Evolutionary analysis of the most polymorphic gene family in *falciparum* malaria [version 1; peer review: 1 approved, 2 approved with reservations]

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**Abstract**

The *var* gene family of the human malaria parasite *Plasmodium falciparum* encode proteins that are crucial determinants of both pathogenesis and immune evasion and are highly polymorphic. Here we have assembled nearly complete *var* gene repertoires from 2398 field isolates and analysed a normalised set of 714 from across 12 countries. This therefore represents the first large scale attempt to catalogue the worldwide distribution of *var* gene sequences. We confirm the extreme polymorphism of this gene family but also demonstrate an unexpected level of sequence sharing both within and between continents. We show that this is likely due to both the remnants of selective sweeps as well as a worrying degree of recent gene flow across continents with implications for the spread of drug resistance. We also address the evolution of the *var* repertoire with respect to the ancestral genes within the *Laverania* and show that diversity generated by recombination is concentrated in a number of hotspots. An analysis of the subdomain structure indicates that some existing definitions may need to be revised. From the analysis of this data, we can now understand the way in which the family has evolved and how the diversity is continuously being generated. Finally, we demonstrate that because the genes are distributed across the genome, sequence sharing between genotypes acts as a useful population genetic marker.

**Keywords**

Plasmodium, var, evolution
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Introduction

*Plasmodium falciparum* is the most virulent of the human malaria parasites, at least in part because it can concentrate in the small vasculature of organs via its ability to adhere to endothelial cells leading to organ dysfunction. It has also developed the ability to maintain chronic infections in humans to maximise the probability of mosquito transmission. A single multi-gene family of ~60 var genes in this parasite is a major determinant of both of these important phenotypes. The var genes encode a family of proteins called *Plasmodium falciparum* Erythrocyte Membrane Protein 1 ( PfEMP1) that are expressed on the surface of infected red cells in a mutually exclusive fashion. On the red cell surface, PfEMP1 proteins mediate both adherence to endothelium and induce host protective antibodies. The latter are evaded through transcriptional switches among the var gene family such that new sequences are periodically expressed. Through their role in immune evasion they have evolved to be extremely polymorphic. The genes are located mostly in subtelomeric regions but are also located in a few chromosome-internal clusters. Analysis of fully sequenced genes has shown that var genes can be subdivided into three major classes ( UPS A, B and C), based on their upstream sequences and chromosomal position, and that they are composed of long 5' exons that encode multiple combinations of two types of extracellular domains, the Duffy Binding Like (DBL) and Cysteine Rich Interdomain Regions (CIDR) and a more conserved 3' exon encoding the intracellular region. The DBL and CIDR regions have been divided into subclasses based on sequence similarity. Most of the data from field isolates on var gene sequences has to date been collected by PCR of a small region of DBLα, eg: var pseudogenes that inevitably assembled as short contigs, for example, var genes containing DC8 and 13 are expressed in parasites causing severe malaria and bind to endothelial protein C receptor (EPCR). A more extensive analysis of full-length genes from the sequenced genomes of seven laboratory-adapted isolates revealed that all domain subclasses can be further divided into a number of subdomain types and that in some cases these can be grouped together into "domain cassettes". Some of these domain cassettes (DC) are associated with binding to specific receptors on endothelial cells; for instance, var genes containing DC8 and 13 are expressed in parasites causing severe malaria and bind to endothelial protein C receptor (EPCR). More recently, a combination of Illumina and long read Pacific Biosciences sequencing has been used to assemble the genomes, including the var genes from 12 Malian isolates and 15 field isolates. This has confirmed that the number and domain organisation of var genes in these samples is similar to that in the 3D7 reference. RNAseq of parasites isolated from severe and non-severe patients in Indonesia has also been used to assemble expressed var gene sequences.

The var gene family is restricted to the *Laverania* sub-genus of malaria parasites that infect great apes and from which *P. falciparum* evolved. However, the fact that the number, genomic location and domain architecture of these genes is very similar in *P. reichenowi* (a parasite of chimpanzees that diverged from the *P. falciparum* lineage around 200,000 years ago) suggests that the evolutionary fitness of the diversity and organisation of this family has been long established.

The central role of these genes in crucial areas of malaria biology has meant that they have been the subject of extensive research interest, but to date the paucity of full-length sequence information has precluded an in-depth global study. Recently, large scale sequencing has produced short-read data for thousands of *P. falciparum* isolates but to date the analysis of these data has been restricted to coding sequences within the central core genome and has not addressed the sub-telomeres or central var gene clusters. Here we have assembled almost complete var gene repertoires from MalariaGEN *P. falciparum* Community Project and Pf3k datasets and present data from 12 countries for which ~60 isolates were available from each. Using this normalised dataset, we performed a detailed analysis of var gene sequence-sharing across the world. Sharing across the dataset is much greater than had been expected and there were clear hotspots of recombination. Subdomains of DBL and CIDR sequences could be identified based on their sequence conservation, but we show that some current definitions are not robust. Many of the sequences that comprise extant var genes can also be traced back to their ancestors in great apes, and because var genes are distributed throughout the genome, we demonstrate that these genes are rich and informative population genetic markers for the genome as a whole.

Results

Using whole genome sequencing data from approximately 2,400 clinical isolates of *Plasmodium falciparum* produced by the Pf3k project (*Extended data: Table S1*), we developed a pipeline (Methods) that enabled us to assemble contigs representing putative individual var exons. Throughout this study, a var assembly refers to its large exon I sequence, rather than the whole gene that also includes a highly conserved and much smaller exon II sequence. The number of LARSFADIG motifs (the only motif shared by all DBLα domains that are present in exon I of all regular var genes) is a proxy for the number of var genes/genome. To create a normalised dataset, we included 714 samples containing between 45 and 90 LARSFADIG motifs to minimise the number of multiple infections present (Table 1). All genomes in our normalised set contained variable numbers of var pseudogenes that inevitably assembled as short fragments and did not contribute to the accurate number of intact genes assembled. By focussing on var genes > 3.5 kb, we observed approximately the expected number of var genes/ genome with a tendency for a higher number from African countries where higher transmission increases the probability of infection with multiple genotypes. This was confirmed in 20 by the presence of multiple genotypes of polymorphic single copy genes. To further validate the data, we used our previously assembled genome sequences of 12 isolates that had been adapted to culture to provide enough DNA for long-read PacBio sequencing. The PacBio assemblies were corrected with short-read sequencing produced using Illumina HiSeq 2500 and
Table 1. Summary of normalised dataset. var1CSA and var2CSA sequences and exon2 are excluded.

<table>
<thead>
<tr>
<th>Country of origin</th>
<th>Analysed isolates*</th>
<th>Average number of LARSFAGID motifs per genome</th>
<th>Average number of sequences &gt; 3kb per genome</th>
<th>Unique shared var genes per genome**</th>
</tr>
</thead>
<tbody>
<tr>
<td>The Gambia</td>
<td>60</td>
<td>63.6</td>
<td>62.1</td>
<td>20.6</td>
</tr>
<tr>
<td>Kenya</td>
<td>55</td>
<td>50.5</td>
<td>48</td>
<td>6.3</td>
</tr>
<tr>
<td>Thailand</td>
<td>60</td>
<td>55.1</td>
<td>54.5</td>
<td>33.2</td>
</tr>
<tr>
<td>Ghana</td>
<td>60</td>
<td>63.3</td>
<td>62.6</td>
<td>6.6</td>
</tr>
<tr>
<td>Cambodia</td>
<td>60</td>
<td>56.1</td>
<td>53.6</td>
<td>29.6</td>
</tr>
<tr>
<td>Mali</td>
<td>60</td>
<td>70.3</td>
<td>70</td>
<td>8.6</td>
</tr>
<tr>
<td>Senegal</td>
<td>60</td>
<td>51.2</td>
<td>46.8</td>
<td>8.6</td>
</tr>
<tr>
<td>Malawi</td>
<td>60</td>
<td>65.5</td>
<td>61.5</td>
<td>6.1</td>
</tr>
<tr>
<td>Guinea</td>
<td>60</td>
<td>59.4</td>
<td>62.6</td>
<td>8.1</td>
</tr>
<tr>
<td>Vietnam</td>
<td>59</td>
<td>57.2</td>
<td>50.3</td>
<td>21.0</td>
</tr>
<tr>
<td>Laos</td>
<td>60</td>
<td>58.9</td>
<td>58.5</td>
<td>27.1</td>
</tr>
<tr>
<td>Congo</td>
<td>60</td>
<td>63.2</td>
<td>60.3</td>
<td>9.2</td>
</tr>
</tbody>
</table>

* Genomes excluded due to contamination or not enough assembled sequences
** unique count of hit against normalised set, hit > 99% identity, > 3.5kb, 80% overlap

the var genes were subsequently manually curated. Comparison of our assembled Illumina and PacBio data did show one systematic error: in 1.5% of var exon I sequences, frame-shifts occurred within homo-polymer tracts that could not be corrected by iterative alignment and error-correction using iTORN. As a consequence, we estimated that our assemblies are 98.5% accurate, vastly exceeding the level required for our downstream analysis of high frequency events.

