronic parasites only reached high parasitaemias when injected into naïve mice, our data also indicate that they are better controlled when the host has previously experienced an acute infection. This pattern was also observed when chronic parasites were collected from B-cell knockout mice (Supplementary Figure 6B). Chronic infections therefore involve the emergence of phenotypically distinct parasites expressing ChAPLs, independently of adaptive immunity.
Thus we have established that there are transcriptional and phenotypic changes observed during chronic *Plasmodium* infections and that they do not stem from classical antigenic variation. We then sought to determine whether these were due to a transcriptional switch at the population level, or to a selection of pre-existing parasites from the acute phase. We isolated single parasites from the acute and chronic phases of infection, initiated clonal infections in C57Bl/6 mice and performed RNA sequencing analyses of these cloned populations of parasites (Figure 3B). We found that one out of ten acute-phase clones was more virulent than the population as a whole (Figure 3B; Supplementary Figure 7). While avirulent clones expressed AAPLs, the virulent clone expressed a ChAPL, as predicted by our finding that ChAPL-expressing parasites are more virulent in naïve mice (Figure 3C; Supplementary Figure 8; Supplementary Table 6). As expected, the clones from the chronic phase were all virulent and expressed ChAPLs (Figures 3B & 3C, Supplementary Figures 8 & 9). This suggests that the establishment of chronic infection involves selection of a small proportion of parasites expressing ChAPLs from a clonally variant population largely expressing AAPLs.

On the basis of these observations, we hypothesized that the proportion of virulent parasites already in the acute phase might affect the success of establishing a chronic infection. In our initial transcriptome experiment, we found that high expression of ChAPLs during the acute phase was associated with a more rapid recrudescence (Supplementary Table 7). We then tested whether chronic-like parasites, expressing L1 *pir* genes, outgrew acute parasites largely expressing AAPLs. We used SBP parasites as a proxy for chronic parasites as they express the L1 *pir* PCHAS_1100300, and have a similar phenotype to ChAPL-expressing chronic parasites. We performed mixed
infections in C57BL/6 mice with different ratios of SBP and acute MT (expressing AAPLs) parasites tagged with different fluorescent markers (Figure 4, Supplementary Figure 10 A-C; Supplementary Table 8). SBP parasites rapidly outcompeted MT parasites (Figure 4; Supplementary Figure 10D), confirming that the more virulent, chronic-like parasites had a selective advantage, as has been shown previously for different parasite isolates. The selection of a minority of iRBCs, expressing virulence-associated pir genes (Figures 3B & 3C), can therefore explain the change in expression from S- to L-pir genes observed in the parasite population between the acute and chronic phases of infection (Figures 1 & 2). As we had hypothesized, higher doses of L1-expressing parasites resulted in an earlier recrudescence confirming that these parasites have a better capacity to establish a chronic infection (Figure 4, Supplementary Figure 11).

When a similar infection mixing SBP and MT parasites was carried out in RAG1-/- mice (lacking both T and B cells) and RAG-/- mice treated with chlodronate liposomes to deplete macrophages (Supplementary Figure 12A), virulent parasites similarly rapidly outgrew AAPL-expressing parasites. Thus, this early selection takes place even in the absence of B cells, T cells, and macrophages indicating that other mechanisms are operating (Supplementary Figure 12B).

Unexpectedly, when mice were infected with $10^4$ virulent parasites plus $10^5$ avirulent parasites, the acute parasitaemia and pathology were lower than in mice infected with $10^4$ virulent parasites alone (Figure 4B, Supplementary Figure 11). This suggests that AAPL-expressing parasites act to lower the virulence of parasites expressing largely virulence-associated L1 pirs and delay the onset of chronicity.
Therefore, the initial composition of parasites expressing different pir genes regulates parasite recrudescence.