Polymorphism

The degree of polymorphism in the assembled var genes can be shown by mapping the short sequence reads back to the 3D7 reference sequence (Figure 1a). The areas of very low or highly variable sequence-coverage coincide with the positions of var genes. Thus, when the reads from one of our sequenced isolates were mapped to a random set of ten var sequences from the genome-reference clone 3D7, similarly patchy coverage was seen (Figure 1b, top panel). Given the known highly polymorphic nature of var gene sequences, we were surprised to observe that in some cases (Figure 1b, bottom panel) reads from a field isolate mapped almost completely to a single 3D7 gene; a gene from the reference clone was therefore almost completely duplicated in an unrelated field isolate.

Conserved var genes

Three types of var gene are known to be unusual in having greater sequence similarity to their orthologues than the repertoire as a whole: var1CSA is expressed late in the cell cycle and does not appear to reach the red cell surface22, var2CSA has an atypical domain architecture, is most expressed in infected women in their first pregnancy and mediates binding to chondroitin sulphate in the placenta resulting in low birth weight and premature delivery23. var3 is a short gene of unknown function, of which there are three highly conserved copies in the 3D7 reference and which have been shown to be expressed on the infected red cell surface23. Before carrying out any further analysis on our global var set, we therefore extracted these unusual var sequences. We found that var1CSA is universally present as two distinct allelic forms that resemble the sequences of 3D7 (38%) or IT (62%) and show no obvious evidence of recombination. The 5’ region of the 3D7 allele showed almost complete sequence conservation (only 0.5% divergence) compared with the same 3.2 kb region of the IT allele (3.6% divergence; Figure 2). A maximum likelihood tree drawn from the full-length amino acid sequences of var1CSA from all 714 isolates reflected the overall bi-allelic structure but showed more complex branching (Extended data: Figure S1). There appears to be no significant change in the ratio of allelic types by region and indeed the same bi-allelic pattern can be traced back more broadly to the Laverania subgenus from which P. falciparum emerged25.

Global sequence diversity and analysis in var2CSA has been recently reported for 1,249 isolates from the MalariaGEN data26 and so will not be described in detail here. However, in addition to the conclusions of these authors, we found very elongated var2CSA sequences of up to 15kb in some isolates (eight >12kb), and some similar extended sequences within the PacBio assemblies. The additional sequences are composed primarily of a single domain type, DBL epsilon (Extended data: Figure S2). Within this latter subset we find four sequences from South East Asia that share >99% identity with a 6kb region of the orthologue from P. praefalciparum, a parasite of gorillas that is the closest relative of P. falciparum.
Figure 1. Mapped-coverage of sequencing reads from clinical isolates. (a) Sequencing reads from a clinical isolate (PA0274) map at high coverage across much of chromosome 4 of the *P. falciparum* 3D7 reference genome. The distal regions of the subtelomeres and discrete interstitial regions contain *var* genes and are clear exceptions with little or no coverage, indicating high sequence polymorphism. (b) Sequencing reads from the PF0389 (upper) and PH0765 (lower) clinical isolates mapped against a reference comprising 10 concatenated *var* genes from *P. falciparum* 3D7 (alternating orange/brown). Few regions of the *var* genes are similar enough between isolates to enable reads to map and just one *var* gene is completely covered in one clinical isolate.

Figure 2. Sequence diversity of *var1CSA* genes from clinical isolates. *var1CSA* genes were split into two major types corresponding to *P. falciparum* 3D7 (top) and *P. falciparum* IT (bottom) *var1CSA* reference genes, based on a phylogenetic analysis (Addition File 2: Figure S1). Using BWA-MEM and SAMtools Pileup, sequence identity and polymorphism were detected and then plotted across each reference. The regions of each gene encoding specific protein domains are indicated. Genes of the 3D7 type have high sequence identity across their entire length but those of the IT type show greater polymorphism, particularly in their 3’ half.

The three *var3* genes in 3D7 show between 96–98% identity. In our normalised dataset we find 680 copies from 714 isolates with ~98% identity suggesting both a high degree of sequence conservation and degree of copy number variation. Indeed, within the PacBio dataset the number of copies varies from zero to six.

**Shared var sequences**

Having excluded genes known to have similar sequences, the most striking feature of the data, bearing in mind the highly polymorphic and recombinogenic nature of these genes, was the presence of long stretches of nucleotide sequence that are almost perfectly shared between *var* sequences of different
isolates. For instance, using stringent alignment criteria (>3.5 kb alignments with >99% identity and >80% overlap, used throughout this study unless otherwise stated) we identified 59,202 pairwise matches within the dataset representing 11,054 genes. We therefore reasoned that sharing of these polymorphic gene sequences at almost base perfect levels must have arisen from recent recombination events. Examining the length of the matches against their sequence similarity revealed a sharp decay in the length of matches when sequence identity dropped below 100% (Figure 3a). The longer matches must therefore represent recent events that have not had time to undergo recombination or SNP accumulation.

As a more visual representation of the way in which stretches of var gene sequence are shared between isolates, we constructed networks from a subset of the data where nodes (isolates) were connected if they shared a single var gene sequence using the same alignment criteria (Figure 3b). The same sequences could clearly be seen to be present across different continents showing an unexpected level of gene flow across the world. This analysis however hides a degree of granularity in the data (shown in Figure 3c) where nodes are only connected if they share at least three var gene sequences. At this more stringent level, the sharing of sequences between isolates is now dominated by South East Asian isolates and by two small groups of mainly African isolates. South East Asia is discussed in more detail later.

Following a more global approach to quantitate the level of sequence sharing, we clustered all of the sequences based on their top similarity hit using OrthoMCL and generated 3,351 clusters containing the 11,054 stringently aligned genes. After classifying the dataset into large geographic areas (West Africa, East and Central Africa, and Asia), we found 26 of the clusters containing 419 genes were present across all areas and 92 clusters containing 508 genes were shared between at least one African and one Asian area. Not unexpectedly, when we decreased the minimum length of the genes within a cluster to >2 kb, there were far more (241) clusters containing 22,466 genes of which 4,002 were present in all areas of Africa and Asia (Extended data: Table S2), reinforcing the conclusion that there is significant intercontinental gene flow.

Figure 3. Extensive sequence sharing between var genes. (a) Boxplot of nucleotide-alignment lengths versus sequence identity between var genes. (b) Network of var sharing between normalized dataset of 714 isolates. Each node represents an isolate, coloured by region. Edges represent isolates sharing at least one var gene (> 99% identity, 3.5 kb overlap and > 80% sequence overlap). (c) Alternative network of var sharing but with nodes (isolates) connected with three shared var genes.
To gain some insight into the distribution and chromosomal position of these shared \textit{var} genes, all of the complete \textit{var} gene sequences from our PacBio assemblies were BLAST-searched against our database and genes of \(>3.5\) kb with \(\geq 99\%\) identity were identified. A small proportion of genes from the PacBio-sequenced isolates (3.2\%) had a comparatively large number of hits (up to 55) and 81\% of these highly shared genes, with \(>10\) hits, were concentrated primarily in internal clusters on chromosomes 4 and 7 and to a lesser extent chromosomes 8 and 12 as well as a subtelomeric gene on chromosome 6 (Table 2). In the former case, they appear to be a consequence of a selective sweep by chloroquine resistance on chromosome 7\(^2\). For reasons that are not entirely clear, the signature of the selective sweep has not been eliminated by recombination in a subset of isolates (22) resulting in a 350 kb region around the selected locus including four \textit{var} genes remaining shared\(^2\). Similar arguments may apply to selective sweeps by anti-folate drugs on chromosomes 4 and 8. The reasons for an apparent additional selective sweep on chromosome 6 are unclear, but the sweep has been noted previously from SNP data\(^2\).

Sequence sharing is most striking amongst the SE Asian isolates (Table 1). In Cambodia, a distinct population substructure has arisen in which resistance to the drug Artemisinin (through point mutations in the Kelch gene) was associated with the expansion of non-interbreeding subpopulations from a recent bottleneck. To determine whether the level of \textit{var}-sequence sharing is due to this recent bottleneck, we built a network from the SE Asian assemblies (Figure 4) clustered by the Kelch

<table>
<thead>
<tr>
<th>Chromosome</th>
<th>(&gt;10) hits</th>
<th>%</th>
</tr>
</thead>
<tbody>
<tr>
<td>Chr 7</td>
<td>19</td>
<td>32.8</td>
</tr>
<tr>
<td>Chr 4</td>
<td>16</td>
<td>27.6</td>
</tr>
<tr>
<td>Chr 6</td>
<td>11</td>
<td>19.0</td>
</tr>
<tr>
<td>Chr 12</td>
<td>5</td>
<td>8.6</td>
</tr>
<tr>
<td>Chr 8</td>
<td>5</td>
<td>8.6</td>
</tr>
<tr>
<td>Other Chromosome</td>
<td>7</td>
<td>12.1</td>
</tr>
<tr>
<td>No data</td>
<td>2</td>
<td>3.4</td>
</tr>
<tr>
<td>Total</td>
<td>65</td>
<td>100.0</td>
</tr>
</tbody>
</table>

Table 2. The chromosomal position of \textit{var} genes. Only \textit{var} genes from the global set, with \(>10\) matches (>3.5 kb, with \(> 99\%\) identity and 80\% overlap) to a \textit{var} gene from twelve PacBio-based \textit{P. falciparum} assemblies were counted.