To determine whether our findings apply more generally, beyond *P. chabaudi* AS infections of laboratory mice, we established chronic infections with alternative host/parasite combinations. We used the more virulent *P. chabaudi* CB strain in C57BL/6 mice and *P. chabaudi* AS in the African thicket rat *Grammomys surdaster*. This is a natural host of rodent *Plasmodium* (Supplementary Figures 13A & 13B). Reduced AAPL and increased ChAPL gene expression during chronic *P. chabaudi* infection were observed in both cases, suggesting that our findings are a general feature of *P. chabaudi* chronic infections irrespective of strain of parasite or host (Supplementary Figures 13C & 13D). We then investigated chronic *P. berghei* infections in Brown Norway rats (Supplementary Figure 14A). The subtelomeric regions and pir repertoire of *P. berghei* are quite different from those of *P. chabaudi*\(^4\). Loci resembling the AAPLs and ChAPLs of *P. chabaudi* are absent. Nevertheless, we observed a general decrease in pir expression during the chronic phase with an increase in expression of L2 pir genes (Supplementary Table 9; Supplementary Figure 14B). The L2 pir genes of *P. berghei* are similar to the *P. chabaudi* L1 genes we found in ChAPLs. They are present in recently copied loci (Supplementary Figure 14C), contain long variable central domains and have a longer N-terminal domain that may contain a second transmembrane helix (Supplementary Figure 15). This suggests our findings are a general feature of malaria parasites infecting rodents.

Our results show that mosquito transmission of *P. chabaudi* gives rise to a clonally variant population of parasites, the majority expressing AAPLs and a minority ChAPLs. During the acute phase of infection, AAPL-expressing parasites are controlled
by the host response, while ChAPL-expressing parasites survive and establish a chronic infection. The more ChAPL-expressing parasites that are present early in infection, the more rapidly the chronic infection is established.

While it is thought that maintenance of chronic *P. falciparum* infection depends on the differential expression of *var* genes, how similar infections are established and maintained by *P. vivax*, *P. malariae* and *P. ovale*, that lack these variant antigens, is unknown. Here, we show that the establishment of chronic *Plasmodium* infection in rodent models is reproducibly associated with phenotypically distinct parasites, expressing particular *pir* gene clusters. Importantly, contrary to what is understood about maintenance of chronic infection in *P. falciparum*, this change in the parasite population is independent of adaptive immunity. Antigenic variation may still operate in maintaining chronicity, but our results clearly show that other mechanisms allow for the establishment of chronic infection in parasites that infect rodents.

Our finding that the initial population of parasites comprises individuals, each expressing different *pir* genes, suggests that the chronic infection is established by a small number of parasites which evade the early immune response. The selective advantage during infection of ChAPL-expressing over AAPL-expressing parasites could be of multiple origins. Although our previous results showed that growth rates of avirulent MT parasites and virulent SBP parasite were similar in RAG1-/- mice, red cell invasion efficiency might still play a role in selection of virulent clones for establishment of chronic infection. We are also investigating rosette formation, red cell deformability, sequestration, and the capacity to escape a macrophage-independent
innate immune response as possibilities. It has been shown that the expression of subsets of *P. falciparum* rif genes (which may be relatives of pir genes) and *P. vivax* pir genes are associated with cytoadherence of iRBCs to human endothelial cells and red blood cells, two processes implicated in parasite survival and pathogenicity. Furthermore, it has previously been suggested that the same molecules are involved in cytoadherence and immune evasion in *P. chabaudi*. Differences in cytoadhesive properties could therefore promote the expansion of more virulent parasites. Characterizing this process and determining the function of pir genes in vivo may identify mechanisms that universally regulate immune evasion by *Plasmodium* and lead to novel approaches to reduce their survival and transmission.
Methods

Mouse and parasite lines used

All experiments were performed in accordance with the UK Animals (Scientific Procedures) Act 1986 (PPL 80/2538) and the guidelines provided by the National Institutes of Health (NIH) Animal Care and Use Committee (ACUC) and approved by the Francis Crick Institute Ethical Committee and by the Institutional Animal Care and Use Committee (IACUC). T cell receptor-α chain knockout (TCRα-/-)31, mice homozygous for a targeted mutation of the transmembrane exon of the IgM μ chain, μMT32, and V(D)J recombination activation gene RAG-1 knockout (RAG-1-/-)33 on a C57Bl/6 background, and C57 Bl/6 Wildtype (WT) mice were obtained from the specific-pathogen free (SPF) unit and subsequently conventionally housed with autoclaved cages, bedding and food at the BRF, of the Francis Crick Institute. The Grammomys surdaster rats used in this study were captured in the wild at Fungurume and Lumata in the Katanga province, south of DR Congo, were bred and maintained in a conventional unit in sterile cages, with autoclaved water and bedding at NIH/NIAID/LMIV 34. Inbred Brown Norway rats were obtained from Charles River Laboratories and housed at the Animal Holding Unit of the Wellcome Trust Sanger Institute. Experiments were performed with 6-8 week old female mice and Grammomys surdaster rats housed under reverse light conditions (light 19:00-07:00, dark 07:00-19:00) and 6-week old female Brown Norway rats housed under normal light conditions, at 20-22°C. Measurements of clinical pathology were taken at 11:00. Core body temperature was measured with a rectal thermometer and erythrocyte density was determined with a VetScan HMII haematology system (Abaxis). Changes in body
temperature, body weight and erythrocyte density were calculated relative to a baseline measurement performed 1 day before infection. An infection was considered virulent when it induced a statistically significant decrease in the mouse temperature, weight and/or red blood cells compared with uninfected control mice.

Cloned lines of *Plasmodium chabaudi chabaudi* AS and CB parasites were used. To initiate infection, mice were either injected intraperitoneally (IP) or intravenously (IV) with 1, $10^4$ or $10^5$ infected erythrocytes or submitted to the bite of 20 infected mosquitoes as previously described$^{12}$. *Grammomys surdaster* rats were infected by I.V. injection of 2500 sporozoites of *Plasmodium chabaudi chabaudi* AS. Brown Norway rats were infected by I.V. injection of 7200 sporozoites of *Plasmodium berghei* ANKA. Parasitaemia was monitored by Giemsa-stained thin blood films on blinded samples. The limit of detection for patent parasitaemia was 0.001% infected erythrocytes.

**Chronic infections in wild type and genetically altered rodents**

Out of 10 wild-type C57Bl/6 mice, 10 μMT mice and 10 TCRα-/- mice infected with *P. chabaudi* AS via mosquito bite, 9 WT C57Bl/6 mice, 5 μMT mice and 3 TCRα-/- mice were randomly selected for transcriptomic analysis. 3 C57Bl/6 mice infected with *P. chabaudi* CB, 3 *Grammomys surdaster* rats infected with *P. chabaudi* AS and 3 Brown Norway rats infected with *P. berghei* ANKA (all via mosquito bites or injection of sporozoites) were also used for transcriptomic analysis. Our aim was to use a minimum of three biological replicates per condition for all experiments, as this is considered to be sufficient for general surveys of differential expression$^{35}$. Blood was extracted from the same rodents at two time points, therefore analysing the changes occurring over time in
the same parasite populations. During the acute phase of infection, 100μL of blood was
collected from the tail after the completion of seven cycles of schizogony. During the
chronic phase, exsanguination was performed between 30 and 40 days post-infection in
μMT mice, at 30 days post-infection in TCRα-/− mice, at day 15 post-infection in Brown
Norway rats or when the parasitaemia reached 0.1% between 27 and 40 days post-
infection in wild type C57Bl/6 mice and G. surdaster rats. After data collection, the acute
sample from wild-type mouse 454 and both acute and chronic samples from μMT mouse
433 were excluded, as they were outliers in a multi-dimensional scaling analysis of the
gene expression data.