Figure 4. \textit{var} genes record pattern of artemisinin resistance in SE Asia. Network of \textit{var} sharing with each node representing an isolate, coloured based on polymorphisms in the \textit{kelch13} gene. Edges represent either (a) \(\geq 15\) shared \textit{var} genes (99\% identity, \(\geq 3.5\)kb and 80\% overlap), or (b) \(\geq 7\) shared \textit{var} genes. Dark blue isolates are where more than one SNP occurs in \textit{kelch13}. Additional samples were used for this figure (see methods).
mutation that the isolates contained. Moreover, by relaxing the network inclusion criteria from >15 var genes to >7, we were able to see likely mixed infections where different Kelch mutations may have come together. The bottleneck that produced these different populations probably occurred around 20 years ago with the introduction of Artemisinin, providing a useful baseline for tracking var gene evolution over time and for calibrating the relationship demonstrated in Figure 3a (Extended data: Figure S3).

Gene duplication
Var gene duplication within a single genotype has been observed in cultured isolates. We performed the analysis on the 12 full PacBio assembled genomes (Extended data: Table S3) and found between 4–21 duplications of at least 2kb. However, within our assembly process, such duplications would be assembled as single genes. We therefore examined intra-genome duplication by an isolate specific analysis of read depth. We chose samples containing a single genotype (defined as containing between 45 and 90 LARSFADIG sequences). We found a mean of 1.35 var genes/genome containing a partial duplication (range 0–10) and an average of 0.6 var genes/genome that were completely duplicated (range 0–8). These frequencies varied between Asian and African samples where we found slightly more duplication in the former (1.54 vs 1.19 per genome for partial duplication and 0.6 vs 0.54 per genome for complete duplicates). Thus, intragenome duplication events are relatively common and more frequent in areas of more limited genetic diversity. Whether this reflects an increased rate of inbreeding or is related to a higher var gene recombination rate in Asian parasites remains to be determined. Partial duplication indicates a meiotic or mitotic recombination event.

Recombination
Definitive evidence of recombination events within the var gene repertoire has previously required genetic crosses. However, the global var database contains deep recombination history. Having established that long stretches of near-identical sequences represent relatively recently shared sequences, we attempted to investigate recombination by looking for breakpoints in long shared sequence blocks. We called a breakpoint when conserved sequence blocks within the whole dataset of >1kb and >99% identity were interrupted by a second sequence block of >500bp with no hit within the database using our stringent criteria (see Methods, Extended data: Table S4) this gave us a total of 42,902 breakpoints in 35,574 var genes. Multiple hotspots are apparent from the data (Figure 5) showing that apparent recombination events within var genes were frequently detected but were not random. Within DBL domains, the most frequent events were in DBLγ (41.7% of total) followed by DBLγ (10.4%), DBLβ (8.4%) and DBLα (6.3%). Next most abundant were CIDRα (5.2%) and CIDRβ (2.8%). A large proportion (20.4%) of breakpoints were outside recognised domains (Extended data: Table S5). We also inferred a much deeper recombinational history from the data. For instance, by mapping the totality of the assembled var genes against 6 exemplar var genes at various levels of sequence-identity (Figure 5), the identity of the breakpoints was reinforced, with some genes having their entire sequence covered by high-identity hits from within the assembled data. To go back further in time, we repeated this analysis using three extant genes and three genes from P. praefalciparum as targets for the mapping (Figure 6). The deep history of these genes is clear with, in one case, the 5’ third of the gene being highly conserved between current P. falciparum and P. praefalciparum sequences.
As an alternative approach to looking at var gene recombinational history, we BLAST-searched our normalised database (with the addition of var2CSA sequences) against the 3D7 reference genome, recorded matches (> 1000 bp, > 99% identity) and reported them by UPS type (Table 3). First, it can be seen that var2CSA has a few 1000 bp hits, indicating that despite the relatively conserved nature of this gene enough SNPs must be present to abort the matches. Second, we can see that by this criterion, over 50% of the var gene sequences of Pf3D7 are unique, highlighting the level of polymorphism that exists. Finally, the upsC type has the most conserved sequences 58%, versus 27% of the other ups types. This class also has more breakpoints, especially in DBLδ (Table 3), suggesting that the high levels of recombination seen in this ups class and subdomain type may be facilitated by sequence homology.

Recombination within var genes of the progeny of a genetic cross has been associated with nucleic acid secondary structure.

Table 3. Number of exon-1 sequences of Pf3D7 var genes categorised by UPS type. Sequences shared with the normalised dataset with > 99% identity and an overlap ≥1 kb are shown. upsE is the var2CSA gene.

<table>
<thead>
<tr>
<th>UPS type</th>
<th>var genes</th>
<th>Total bases</th>
<th>% genes covered once</th>
<th>breakpoints per var gene</th>
</tr>
</thead>
<tbody>
<tr>
<td>UPS E</td>
<td>1</td>
<td>8,004</td>
<td>22.1%</td>
<td>0.0</td>
</tr>
<tr>
<td>UPS A</td>
<td>6</td>
<td>50,661</td>
<td>23.6%</td>
<td>1.8</td>
</tr>
<tr>
<td>UPS B</td>
<td>37</td>
<td>20,9940</td>
<td>29.8%</td>
<td>1.2</td>
</tr>
<tr>
<td>UPS C</td>
<td>13</td>
<td>70,740</td>
<td>57.9%</td>
<td>2.8</td>
</tr>
</tbody>
</table>

Figure 6. Ancient patterns of recombination within the Lavernia sub-genus. Similar to Figure 5, but P. falciparum var genes (blue) were selected that hit against a P. praefalciparum var gene (orange). The ribbons show matches of ≥ 99% identity 99% and a minimum length of 500 bp.
We used exactly the same approach (using RNAfold v2.1.8) to assess the secondary structure across the 100 bp regions centred on our apparent recombination sites. Compared with 100-bp control sequences sampled from all 66 var sequences of the PU0134-C sample and the PF3D7 reference, the free energy distributions look indistinguishable (Extended data: Figure S4) suggesting that the sites we identified are not specifically associated with DNA secondary structure. To determine whether specific sequence signatures define breakpoints, we used MEME to analyse 50bp either side of a random sample of 4,000 detected breakpoints. While there was no single consensus sequence in this analysis, a number of motifs occurred at high frequency and overall had a high GC content (43%) relative to the genome as a whole (19%) and var genes in general (36%). One highly significant motif accounted for over ten percent and only nine motifs accounted for nearly fifty percent (2,336 events; Extended data: Figure S5).

Domains and domain cassettes
It is clear from analysis of the Laverania subgenus that the var gene family is at least a million years old19. This subgenus is split into two clades (A and B). Within clade B, P. reichenowi and P. praefalci-parum show very similar gene numbers, chromosomal organisation and domain types and domain numbers to P. falciparum. Clade A parasites (P. gaboni and P. adleri) however show strikingly different CIDR and DBL subdomain content and organisation. Thus, the appearance of what closely resembles the current var gene content, organisation and domain structure emerged between 50,000 and 200,000 years ago and has shown no tendency to change significantly since this time.

The detailed analysis of PfEMP1 proteins to date has been based on data from seven genomes9. In view of the vastly increased volume of data presented here we undertook a reanalysis. Within our expanded dataset, as expected, existing definitions of DBL and CIDR domain types were concordant with existing data in P. falciparum. These domain types are also concordant with data from the much more anciently diverged sequences of the group A Laverania parasites, confirming that these sequence arrangements are both ancient and evolutionarily stable. We first carried out an analysis of main domain neighbours within the normalised dataset (Extended data: Figure S6). There are a number of highly enriched domain pairs, DBLα–CIDRα as previously reported and DBLζ–DBLε (>99%). Others that occur at high frequency include DBLε–DBLε (56%) and DBLβ–DBLγ (64%).

In view of the data using existing subdomain definitions that relates expression of particular domain cassettes with parasite phenotypes and disease severity we undertook an analysis of domain cassettes within our data. The complete analysis is presented in Extended data: Table S6 in which combinations of at least three subdomains that occur at least 10 times are shown. We found a total of ~86,000 triplets representing 1,567 different combinations that clearly included the previously defined domain cassettes. Four novel domain cassettes occurred >1,000 times and comprised ~7% of the total. We could not find a significant association of domain types with geographical regions.

Sub-domain definitions are currently based on trees drawn from multiple alignments of sequences of seven genomes. However, tree-based approaches are not ideal for showing relationships between highly recombinogenic sequences and have produced some anomalous results81. As an alternative, we used MEME to identify 256 six-nucleotide motifs within each domain or domain type and then clustered these by their co-occurrence within a single sequence. As an example of the utility of this approach we used it to classify the DBL and CIDR domains from the Laverania25 (Extended data: Figure S7). The robust identification in the Laverania of the main domains from P. falciparum reinforces the conclusion that the main domain architecture within the Laverania has been stable over long evolutionary time scales. We next applied the same approach to subdomains of P. falciparum. For example, Figure 7 shows the clustering of 1,000 DBL epsilon domains projected onto existing subdomain classifications. It is clear that there are a number of anomalies. For example (Figure 7, top box), DBLε10 and DBLε4 appear to be a single subdomain and a small portion of DBLε10 clusters with DBLε1 and DBLε11 (Figure 7, bottom box). By contrast, when we carry out the same set of analyses on DBLε6 sequences, the clustering (Extended data: Figure S8) clearly distinguishes between the major subclasses (0, 1 and 2) but does not reveal the same degree of substructure. The MEME analysis therefore suggests much more heterogeneity in the data making it difficult to assign robust sub-domains in this case.