**Phenotypic analysis of acute and chronic parasites**

Donor mice were infected via mosquito bites. Parasites from donor mice (n = 3) were
blood-passaged into recipient mice at 1, 3, 4 and 6 weeks post-mosquito transmission.
Recipient mice (n=5 per donor mouse, the minimum number required for an 80% chance
of significant differences (p<=0.05) in parasitemia, weight and erythrocyte counts) were
infected by IP injection of $10^5$ infected erythrocytes. Parasitaemia and signs of pathology
were monitored in recipient mice on blinded samples.

**Cloning of single parasites from acute and chronic phases**

One mouse was infected via mosquito bite. Infected blood was then collected from the
tail at two time points: 1 week post-infection (acute phase) and six weeks post-infection
(chronic phase). Both times, parasites were cloned by dilution and single parasites were
injected IV in wild type mice. After expansion, each cloned line was blood passaged into
eight experimental mice by IP injection of $10^5$ infected erythrocytes. Five mice were used to monitor parasitaemia and signs of pathology and three mice were sacrificed and exsanguinated after the completion of seven cycles of schizogony for RNA extraction.

**RNA extraction**

During all RNA extractions experiments, blood-stage parasites were isolated at 11:00, when >90% of the parasites were at the late trophozoite stage of development (Table S8). Parasite RNA was isolated as previously described. Briefly, whole blood was depleted of leukocytes by filtration (Plasmodipur, EuroProxima) and from globin RNA and red cell debris by saponin lysis (Sigma) and centrifugation. Purified parasites were resuspended in TRIzol, snap-frozen on dry ice and kept at -80°C until use. RNA was then extracted, resuspended in water and its quantity and quality were determined on an Agilent 2100 Bioanalyzer RNA 6000 Nano or Pico chip.

**RNA-seq library prep and sequencing**

The majority of RNA was used to make 200-450bp fragment libraries using Illumina's TruSeq RNA Sample Prep v2 kit, with 10 cycles of PCR amplification using KAPA Hifi Polymerase rather than the kit-supplied Illumina PCR Polymerase. Two samples required 14 cycles of PCR (T1_46-4 and T1_47-4). Mosquito-transmitted *P. chabaudi* CB samples (MTCB) were subjected to 15 cycles of PCR. The libraries were sequenced using an Illumina HiSeq 2000 v3 with 100bp paired-end reads (samples beginning ‘T’), an Illumina HiSeq 4000 with 75bp paired-end reads (MTCB samples) or an Illumina HiSeq 2500 with 75bp paired-end reads for the rest.
RNA from cloned parasites was used to make 100-300bp fragment libraries produced using Illumina’s TruSeq Stranded mRNA Sample Prep Kit with 10 cycles of PCR amplification using KAPA Hifi Polymerase. These libraries were sequenced using an Illumina HiSeq 2500 with 75bp paired-end reads. A description of the libraries can be found in Supplementary Table 10.

Analysis of gene expression

Tag sequences were removed from sequenced reads. Where individual samples were run over multiple lanes, the reads from these were merged before mapping. Reads were then mapped against spliced gene sequences (exons, but not UTRs) from the v3 *P. c. chabaudi* AS (this work) or the v3 *P. berghei* ANKA (Fougère et al., under review) reference genomes using Bowtie2 v2.1.0\(^{-36}\) (-a -X 800 –x). Read counts per transcript were estimated using eXpress v1.3.0\(^{-37}\), with default parameters. Genes with an effective length cutoff below 10 in any sample were removed. Summing over transcripts generated read counts per gene.

Differential expression analysis was performed using edgeR v3.8.6\(^{-38}\) on genes with \(\geq 3\) counts per million. The Fisher’s exact test was used with cutoffs FDR < 0.01 and fold change \(\geq 2\), except for the *P.berghei* analysis where FDR < 0.1 and fold change \(\geq 1\) was used. Functional categories of genes were identified by orthology using GeneDB \(^{39}\) from several different *P. falciparum* datasets: invasion genes\(^{40}\), sexual genes\(^{41}\) and subtelomeric (by manual inspection of chromosomes).
Gene expression clustering was used to identify which genes within ChAPLs and AAPLs were co-expressed. This was done by generating a toroidal 5x5 Self-Organising Map (SOM) using the Kohonen package in R\(^4\). FPKM values for each gene were logged (base 2) and mean-normalised per gene. The SOM was trained in 100 iterations.

**Sequencing and assembly of *P. c. chabaudi* AS v3 genome**

High molecular weight DNA was prepared as follows. Heparinised blood from two mice infected with *P. chabaudi* AS (blood passaged parasites, days seven & eight post-infection) was pooled, diluted 1:3 in PBS, and passed through a Plasmodipur (Euro-Diagnostica) filter. Erythrocyte membranes were removed by saponin lysis (0.15% in ice cold PBS). Parasitised erythrocytes were recovered by centrifugation (2000g 10 min 4 °C) and washed twice in ice cold PBS. The cell pellet was taken up in 50mM Tris HCl pH 7.5, 50mM EDTA pH8.0, 100mM NaCl, 0.5% SDS and digested with RNase A (1mg/ml, Life Technologies) for 30 min at 37°C. Proteinase K (Roche) was added to a final concentration of 1mg/ml, incubated at 45°C overnight, followed by extraction with phenol chloroform and ethanol precipitation.

For preparation of long-read sequencing libraries, 5μg of *P. c. chabaudi* AS genomic DNA was sheared to 20-25kb by passing through a 25mm blunt ended needle. SMRT bell template libraries were generated using the Pacific Biosciences issued protocol (20 kb Template Preparation Using BluePippin™ Size-Selection System). After 7kb-20kb size-selection using the BluePippin™ Size-Selection System (Sage Science, Beverly, MA) the library was sequenced using P6 polymerase and chemistry version 4 (P6C4) on 5 single-molecule real-time (SMRT) cells, each with a 240 minutes movie.
The five SMRT cells were processed using a Pacific Biosciences RSII. Reads were filtered using SMRT portal v2.2 with default parameters (minimum subread length 50, minimum polymerase read quality 75, minimum polymerase read length 50). The yield was 258214 reads totaling 1.51Gb, with read N50 9003bp and read quality 0.833. The reads were assembled using HGAP v2.2.0 to identify sequences relating to nuclear, mitochondrial and apicoplast genomes. Four unitigs were found to be mouse contamination. Fourteen unitigs represented the fourteen chromosomes from the v2 assembly including most of the unplaced contigs from v2 and some additional new sequence. All fourteen had telomeric repeats at both ends. The only major rearrangement between the assemblies was between the ends of chromosomes 6 and 13. We mapped Quiver-corrected PacBio reads against v2 and v3 assemblies and confirmed that the v3 assembly was correct. However, we noticed base errors and in particular many indels in the v3 assembly due to the relatively high error rate of PacBio sequencing. The v2 assembly had had the benefit of Sanger and Illumina technologies such that at the base level it was more accurate than the PacBio assembly. Therefore we generated 2x250bp paired-end pseudoreads from the v2 assembly with a fragment size of 600bp to a depth of 50x and used these to correct the v3 assembly using iCORN2\(^4\). We transferred gene models from the v2 assembly to the corrected v3 assembly using RATT\(^4\). We then trained Augustus v2.5.5\(^5\) using the v2 gene models with default parameters and predicted a new set of models for the corrected v3 assembly. We kept only those, which did not overlap with the RATT-transferred gene
models. We then manually assessed the transferred models, annotated the new Augustus models and renamed the locus tags. In general we added a zero to the end of the v2 locus tags, but where a gene did not exist in v2 or had moved we added an odd number at the end. We maintained numerical ordering of locus tags across each chromosome so that they reflect relative chromosomal location.