Another method of viewing the high dimensionality of this data is a nonlinear t-SNE projection in two dimensions (Figure 8). For DBLε the tight clustering of the data into groups is again apparent (Figure 8a). Since the main goal of such sub-classifications is to relate sequence to phenotype, we looked closely at CIDRα1 sequences, the majority of which are known to bind to EPCR and whose expression has frequently been linked to severe malaria. When we do this, we find a clear discrete clustering (Figure 8b, boxed) of subtypes CIDRα1.2 and 1.3 that are the only two that show no binding to EPCR. The other subdomain that shows variable EPCR binding is CIDRα1.5 and this splits into two groups (circled). The rest of the projection clearly shows that while some of the data cluster consistently with current subdomain definitions, many do not, suggesting that the current CIDR subdomain definitions are not robust when analysed in this manner. The same projections for other DBL and CIDR domains are presented in Extended data: Figures S9-S11. The general conclusion from these figures is that while some existing subdomain definitions look robust to this analysis many do not. The most clearly resolved clusters can be seen in DBLε and DBLζ. DBLε is most closely related to the original and likely the most ancient DBL domain since it is found in many species and is present in proteins involved in red cell invasion. DBLζ is most abundant in the clade A Laverania that diverged from current P. falciparum populations around 1 million years ago23. This suggests that maybe because of their age, the subdomains in these classes may have functionally diverged (for example DBLε has diverged from other DBL domain types to bind IgM28), such that recombination between them is not favoured. In contrast, in the majority of other cases, functional diversification is not yet complete such that limited recombination is still going on. However, it should be noted
Figure 7. **Visualisation of motifs in DBLe domains.** Presence of individual MEME motifs (columns) in each DBLe domain (row) is shown (red). For each domain, annotation of subdomains, according to the VarDom server, is also shown (blue). The top dotted box shows the similarities between DBLe4 and DBLe10 and thus that they should be combined into a single subdomain. The bottom dotted box shows that a small sample of DBLe10 clusters with either DBLe1 or 11.

Figure 8. **Meme motifs DBLe and CIDRα domains.** Matrix of MEME abundance (from Figure 7) visualized using t-SNE plots for (a) DBLe and (b) CIDRα.
that because the granularity of the output is dependent on
the length of the MEME motifs included in the analysis, the
parameter space is too large for us to explore all possible
outputs but suggests that once a strong phenotype can be
associated with a given domain type then this approach may
be valuable in tying sequence variation to phenotype.

As an alternative approach to examining the way in which the
main domain types have evolved and continue to do so we
examined their relative diversity in two ways. First, we
plotted accumulation curves of the number of new sequences
with increasing sample size to show that the level of world-wide
sequence diversity varied significantly with domain type
(Figure 9). Most diverse were CIDRα, DBLα and DBLδ while
the least diversity was seen in DBLζ and DBLε (consistent with
the results above). Next, we took each domain individually and
carried out an all against all BLAST. The results were then
visualised as box-plots either of the overall diversity or as
the %identity of the top BLAST hit (Figure 10a). The mean
BLAST similarity varied from 30–80% with the most conserved

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**Figure 9. Diversity projection of domains.** Accumulation plots of the number of domains versus the number of unique domains. Only hits with ≥ 99% amino acid sequence identity over their full lengths included.

**Figure 10. Diversity in domains.** Boxplots showing sequence identity for 1,000 randomly selected sequences from each domain type based on an all-against-all BLAST analysis. In (a) all results are shown, in (b) the sequence matches for the top hits are shown.
domains being those that constitute var2CSA and the most diverse being DBLε. These values are similar to those described by 13. However, when the identity of the top hit was plotted (Figure 10b) a different picture emerges with DBLε having the highest identity. This is again consistent with the results above because of the well separated subgroups: within the subgroups identity is high but the groups are very diverse from each other.

**Discussion**

While there have been a large number of studies that have attempted to analyse the local or regional distribution of var gene sequences, these have to date mainly been limited to amplicons containing DBLα sequences, eg:1–9. Some smaller studies have included full length sequences but to date these have not involved sampling across the world. This therefore represents the first large scale attempt to catalogue the worldwide distribution of var gene sequences. We draw a number of conclusions from this study. First there is an unexpected level of var gene sequence sharing across the globe that falls into three main categories:

1. A very high degree of full-length sequence sharing in South East Asian isolates that is likely due to recent selection for parasites resistant to artemisinin;
2. A small proportion of sequence sharing that appears to originate from drug induced selective sweeps;
3. Sequence sharing that appears to have resulted from the recent intercontinental movement of parasites

In the case of (1) we find that the analysis of the var gene content alone is sufficient to partition the population into the same subgroups as well-established SNP-based population genomics approaches1 and if we increase the resolution by determining the number of isolates that share >50% of their var gene repertoire it becomes clear that there has been an almost clonal expansion of certain genotypes in some areas of this region (Extended data: Figure S3). By adding the date of collection to this analysis we can confirm that selection is still continuing, by the implementation of artemisinin combination therapy where resistance to the partner drugs is already appearing (Extended data: Figure S3). Thus, because var genes represent markers across the genome they can be used independently of SNP analysis to study recent selection events.

Second and somewhat surprisingly we can see in the distribution of var gene sequence the remnants of much older selective events involving chloroquine and anti-folate drugs on chromosome 7, 4 and 8. Why this signature has been retained in some isolates and not others is entirely clear. In the case of chromosome 7, those isolates that have retained six identical genes in the internal cluster are characterised by the presence of one (the fifth var gene) on the opposite strand that may have restricted recombination. Nevertheless, the fact that this event is present in 22 isolates suggests that there must have been some restriction on recombination since selection. Why the signature has presumably been retained on chromosomes 4 and 8 in some isolates only is unclear. With regard to chromosome 6, a local selective sweep of unknown cause has been reported in West Africa20 and this may well be the same signature that we are detecting. The origin of the var gene sharing on chromosome 12 is at present unclear. Since we only had 12 PacBio genomes with which to search our database, it is likely that a much higher proportion of these shared genes would be revealed as remnants of selective sweeps if we were to search a larger number.

Third, there is an additional transcontinental level of var gene sharing that does not appear to be chromosome enriched and therefore does not show evidence of a selective sweep. Our analysis suggests (because the length of sequence shared is related to the percent identity) that these simply represent recent recombination events that have resulted from parasite movement as a result of international travel. This observation reinforces the worry that there is background level of global parasite movement that may have consequences for the spread of drug resistance. These three different types of sharing are clearly evident in Figure 3b and c.

By analysing our database as a whole we were able to identify likely recombination events involving breaks of two blocks of identical sequence. These events are not random but instead are most common in DBLα domains (41.7%) and outside of DBL domains (20.4%). It is encouraging that the degree of domain structure revealed in the MEME t-SNE plots seems inversely related to the amount of recombination observed reinforcing the conclusion that some domain types are now relatively stable while others are still evolving. If we concentrate on those SE Asian isolates that share a high proportion of their var gene repertoire we can readily detect recent recombination events that appear to be more frequent in genes of the UpsC type (Extended data: Figure S3). In contrast with a previous report21 we do not find evidence of nucleic acid secondary structure associated with the breakpoints but do find a limited number of MEME motifs with a high GC content that account for ~50% of the breakpoints.

With regard to the unusual var genes that are relatively conserved, we reinforce the observation that var1CSA has two alleles that first separated within the Laverania infecting apes22. Within both major subtypes there has been a degree of diversification (Extended data: Figure S1) but no evidence of recombination suggesting that they are being maintained within the population by balancing selection. However, since the function of this gene is unknown, the source of this selection cannot yet be identified.

var2CSA sequences have been analysed elsewhere23, however, unexpected in our data was the presence of a number of highly elongated sequences extended by large numbers of DBLε domains and a region of ~6 kb in some isolates that is identical to a region of the gene from P. praefalciparum (Extended data: Figure S2). While the function of this gene in placental cytoadherence is well established it has also been reported to be involved as a central switching intermediate in antigenic variation24. Some DBLε sequences are known to bind IgM but whether this is pathologically relevant, what role the extended
sequences might play and why some have been maintained since the *P. praefalciparum* split are unknown. *var3* sequences show high sequence homology (~98%), a degree of copy number variation and some rare variants containing an alternative NTS sequence and a DBLα domain.

The cataloguing of different domain and subdomain types has been important in *var* gene research since it has been established that certain domain and subdomain types as well as combinations of them (known as domain cassettes, DC) are involved in binding to specific host receptors and associated with severe disease. Whether these DCs are functional units or not is however unclear since in the best studied case of DC8 and DC13 binding to EPCR, binding appears to be mediated by CIDRα alone\^35. While we had no clinical phenotype data here and have analysed genomic sequence rather than transcript data, our re-analysis nevertheless revealed some novel interpretations. While some domain types (particularly DBLε and DBLζ) show robust clustering into set of clear subdomains, the clustering of other DBL domains is less clear. The former two are likely the most ancient as they are the most numerous in clade A *Laverania*. In addition, DBLε is closest phylogenetically to the DBL type involved in other functions such as invasion, present across the genus and therefore likely to be ancient. They therefore appear to have segregated into what are presumably functional subtypes that now show little propensity for recombination to split them. In the case of the other types, we know that they have been long established since they are present in their present form in clade B *Laverania*. However, the inability to cluster them into unambiguous subtypes in most cases shows that a significant degree of recombination between them is still occurring. Indeed, our conclusion that this is correct is strengthened by the fact that the domain in which we found the most recombination events (DBLζ) coincides with the MEME analysis in which we found the least structure. The fact that they have not segregated into such distinct classes suggests that there is an ongoing process of optimisation possibly driven by host polymorphism in the host proteins with which they interact.