Sequence analysis of pir genes

*Pir* gene subclade assignments were initially made using Hidden Markov Models (HMMs) built from the rodent malaria *pir* subclades published in Otto et al.\textsuperscript{14}. There were some discrepancies between the two sets, e.g. where the clade in the published tree did not agree with the top HMM hit. In particular we identified three members of the S3 subclade, which was not previously thought to be present in *P. chabaudi*. We used hmmsearch from HMMer i1.1rc\textsuperscript{36} with an E-value cutoff of 1e-10 and took the best hit to a subclade. We thus identified 207 *pir* genes which we could assign to a rodent malaria parasite (RMP) *pir* subclade, while a further five genes could not be classified into existing subfamilies (PCHAS_0401400, PCHAS_0602000 and PCHAS_1247000).

HMM assignments of L2 *pirs* in *P. berghei* were consistent with the original tree-based classification presented in Otto et al. (2014).

We used FIRE v1.1a\textsuperscript{47} to identify putative promoter sequences upstream of genes in the AAPLs and ChAPLs. Firstly we used RNA-seq data from the study to determine 5' untranslated regions (UTRs) of *pir* genes and then manually identified untranscribed regions upstream of each UTR, reaching until the proximal UTR of the next upstream gene. Where there was insufficient RNA-seq data or other ambiguity, we did not identify
a region. This was successful for 27/32 ChAPL genes and 22/28 AAPL genes. We then defined AAPL and ChAPL genes as two distinct expression clusters and ran FIRE with --exptype=discrete and --nodups=1 options.

Sequence alignments were performed using Muscle\textsuperscript{48} with default parameters and visualized using Jalview\textsuperscript{49}.

**Regression analysis of pir genes against parasitological parameters**

DESeq2 was used to determine whether variation in gene expression between parasite populations in different mice was correlated with parasitological parameters such as parasitaemia and time of recrudescence\textsuperscript{50}. In particular we found that the expression level during the acute phase of 152 genes was correlated with the time to recrudescence across nine mice (p-value < 0.01; fold change $\geq$ 2). We discretized the time in days at recrudescence as follows: mice 214 (32 days), 421 (31 days), 427 (30 days) were short, 2014 (34 days), 2114 (36 days), 452 (33 days) and 456 (34 days) were medium, 211 (43 days) and 212 (39 days) were long. We then used this single discrete factor in the DESeq2 design formula. Those genes where expression level during the acute phase was negatively correlated with time at recrudescence were, with a single exception, *pir* genes from 3L, 6L and 6R subtelomeres (Table S6). These are the three ChAPLs and another L1-rich locus (4R). Interestingly, the strongest effect was from an exported gene of unknown function (PCHAS\_1201300). The strongest positive effects were from *pir* genes in AAPLs.

**Mixed infection**
Transgenic lines of *Plasmodium chabaudi* AS expressing mCherry or mNeonGreen under the control of the constitutive promoter EF1α were generated by transfection with the plasmid *PcEFp230p_mCherry* and *PcEFp230p_mNeonGreen*, respectively. These plasmids targeted the silent 230p genomic locus, and were generated by replacement of the ssu rRNA locus of *pPc-mCHCAM*, described in \(^51\), with the *P. chabaudi* p230p locus (*PCHAS_03_v3 276,451-283,284*). For *PcEFp230p_mNeonGreen*, the mCherry tag was replaced with mNeonGreen\(^52\). Parasites were cycled ON and OFF acid water containing pyrimethamine to select for stably transfected parasites, and cloned by serial dilution. The insertion was verified by PCR amplification and Southern blot analysis and fluorochrome expression by Flow cytometry analysis (Figure S9). mNeonGreen-expressing parasites were then serially blood passaged (SBP) 10 times while mCherry-expressing parasites were mosquito transmitted (MT) into donor mice. Both parasites were then mixed at different ratios and \(10^5\) infected erythrocytes were injected i.p. into ten recipient mice per group. Competition between parasite lines as well as parasitaemia and pathology were then monitored in recipient mice. To determine the ratio of mNeonGreen- and mCherry-expressing parasites during infection, tail blood was collected and stained with Hoechst 33342 (New England Biolab). Samples were acquired on a BD LSRII flow cytometer and data were analysed with FlowJo software (TreeStar).

**Clodronate liposomes treatment**

B6.RAG-1/- mice were treated with 200uL i.v. of either saline (Sigma; negative control) or clodronate liposomes (clodronateliposomes.org). Flow cytometric analysis was carried out to assess the depletion of phagocytic cells. Single-cell suspensions of
splenocytes were prepared, , and cells enumerated 24h after in vivo treatment. After staining with Zombie Aqua (Biolegend) for live/dead discrimination, cells were stained with monoclonal antibodies using appropriate combinations of fluorochromes (CD11c-BV786, CD169-BV605, F4/80-BV421, Ly6G-PerCPCy5.5, MHCII-FITC, Ter119-PE-Cy7, CD68-PE, Ly6c-APC-Cy7, CD11b-AF647, all from Biolegend). The samples were acquired on a BD Fortessa/X20 (BD Biosciences) using Diva acquisition software (BD Biosciences). FlowJo (Tree Star) software was used to analyze the data.

To determine the role of T-, B- and phagocytic cells in the selection of virulent parasites, B6.RAG-1/- mice were treated with saline or clodronate liposomes as described above at days -1 and 4 after i.p. infection with 10^5 iRBCs containing a mix of Neon-Green-expressing SBP parasites and mCherry-expressing MT parasites (see "Mixed infections" section).

**Data availability**

RNA-seq datasets used in this study have been submitted to the European Nucleotide Archive (ENA) under study accession ERP017479. These datasets are described further in Supplementary Table 10. The Pacific Biosciences RSII genomic sequencing reads, used to generate the *Plasmodium chabaudi chabaudi* AS v3 genome sequence, have been submitted to the ENA under experiment accession ERX613966. The assembled genome sequence and annotation can be accessed from GeneDB (http://www.genedb.org/Homepage/Pchabaudi) and PlasmoDB (http://plasmodb.org/plasmo/). The authors declare that no competing interests exist.
References


Conteh, S. *et al.* Grammomys surdaster, the natural host for Plasmodium berghei parasites, as a model to study whole organism vaccines against malaria. *American Journal of Tropical Medicine and Hygiene* (in press).


Waterhouse, A. M., Procter, J. B., Martin, D. M., Clamp, M. & Barton, G. J.
Jalview Version 2--a multiple sequence alignment editor and analysis workbench.

Love, M. I., Huber, W. & Anders, S. Moderated estimation of fold change and
dispersion for RNA-seq data with DESeq2. Genome Biol 15, doi:ARTN 550

Spence, P. J. et al. Transformation of the rodent malaria parasite Plasmodium
Shaner, N. C. et al. A bright monomeric green fluorescent protein derived from
(2013).

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Author contributions
T.B., A.J.R., C.N., M.B. and J.L. (last author) designed the study. T.B. designed and performed the mouse experiments with the help of I.T., G.K., S.M. and P.S. A.J.R. designed the sequencing experiments and performed the bioinformatic analyses. M.S. coordinated sequencing experiments. U.B. manually annotated the *Plasmodium chabaudi* AS genome sequence. D.C. and J.L. (7th author) created transgenic parasites. P.E.D. and S.C performed thicket rat experiments, T.M, E.B. and O.B carried out the experiments in the Brown Norway rats. T.B., A.J.R., C.N., M.B. and J.L. (last author) wrote the manuscript.

**Competing financial Interests**

The authors declare no competing financial interests.

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Figure legends

Figure 1: Chronic infections modify the transcriptome of *P. chabaudi* independently of the adaptive immune response.

Parasitaemias over the course of infection in (A) 10 wild-type C57Bl/6 mice, (B) 10 B6.μMT-/- mice and (C) 10 TCRα-/- mice are shown on the left panels. Parasite mRNAs were collected from 9 wild-type C57Bl/6 mice, 5 B6.μMT-/- mice and 3 TCRα-/- mice selected randomly at the time points indicated by the red arrows. On the right panels, hot pie diagrams show expression levels (black and white inner circles) and fold changes (coloured outer circles) for genes expressed more highly (FDR <= 0.01, fold change >= 2) in the chronic and acute phases. Genes are classified into groups according to several categories including: subtelomeric genes (subtel.); genes associated with red blood cell invasion (invasion); genes associated with gametocytogenesis (sexual) and *pir* genes (highlighted in brown).