Conclusion

We have assembled essentially complete repertoires of *var* genes from global isolates of *Plasmodium falciparum*. This new resource enables key aspects of the evolution of this important multi-gene family to be unravelled. The repertoire has ancient routes since the number, domain content and genomic organisation have been established for more than 500,000 years\^35. Within that time frame, some domain types appear to have evolved to a relatively stable state that is structured but still polymorphic and that we hypothesise is related to the optimisation of their functions. Most domain types however appear to be still evolving as evidenced by the detection of recombination breakpoints and by the absence of defined structure in the t-SNE analysis. These breakpoints are characterised by a limited number of sequence motifs. Thus, while the evolutionary fitness of the family as a whole must be very high because of its longevity, optimisation is still occurring presumably driven either by function or by immune pressure.

Methods

Samples

Illumina sequence data were obtained from the MalariaGEN *Plasmodium falciparum* Community Project and the Pf3k project\^36,37. In addition, we included 12 unpublished Kenyan samples (PFKE01 - PFKE12) that will be described in more detail elsewhere. All samples (except the latter 12) were obtained directly from patients and sequenced using Illumina technology with read lengths of 75 or 100 base pairs.

Pipeline for assembly and annotation of *var* contigs

Sequence data for each sample were mapped against the reference genome (PGD7 version 3 from GeneDB)\^38. From the resulting mapping file (in BAM format), reads were extracted that either did not map to the reference genome or that mapped to *var* coding sequences, plus 500 bases up- and down-stream. Sequences were assembled and annotated using bespoke Perl and BASH scripts that executed the following steps: (1) Reads were assembled *de novo* using Masaucz\^39. (2) Overlaps (> 200 bp) between contigs were determined using MegaBLAST (parameters: -W 40 -F -m 8 -e 1e-80) and contigs were excluded if they overlapped for more than 90% of their lengths, with identity ≥ 99%. (3) Illumina sequencing reads were mapped using SMALT (version 0.7.4, default parameters) to the full set of contigs. Contigs were merged with a Perl script (findoverlaps_ver3.pl, [https://github.com/ThomasDOtto/IPA/](https://github.com/ThomasDOtto/IPA/)) if the overlap region was > 500 bp and if the coverage of mapped reads across the overlap region was 50% of the median coverage value, determined from the whole assembly. (4) Contigs were joined into larger scaffolds using SSPACE\^40 (version 2.0, parameters -n 31 -x 0 -k 10). (5) Gaps were closed between contigs using IMAGE\^41, for two iterations with k-mers of 71, and minimum mapping scores of 65. (6) Small errors were corrected with iCORN\^41 (version 2) for two iterations from reads aligned using Bowtie2. (7) Miss-assemblies were detected and broken based on aligned paired reads, using REAPR (version 0.7.4\^41, parameter -a). (8) Coding sequences were identified in each contig using Augustus\^41 (version 2.7) trained on the *var* genes of *P. falciparum* 3D7 version 3 and *P. reichenowi* PrCDC\^41. (9) The identified coding sequences were annotated with functional descriptions based on BLASTp, and Perl scripts were used to write out the nucleotide and amino acid sequences. (10) Estimation of coverage was performed by mapping the Illumina reads back against the extracted *var* coding sequences with SMALT (as above), read depth was counted as normalised to FPKM values.

Evaluation of assemblies

During development, the assembly algorithm was tested using paired reads (75bp, 150bp and 250bp) produced from the Pf3D7 reference (DNA was from the same stock as the reference genome project ), with a fragment size of ~500 bp, slightly longer than the Pf3k samples (mean 380 bp).
For quality-control purposes, whole-genome assemblies were used that were produced using single molecule real time sequencing (Pacific Biosciences) as part of the Pf3k project48.

A “reference set” of var genes, from both Pf3D7 and 15 Pf3k reference genomes, was used to evaluate the var genes assemblies (hereafter termed “Illumina set”). The DNA used for the Pf3k reference genomes was also sequenced on the Illumina HiSeq platform, using the same protocol as for the field samples (100bp paired reads48). The Pf3D7 DNA was from the same stock as the reference genome project and was sequenced on a MiSeq with 250 bp paired reads from a PCR-free library.

All predicted var genes ≥ 3kb from the Illumina set were compared, by BLAST-searching, against the coding sequences and raw assembly (output of HGAP48) of the reference set. Matches identity of 99.5% identity, covering > 95% of the shortest sequence were counted. LARSFADIG is a conserved amino acid sequence known to be present in the DBLα domain of var genes and its presence was used to crudely estimate the number of var genes in the uncorrected and final Illumina sets. Two samples from the reference set were known to contain mixed infections. However, these samples did not contain the full-expected number of var genes; several were present in a partially assembled form on contigs < 5kb that had been excluded from the Pf3k reference genome pipeline.

Pacific Biosciences reads were used to validate the contigs of the Illumina set for each isolate. First filtered raw reads (output of HGAP48) were chunked in 500 bp pieces and mapped independently using BWA-MEM46 (parameters: -x pacbio -t 16 -a -S -P). Each 500 bp sub-read was mapped as often as possible (parameter: -a) and discontinuities in the mapping of sub-reads were checked manually using BAMview47. For a correctly assembled var gene, the sub-reads should map continuously over the full length of the var gene. Errors were defined, if the first hit of a gene in the Illumina set had a non-mapping region > 200 bp, and the long mapping or validation approach confirmed the error.

The completeness of each assembly was calculated as the sum of all matches (≥ 500 bp overlap ≥ 99.5% identity) between the genes (≥ 500 bp) of the Illumina set and the genes (≥ 3 kb) of the reference set, divided by the total length of all var genes ≥ 3 kb in the reference set.

To establish the error rate, we divided the number of incorrect var genes from the 16 Illumina var gene assemblies by the total number of var genes > 3kb.

For the further analysis we used 12 PacBio Reference genomes and Pf3D7.

Normalized dataset
As the number of samples per country varied across the whole dataset, a dataset comprising assemblies from 60 isolates, from each of the 12 countries, was extracted. Where possible, samples were selected with close to the expected number of LARSFADIG motifs for a single parasite genotype (between 45 and 90 LARSFADIG sequences). Where this was not possible, the number of LARSFADIG sequences were increased. If still not enough samples could be retrieved, the lower limit was decreased. As we were interested in the variable first exon, rather than the more conserved exon 2, the 3’ end of each var gene was excluded if it matched (BLASTx -m 8 -e 1e-30 -b 5 -v 5) the exon 2 amino acid sequences from Pf3D7 or P. reichenowi PrCDC.

Conserved var sequences, like var1CSA, var2CSA and var3 (< 4kb), see below, were also excluded. The remaining var genes were compared with a MegaBLAST (-F F -e1-80). High sharing between several Gambian samples was observed, possibly due to repeated infections of the same vector in low endemic regions48. In four Kenyan samples (PC0016-C, PC0025-C, PC0080-C and PC0083-C), identical hits to PfDd2 were identified and these samples were also excluded.

In total, a normalized set of 714 genomes from 12 countries was generated that comprised 39,119 sequences ≥ 3kb from exon I. This dataset was used for most of the analysis, including general sharing, recombination, duplications. A further data set was created for use in Figure 4 that comprised additional sequences from Cambodia and Thailand, where they contained at least 40 LARSFADIG sequences and if they had a least 20 shared var genes to the Cambodian samples.

var1CSA, var2CSA, var3 analysis
To identify var2CSA copies across the data set, Pf3D7 var2CSA (Pf3D7_1200600) was BLAST-searched against the normalized var gene set. Contigs with an identity >95% were considered to be var2CSA candidates. Contigs > 3kb were further verified by checking for the presence of known var2CSA-specific domains (DBLpam1-3 and CIDRpam).

Using the three var genes from Pf3D7 as query sequences (PF3D7_0100300, PF3D7_0600400 and PF3D7_0937600), copies of var3 were detected in the global set using BLASTn with a minimum of 95% identity, an alignment length of ≥ 1kb, and a maximum gene length of 4kb.

The detection of var1CSA was more difficult due to its similarity with other var genes. However, the first 3.2 kb, from the 5’ end, is particularly conserved and this region was used to discriminate var1CSA from other var genes. A Maximum Likelihood tree was constructed from 5’ 3.2 kb regions of var1CSA. Alignments were generated with MUSCLE49 and cleaned in seaview50 first manually, then GBLOCKS51, default parameters). Trees were drawn with PHYML52. Alignments were visualized in JalView53. To analyse sequence variation across the lengths of var1CSA, copies were aligned with ≥ 80% overlap with BWA-MEM against either of the two var1CSA types, and variants called with SAMtools Pileup and BCFtools.

Domain analysis
Domains were first defined in the Plasmodium falciparum genome as part of the genome project1. The domains were refined
subsequently using var genes from seven lab-strains\textsuperscript{13} and are accessible through the VarDom server. From there HMMer models\textsuperscript{2} were obtained. Amongst those HMMer models, is the “duffy binding like” domain defined by Pfam\textsuperscript{3}. The HMMer models for the subdomains were built from the amino acid sequences extracted from the VarDom server. In all var genes longer than 3kb, the positions that encoded domains were identified using hmmmSCAN (parameters: -noali -E 1e-6 -domE 1e-6\textsuperscript{7}), and the HMMer models from the VarDom server.

Domains were assigned to the sequence, first by the highest score and ensuring that domain coordinates did not overlap by more than 10bp. The output of HMMer was parsed into a GFF file and a list of domains per gene. This enabled the domain order and distance between domains to be determined.

Domains from a further six Plasmodium spp from the Laverania subgenus were obtained in a previously published dataset\textsuperscript{25}.

Domain evolution
The domain diversity was investigated by searching for conserved motifs. MEME motifs\textsuperscript{60}, of 8–15, four or six amino acids were identified, rather than the perfect matches that have previously been reported\textsuperscript{13}. Basically, the programme meme\textsuperscript{66} search for the 96 or 256 most abundant motifs amongst CIDR or DBL domains from the seven Laverania genomes and 1000 random P. falciparum domains from each of the 18 domain types (ATS, CIDR\textalpha, CIDR\beta, CIDR\textgamma, CIDR\textdelta, CIRPam, DBL\textalpha, DBL\beta, DBL\gamma, DBL\delta, DBL\textepsilon, DBL\textgamma, DBL\textdelta, DBL\textepsilon, DBLpam1, DBLpam2, DBLpam3, DBL\textepsilon, Duffy\textunderscore binding\_like, NTS, NTSPam). For visualisation, the domains for each given gene were parsed from the meme\textunderscore txt output file into a binary matrix. Each row represents a different var domain and the column show the different meme motifs. A matrix entry is set to 1 if a meme motifs occurs in a domain. Using the heatmap2 function (ggplot2 package\textsuperscript{67}) and the average hierarchical clustering method in R\textsuperscript{68}, domains were visualized per sequence as a clustered distance matrix. In some cases, metadata like domain type or species were attributed as a barcode. To generate the t-SNE plots the R function Rtsne (parameter: perplexity=30, max\textunderscore iter = 2500) from the Rtsne library was used.

Recombination analysis
Breakpoints were obtained from successive filtering of Blast comparisons. First all the exon1 sequences of the normalised dataset were compared with themselves using megablast (-F F).

Alignment breaks as defined by BLAST matches are breakpoints if following criteria are met: high quality match (>1kb and >99% identity) between query and subjects, the sequence match does not span a gap, and the sequence of query and subject extends at least 200bp further. Also, the query and the subject cannot have a subsequent hit (identity > 95%) within 500bp of the sequence match. The potential presence of stable secondary structures was investigated by calculating free energy using RNAfold\textsuperscript{69}. From each predicted break points, 50 bp sequences up- and downstream were passed to RNAfold (version 2.1.8, parameters --paramFile=dna\textunderscore mathews1999.par --noGU --noLP --noPS --quadr --noTetra), and minimum energy was taken, as described in \textsuperscript{30}. As a control, the free energy of 100 bp sequences of all 66 exon1 sequences of PU0134-C (66 in total) were generated and the minimum energy was taken for each 100bp sequence. Histograms were plotted in R. This was repeated on the var genes of the Pf3D7 reference to generate a figure (Extended data: Figure S4).

Duplication analysis
To estimate duplication within a sample, the calculated RPKM of genes longer than 3kb were used. First multiple infections were excluded (as samples with more than 90 LARSFADIG motifs were excluded in the normalised dataset). If a var gene had an RPKM > 1.85 times of the median, the gene was considered duplicated. If the RPKM was between 1.35 and 1.85 of the median, the var genes was considered partially duplicated. Those values were manually checked in BAMview\textsuperscript{87}.

Analysis of var sharing between isolates
OrthoMCL\textsuperscript{60} was used with default parameters to cluster var sequences. The BLAST input was generated with MegaBLAST -F F and the identity and overlap were filtered with awk to generate clusters with different specific cutoffs.

Networks isolated linked by shared var genes were generated with Gephi\textsuperscript{86} clustering performed through the Fruchterman algorithm\textsuperscript{82}.

Analysis of var genes from Southeast Asia
The Southeast Asia dataset was visualized with Gephi, see above. To colour the nodes, the position of the mutation in the Kelch13 gene (PF3D7_1343700) was used. It was called from the BAM files using mpileup (SAMtools\textsuperscript{90} and BCFTools. Metadata for time and location were used from ftp://ngs.sanger.ac.uk/production/pf3k/release_5/pf3k_release_5_metadata_20170804.txt.gz.

Ethics approval and consent to participate
All samples were previously published and had ethical consent.

Data availability
Underlying data
Zenodo: Extended data for “Evolutionary analysis of the most polymorphic gene family in falciparum malaria”, http://doi.org/10.5281/zenodo.3549732\textsuperscript{64}.

This project contains the following underlying data:
- Additional File 1: Table S7: Overview of all assembled samples with >10 LARSFADIG motifs. Key metadata are shown, along with the number of LARSFADIG motifs and var genes at different length cut-offs available.

The accession numbers of the raw data from 12 previously unreleased Kenyan samples are in Additional file 1: Table S7. All other accession numbers can be found from the MalariaGEN P. falciparum Community Project Pf6 release at https://www.malarigen.net/resource/26 or the Pf3k release at
ftp://ngs.sanger.ac.uk/production/pf3k_release_5/pf3k_release_5_metadata_20170804.txt.gz

Code used to generate the var genes: https://github.com/ThomasDOTto/varDB.

Archived code as at time of publications: https://doi.org/10.5281/zenodo.354977

Data can also be accessed by BLAST-searching from https://www.sanger.ac.uk/action/BLAST

Extended data

This project contains the following extended data:
- Additional File 1: Tables S1-S6.
- Additional File 2: Figures S1-S11.

Data are available under the terms of the Creative Commons Attribution 4.0 International license (CC-BY 4.0).

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This is a review of “Evolutionary analysis of the most polymorphic gene family in falciparum malaria” by Otto et al. The other reviewers have summarized the paper in their comments, so I will not rehash their summaries.

In short, this highly anticipated analysis represents a fantastic amount of work, assembling 2400 var repertoires from a globe-spanning set of *P. falciparum* genomes. The analysis is thorough, well documented, interesting, and revealing. It is written at a level that is easily readable by the var community while also being interesting (I think) to the wider community.

Still, I have a few suggestions and comments which I hope will increase the impact and improve the readability of the manuscript.

Limitations:
- I found this paper to be really strong, and I have no doubt that it will inspire substantial downstream research from many groups. To that end, I'd like to encourage the authors to include, at the end of the discussion, some thoughts on the limitations of the study and its interpretations of the data.

Multiple infections:
- I am mildly concerned about the impact of the MOI>1 samples. If the goal of imposing the 90 LARSFADIG threshold is to avoid MOI>1, but the reason that the African isolates have higher average numbers of LARSFADIGS is due to higher MOIs, this makes me wonder if the cutoff is too permissive. It seems self-contradictory to say that a cutoff was imposed to limit MOI, but that higher average counts were because the cutoff let higher MOI samples through. At the same time, there doesn't seem to be anything to do other than to possibly vary the cutoff, which (I'm guessing) would result in a statement like “we varied the cutoff.
and the results do not substantially change."

- However, this came up again during the Gene Duplication section on Page 8. What fraction of the gene duplications or partial duplications could be based on multiple-infection samples? It seems like some of the observed duplications here could be attributable to both (1) actual partial or complete duplication and (2) multiple infection. I don't know what the right null model would be for the scaling of duplications with number of vars in confirmed MOI=1 infections, but it would be interesting to see the extent to which larger repertoires were predictive of these partial and complete duplication observations.

- Perhaps relatedly, or perhaps not, the Evaluation of Assemblies section of the Methods refers to "Two samples from the reference set" which contained mixed infections. I didn't understand the implications of that paragraph for MOI and QC more broadly.

**Be careful with tSNE:**
- tSNE is fantastic but lacks interpretability in ways that are familiar from PCA or embedding methods. I won't recapitulate the issues with it, but I would like the authors to check that other tSNE perplexity values don't change their visually appealing results and interpretations. This page recaps issues with tSNE in a visually compelling set of examples. [https://distill.pub/2016/misread-tsne](https://distill.pub/2016/misread-tsne)

**Minor things:**
- The copy number variation of var3, noted on Page 5, struck me as interesting. Is there anything interesting that can be said of the copy number distribution?

- Figure 3b is difficult to make sense of, but I wondered what it would look like as a heatmap of the network's weighted adjacency matrix, organized by region? I'm guessing that there's plenty of structure, but the sheer number of edges in the network obscure it.

- On Page 8 it says, "The bottleneck that produced these different populations probably occurred around 20 years ago." Is this an actual probabilistic statement or Occam's Razor reasoning based on the patterns in Figure 4? I got thrown off by "probably," in other words.

- On Page 8, in the Recombination section, it says that definitive evidence has previously required genetic crosses. Is the Claessens et al. (2014) work not definitive? (This is an honest question – it is not meant to be rhetorical.)

- On Page 10, in the discussion of Domain Cassettes, you note some enriched domain pairs (DBLz-DBLe). First, what do the parenthetical 99%, 56%, and 64% mean, exactly? Second, are there also domain pairs that are obviously precluded, or domains that are mutually exclusive?