Figure 2: Chronic infections are characterized by the expression of distinctive clusters of *pir* genes in *P. chabaudi*.

(A) Chronic Associated *Pir* Loci (ChAPL), and (B) Acute Associated *Pir* Loci (AAPL) are shown in the context of the subtelomeric region from the telomeric repeats to the last *pir* or *fam-a* gene. Those genes co-expressed across mice over the course of infection are highlighted in orange. (C) Heatmap of *pir* gene expression at ChAPL and AAPL loci, showing mean-normalised RPKM values for 10 wild type C57Bl/6 mice, four B6.μMT-/- mice and three TCRα-/- mice infected by mosquito bites. Each column represents values obtained from an individual mouse. Samples were collected from the same mice during
the acute and chronic (chro.) phases of infection. On the side, numbers indicate chromosomes and letters indicate the location on that chromosome: L for left hand end, R for right hand end. ChAPLs and AAPLs are highlighted. For each gene, red represents its maximal expression, green its minimal expression. The absolute maximum and minimum values for each gene are shown in a separate heatmap.

Figure 3: Chronic infections select for virulent *P. chabaudi* parasites expressing distinctive clusters of *pir* genes.

Donor mice were infected by mosquito bites. (A) Parasites were subsequently passaged into naïve recipient mice after one, three, four or six weeks (n=15 mice per group). (B) Individual parasites were cloned during the acute and chronic phases of infection. After their expansion, clonal populations of parasites were used to infect recipient mice (n= 5 mice per clone). In (A) and (B), we show maximum parasitaemia, temperature change and weight change for recipient mice compared to mice directly infected by mosquito bites (MT; blue dotted line) or with serially blood passaged parasites (SBP; red dotted line). Each dot represents the mean ± standard error of the mean (SEM) of 5 individual mice. Each group has been compared to mice infected with 10⁵ MT parasites (*P<0,05; **P<0,01; ***P<0,001, two-sided Mann Whitney Test). (C) Heatmap of *pir* gene expression at ChAPLs and AAPLs in three acute and three chronic clones. The first three columns represent cloned parasite lines derived from from the acute phase (f, a, b), the next three columns represent cloned lines from the chronic phase (k, n, p) from 3 individual mice. On the side, chromosomes numbers are indicated as well as the location on that chromosome: L for left hand, R for right hand. ChAPLs and AAPLs are
highlighted. For each gene, red represents its maximal expression, green its minimal expression. The absolute maximum and minimum values for each gene are shown in a separate heatmap.

**Figure 4: Chronicity and virulence of infection are dictated by the initial composition of *P. chabaudi* population.**

Parasites tagged with mCherry (Fluo 1) were Mosquito-Transmitted (MT), and mNeonGreen (Fluo 2) tagged parasites were Serially Blood-Passaged (SBP). Different proportions of the two were used to infect recipient mice by i.p. injection of $10^5$ parasites ($n=10$ mice per group). As a control, a group of 10 recipient mice was infected by i.p. injection of $10^4$ SBP parasites. The mean percentage of parasites being SBP (as determined by flow cytometry) is indicated (upper panel). We show maximum parasitaemia, temperature change, weight change (middle panel) and time of recrudescence (lower panel) for each group compared to mice infected with 100% MT parasites (blue dotted line) or 100% SBP parasites (red dotted line). Each dot represents the mean ± standard error of the mean (SEM) for each group of 10 mice. Each group has been compared to mice infected with $10^5$ SBP parasites ($*P<0.05$; **$P<0.01$; ***$P<0.001$, two-sided Mann Whitney Test).
Acute parasitemia (%)

Acute temperature change (%)

Acute weight change (%)

1st day of recrudescence (days post-infection)