- Figure 7 was a mystery to me. My general impression is that the MEME motifs - the horizontal axis label has inconsistent capitalization of MEME, by the way - support the DBL types but not the subtypes. In other words, the argument is that the previously constructed subtypes were overfitting the data, and dividing domains into substructures that aren't truly meaningful. That's fine, but the figure itself remains confusing. What are all the dotted lines? Which ones do match? Can the font sizes be increased, too?
In the Discussion on Page 13, you write that "These events are not random but instead..." and then there are parenthetical percentages. What do these percentages mean?

**Trivial/Typographical suggestions:**
- Abstract: periods at the ends of the first two paragraphs?
- Results: "from approximately 2400" -> 2398. No need for rounding in line one of the results.
- Figure 1: What are the vertical axes? Annotations are hard to read. Could colors that are higher contrast than orange/brown be used, by chance?
- Figure 2: Are higher resolution (or vector) formats available?
- Page 4, last paragraph: "epsilon" is spelled out, inconsistent with the rest of the text.
- Page 7, left column: The sentence that contains "our PacBio assemblies were BLAST-searched against our database" was confusing to me. Something about the wording and the "our"s made me question whether I understood exactly what was meant by our PacBio assemblies and our database.
- Same as above. "In the former case" confused me, given the complexity of the previous list. Could you be explicit about the case?
- Same as above. "in a subset of isolates (22)". I printed the paper to read it and couldn't tell if this was a parenthetical reference or a number of isolates.
- Page 7, right column: "clustered by the Kelch mutation" - I think this is not exactly correct. The nodes are *colored* by their Kelch mutations, but visually clustered by their var repertoire overlap through the network layout, right?
- Page 8: "an isolate specific analysis" -> "an isolate-specific analysis".
- Figure 5 is very interesting, but I'd love to see it larger, like Figure 6 is.
- Page 9, left column: "First, it can be seen...must be present to abort the matches." I didn't understand this sentence. Could it be clarified?
- Figure 8 has annotations that are described in the text, but not in the figure caption - but the figure precedes that main-text description.
- Figure 9: Can you just annotate the curves directly? I bounced back and forth between the legend and curve and had to just draw the annotations on the paper with a pen in the end.
- Page 14: "Miss-assemblies" -> Misassemblies?
References

Is the work clearly and accurately presented and does it cite the current literature?
Yes

Is the study design appropriate and is the work technically sound?
Yes

Are sufficient details of methods and analysis provided to allow replication by others?
Yes

If applicable, is the statistical analysis and its interpretation appropriate?
Yes

Are all the source data underlying the results available to ensure full reproducibility?
Yes

Are the conclusions drawn adequately supported by the results?
Yes

**Competing Interests:** No competing interests were disclosed.

**Reviewer Expertise:** Complex networks, statistical inference, mathematical models, var genes

I confirm that I have read this submission and believe that I have an appropriate level of expertise to confirm that it is of an acceptable scientific standard, however I have significant reservations, as outlined above.

Reviewer Report 19 December 2019
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Thomas Otto et al. here presents their long-anticipated analysis of near-full length de novo assembled *P. falciparum var* genes from genome sequencing of almost 2400 parasite isolates originating from across the world. Understanding the genetic evolution of *var* genes and the
organization and variation of the encoded PfEMP1 protein family is an important prerequisite for understanding malaria immunity and pathogenesis. The authors have succeeded in the quite daunting task of condensing the extensive dataset into easily read and interpreted figures explaining their main observations of the gene families organization as well as novel demonstration of var gene sharing and flow linked with spread of drug resistance. The data set itself constitutes an important contribution to the scientific field, and I expect it will form basis of multiple subsequent analyses of car genetic evolution and PfEMP1 function.

Through evolution, var genes have diversified through acquisition of point mutations and recombination. Despite their extensive diversity, the var genes are organized chromosomally, in their recombination patterns and in their encoded domain composition in a manner that is associated and facilitating, functional diversification of the protein family. The higher order of var gene division into the UPSB and UPSA genes located on opposite strands in the telomeric regions of the chromosomes and the UPSC genes in the chromosome centromere, and the linkage of this division with the type and phenotype of encoded N-terminal domains (UPSB and UPSC var encoded PfEMP1 binds CD36 through their CIDRa domains, UPSA binds EPCR through CIDRa1 or unknown receptors through CIDRg/d/b domains, and the subtelomeric UPSB chimeric B/A type PfEMP1 aka DC8 binds EPCR via CIDRa1), have been described previously, including in several reviews of PfEMP1. Similarly, the typical domain composition of PfEMP1 and PfEMP1 repertoire of each parasite genome, has been described before by the authors. Thus, while these traits are firmly confirmed in the present study, they are understandably not the focus of this study. However, they are key to understand and interpret the observations made throughout the paper, and so I suggest a few simplifications to the introduction as well as some corrections of terminology to ensure consistency with previous papers and help newer readers along (see minor comments below).

On shared var genes:
Although the present data assemblies cannot directly infer chromosomal localization of the sequenced genes, most of the genes found to be shared across parasite genomes appear to be centromeric UPSC genes (perhaps calculate and give the proportion of the 11,054 shared genes with high similarity to known centromeric genes of the 15 pacbio reference genomes). As the data suggest, for unknown reasons, these var genes may not recombine as frequently as telomeric var genes, or at least that is as possible explanation for the unexplained conservation of the var genetic signature linked to the drug selective sweeps.

On Recombination:
The mechanism of var gene recombination has been the center of several studies incl. the referred paper by Sander et al. (2014)\(^1\), in which chimeric genes were observed in genetic cross progeny clones resulting from ectopic mitotic recombination, i.e. new var genes were results of recombination between var genes of different chromosomes within the same parasite clone (no meiotic rec. events was found in this study). Characteristic of the chimeric genes was multiple closely spaced homologous recombination events. Apart from sequence homology, homologous recombination requires proximity of recombining genes and an initiating event such as DNA strand break or stalling of transcriptional or replication forks (reference) which can cause template switching such as the one observed in the mitotically recombined var genes. The observation that DNA regions with high propensity to form DNA secondary structures (DSS) were condensed around recombination hotspots, in variable sequence regions of var genes and predominantly in recombining PfEMP1 domains, lead to the hypothesis that var genes recombine through
homologous recombination initiated by replication-impeding DSS and and facilitated by bouquet formations of chromosome ends during cell replication Freitas-Junior et al. (2000).²

The identification of recombination sites, in the present study will probably not identify the shortly distanced recombination sites similar to those observed in the parasite cross progeny, and would include meiotic recombination events as well. In addition, the recombination events found appear to be dominated by events within UPSC genes, which appear to recombine less frequently and possibly by a different meiotic mechanism. Altogether, this may explain the suggested discrepancy between this and the Sander et al. study in linking DSS to var recombination. However, as homologous recombination sites following replication fork stalling can occur long distances from the initiating event, I would guess the present extraction of recombination sites should capture many of the events leading to the previously determined recombination hotspots and the proposed link to DSS.

Indeed the authors find that specific recombination sites are not located randomly in the genes, but found in “Hotspots” on the specific gene, which again is non-randomly located, found in particular locations in the var genes, e.g. in the DBLd domains. The data presentation does not allow precise evaluation whether the gene variant specific hotspots aggregate around a specific location in the genes in general e.g. the “mid var genes”, at the DBLd 5’ end. However, the identification of a few short DNA motifs that are common near the majority of recombination sites shows that this is probably the case. A quick search of the localization of the most frequent DNA motifs shows that indeed these are found at the previously identified hotspots of var gene recombination inferred from domain and subdomain homology breaks, i.e. at the 5’ end of DBLd domains and the border between DBLa subdomain 2 and 3 (MEME motif #2 appears to encode homology block 2 of DBLa, a conserved functional site in DBL domains). Please provide an analysis or statement whether this is true, as this is would be an important result in itself to be highlighted.

This will also allow the authors to refer to at least these hotspots association with DSS positions, as the two hotspots are the exact same locations which was associated with high frequencies of DSS in Sander et al. It is possible that the authors could not find this association as the exact recombination sites +/- 50 bps is skewed somewhat from the hotspots used in Sander et al. (HB2 and var mid-point), or that adding meiotic recombination events disturbs the analysis. However, DSS mathematics around these strikingly similar sites should add up. It is unclear what the use of the reference sequences were and in relation to figure S4 showing examples of UPSC var gene hotspots. The authors do not report any statistical test or method, and appear not to have applied the same statistics as Sander et al. to test positional association of high propensity DSS with the newly defined recombination hotspots. In Sander et al. a cutoff for lowest energy DSS was applied (e.g. the 1st or 3rd percentile of the 50mers with strongest prediction of forming DSS), and the distance of these DSS was tested for proximity association with the observed recombination hotspots.

In any case, it is unclear which specific role the authors suggest that the conserved sequence motifs should play in recombination, if not as landmarks to the general hotspots?

On domains and domain cassettes:
Each main PfEMP1 domain type exhibits its own distribution of sequence diversity. These domain specific differences in sequence diversity homogeneity and the fact that many domain subclass intermediates exist, makes unbiased and strict domain classification difficult. In the lack of better
and to give an operational nomenclature, Rask et al. applied a simple empirical rule to name all domain possible classes, which was to assign a domain subclass to any group of whole domain sequences containing at least 3 of 7 genomes investigated using maximum likelihood trees. As the present data very nicely show, this approach is not ideal for recombining genes, but still robust for some domain classes, again probably reflecting the different recombination frequencies within these.

I think the authors have made a wise choice in not attempting to re-name the domain classes, and instead provide beautiful t-SNE plots of each main domain class. This will ease future studies of domain function. For this, I appeal to the authors to include a fasta file with each domain DNA sequence annotated with its “suggestive” subclass type domains for the full dataset. Also, if possible please provide high-resolution images of these t-SNE plots to provide better readability of the legend, and move legend away from spots.

In some cases, the authors understate the robustness of their data. E.g there is no mentioning of CD36 binding CIDR and their clear division from the EPCR binding CIDRa1 in Figure 8 and S7. Legend in Figure S8 states DBLa subtypes does not cluster, although they do into DBLa1 (USPA) and DBL0 (UPSB/C), as started in main text.

There are some inconsistency in the mentioned length of MEME motifs – 256 six bp motifs stated in main text and Figure S8, whereas Figure S7 states 15-30 bps?

Minor comments and suggestions:

1. There is an inconsistent use of “main”, “subclasses”, “subtypes”, “type” and “subdomain classes” in the manuscript.
   - To follow previous terminology I suggest that DBLa,b,g,d,z and CIDRa,b,g,d domians are referred to as “main” classes; e.g. in the sentence “The DBL and CIDR regions have been divided into few main classes DBLa,b,g,d,z and CIDRa,b,g,d based on sequence similarity.” Or alike.

   - I suggest not to use the term “subdomain” but instead use domain “subclass” of the main DBL classes. In PfEMP1, and other proteins, subdomains refer to a smaller functional unit within the domain. E.g. the DBL domain structure is conventionally divided into three subdomains, S1-3., Singh et al. (2006).

2. The sentence: “Analysis of many thousands of such amplicons first produced a classification system for members of this domain into six classes based on cysteine content, amplicon length and the presence of certain sequence motifs. Surprisingly, considering the fact that these amplicons only represented a tiny fraction of the total gene sequence, associations have been found with the expression of particular DBLa types and parasite phenotype.”.....is not helpful to the reader. The DBLa tag nomenclature is not used further in the paper, and in fact the ambiguous linkage between Cys2/group A-like DBLa domains and EPCR and non EPCR binding CIDR domains in group A and Cys4 group B like DBLa0 tags and CD36 and EPCR binding in group B and C genes, may appear confusing to readers. I suggest the paragraph is used to present the functional diversification of the majority of PfEMP1 into those binding CD36 via and those binding EPCR via their specific variants of N-terminal CIDR domains, also associated with chromosomal var location, which will help interpretation and understanding of the distribution of genetic diversity presented
later.

3. And in the following paragraph: I do understand the need to present the domain cassette idea. However, this could perhaps be introduced later under the section investigating putative domain cassettes or changed not to relate to EPCR/or to reflect current it has been clarified that the EPCR binding phenotype is not tied specifically to the DC8 and DC13 domain cassettes, defined from the 7 first analysed genomes Lau et al. (2015)⁵.

For molecular evolution studies and for developing phenotype specific molecular tools the domain cassette concept has been valuable, and I think the present study's clarification of the consistency or rather inconsistency of the previously defined domain cassettes is a welcoming opportunity to clearly stipulate that many of these cassettes (outside var1-3, DBLa-CIDR, DBLd-CIDR, perhaps DC5) are more likely results of recent genetic inheritance than functional units.

4. I suggest correcting the sentence: “gene...:var1CSA is expressed late in the cell cycle and does not appear to reach the red cell surface22” to “var1CSA is expressed throughout the cell cycle and its product does not appear to reach the red cell surface.” (Var1 does not appear to have a late stage specific expression but rather a cell cycle detached expression).

5. I suggest changing “The latter are evaded through transcriptional switches among the var gene family such that new sequences are periodically expressed. Through their role in immune evasion they have evolved to be extremely polymorphic” to “On the red cell surface, PfEMP1 proteins mediate both adherence to endothelium and are targets of host protective antibodies. To escape immune recognition the PfEMP1 family have evolved to be extremely polymorphic and the parasites can switch var gene expression during asexual replication.” – or similar clarification/simplification.

6. I don't think the conclusion can be drawn that “functional diversification is not yet complete” from the observation that a domain class is highly recombinogenic. This somehow implies that the parasite is actively seeking for receptors to bind or alike, which does not fit with ancient nature of the protein family. High diversity or recombination rates within specific domain classes, may be their prime function, or it could be that variants have been selected for function and indeed are functional within the sequence space of a long gone settled steady state of recombination (last paragraph page 10).

7. Likewise I suggest rephrasing: “the conclusion that some domain types are now relatively stable while others are still evolving” – to “the conclusion that some domain types are relatively stable while others are highly recombinogenic”.

References


Is the work clearly and accurately presented and does it cite the current literature?  
Yes

Is the study design appropriate and is the work technically sound?  
Yes

Are sufficient details of methods and analysis provided to allow replication by others?  
Yes

If applicable, is the statistical analysis and its interpretation appropriate?  
Partly

Are all the source data underlying the results available to ensure full reproducibility?  
Yes

Are the conclusions drawn adequately supported by the results?  
Yes

**Competing Interests:** No competing interests were disclosed.

**Reviewer Expertise:** Antigenic diversity, malaria.

I confirm that I have read this submission and believe that I have an appropriate level of expertise to confirm that it is of an acceptable scientific standard, however I have significant reservations, as outlined above.

Reviewer Report 11 December 2019

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The submission by Otto and colleagues describes a long-awaited study on the evolution of the var gene family in the Laverania lineage of malaria parasites. The analysis is a tour de force, including var gene sequences from a huge number of parasites isolated from all over the world. The massive number of sequences enabled the authors to infer evolutionary relationships within the gene family and the existence of universally conserved var genes. In addition, by examining the flow of var gene sequences between parasites, the authors could clearly identify selective sweeps that have made their way through global parasite populations, including recent selective sweeps caused by the spread of artemisinin resistance. The paper describes a valuable resource and a comprehensive analysis of this important gene family that has implications for public health as well as our understanding of parasite evolution. Below are comments regarding a few issues of clarity and presentation.

Figure 1a provides a nice visual representation of the degree of sequence conservation across chromosome 4 comparing isolate PF0389 with the reference 3D7. The polymorphic nature of the var gene clusters is quite easily observed. However, for the non-aficionado a couple of aspects of the figure are not clear. Specifically, what is shown on the vertical axis (there is no scale)? What are the sharp spikes in sequence conservation that are found flanking the “left” subtelomeric region and the internal var clusters? These regions appear to be much more conserved than even the core genome, although without any scale it is difficult to say how much more conserved. A more precise description might be available in the methods section, although a short description here would be helpful.

The authors provide interesting data regarding var1csa, in particular identifying two distinct alleles and confirming its conservation throughout the globe. This gene has been a bit mysterious since it was first described. The gene is annotated as a pseudogene in 3D7 due to its premature truncation, and it similarly displays frame-shift mutations (often in exon 2) in the full gene sequences of numerous isolates previously published by this group. Do the data from this study enable the authors to determine if the gene typically (or universally) displays frame-shift mutations in its open reading frame? The intron of var1csa is similarly short and appears to be missing the region associated with transcription of noncoding RNAs that have been implicated in var gene transcriptional regulation. Is this similarly a universal characteristic of this gene?

Figure 4 provides a beautiful display of var gene sharing as a result of the development and spread of artemisinin resistance in SE Asia. This figure (along with figure 3) demonstrates visually how drug resistance spread. While beyond the scope of the current report, I am curious if there are additional analyses that could be relatively easily derived using this approach. For example, is there additional information associated with each isolate that would enable the authors to incorporate into figure 4 the precise geographic location from which each isolate was obtained? In other words, could these data be used to show where resistance first arose and how it spread throughout SE Asia? Similarly, drug treatment failure in SE Asia is often associated with failure of the partner drug that is supplied with artemisinin, most frequently piperaquine. Would analysis of piperaquine resistance likely show a similar display or would there be a different pattern?

Figure 7 displays Meme motifs, domain and DBL types as a single figure. The image shows clustering, however this was not initially intuitive to me. This might be easier to understand if a cluster tree was displayed on the left as is often done with heatmaps. More importantly, the
dotted box near the top of the figure is described as showing that DBLe4 and DBLe10 show significant similarities and might better be considered as a single subdomain. I gather this conclusion comes from the organizations of the DBL types in blue on the right of the figure. However, the top dotted box seems to include DBLe10 with DBLe3 rather than DBLe4. Is this an error in how the labels on the horizontal axis were aligned?

**Is the work clearly and accurately presented and does it cite the current literature?**
Yes

**Is the study design appropriate and is the work technically sound?**
Yes

**Are sufficient details of methods and analysis provided to allow replication by others?**
Yes

**If applicable, is the statistical analysis and its interpretation appropriate?**
Yes

**Are all the source data underlying the results available to ensure full reproducibility?**
Yes

**Are the conclusions drawn adequately supported by the results?**
Yes

**Competing Interests:** No competing interests were disclosed.

**Reviewer Expertise:** Transcriptional regulation and DNA recombination in P. falciparum.

I confirm that I have read this submission and believe that I have an appropriate level of expertise to confirm that it is of an acceptable scientific standard.