Paths to a malaria vaccine illuminated by parasite genomics

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More human death and disease is caused by malaria parasites than by all other eukaryotic pathogens combined. As early as the sequencing of the first human genome, malaria parasite genomics was prioritized to fuel the discovery of vaccine candidate antigens. This stimulated increased research on malaria, generating new understanding of the cellular and molecular mechanisms of infection and immunity. This review of recent developments illustrates how new approaches in parasite genomics, and increasingly large amounts of data from population studies, are helping to identify antigens that are promising lead targets. Although these results have been encouraging, effective discovery and characterization need to be coupled with more innovation and funding to translate findings into newly designed vaccine products for clinical trials.

The need for a malaria vaccine

The drive to sequence malaria parasite genomes began in the 1990s with a bold vision for ‘vaccinomics’ which would showcase the benefits of genomics to society [1]. This was well conceived, and remains relevant because malaria causes at least half a million deaths and 200 million clinical cases each year, with several species of *Plasmodium* parasites infecting humans via transmission by *Anopheles* mosquitoes [2]. The disease occurs in most tropical areas of the world, with the greatest burden being on children and pregnant women in Africa where *Plasmodium falciparum* is most prevalent. Existing interventions need to be more widely applied, including insecticide-treated bed nets to reduce mosquito biting, antimalarial drug prophylaxis targeted to vulnerable groups, and access to prompt diagnosis and treatment. An effective vaccine could be a cost-effective additional means of controlling malaria [3] and might enable its elimination from some areas.

A key issue for malaria vaccine design is selecting the most appropriate parasite life cycle stage to be targeted (Box 1) [4,5]. A vaccine to prevent blood-stage infection from occurring could target the parasite at the initial pre-erythrocytic stage, but would need to be fully effective because even a small number of parasites emerging from the liver can initiate a severe blood-stage infection. By contrast, a vaccine targeting the asexual blood-stage directly could be effective in suppressing the replicating parasites and thereby preventing most disease, even if it did not achieve absolute sterile immunity. As a potential complement to either of these, vaccination against parasite sexual stages would not prevent disease directly but might reduce transmission to mosquitoes, thereby possibly having a beneficial effect at the population level. Molecular characterization of these developmentally differentiated stages was an early goal of genomic and transcriptomic studies.

The first genome sequence of a malaria parasite was published in 2002 for a cultured strain of *P. falciparum* containing approximately 5500 genes encoded on 14 chromosomes with a complete haploid genome size of ~23 Mb [6] (Figure 1). Genome sequences of diverse malaria parasite species, including *P. vivax*, the second most important cause of human malaria [7], the zoonotic *P. knowlesi* [8], other primate malaria parasites [9,10], and parasites of rodents which can be studied in laboratory mice [11], show similar core genome organization. Each of the chromosomes shows a largely conserved syntenic gene order [12], except in the subtelomeric regions which contain large gene families that have diverged substantially between species [10,12,13]. Within each species, meiotic recombination occurs in a brief diploid stage within the mosquito midgut after fertilization between haploid male and female gametes in the blood meal. Recombination occurs at an average of 1 cM per 10–30 kb in experimental crosses of *P. falciparum* [14], although effective recombination rates in natural populations can be lower due to self-fertilization [15]. The diversity of parasites within infections directly influences the rate of outcrossing, which is greatest in highly-endemic populations where superinfection by unrelated genotypes commonly occurs [16]. Subtelomeric chromosomal regions can additionally undergo recombination during mitotic replication in the asexual blood stage of infection, with crossing-over occurring between non-homologous chromosomes in the same genome [17]. For vaccine design, the genetics and development of the parasite need to considered in the context of host immune responses that might suppress or eliminate an infection.

Feasibility of vaccination

A degree of immunity to malaria is generally acquired by experience of the infection, as shown by experimental studies in the first half of the 20th century on patients with neurosyphilis. Fevers induced by administering malaria parasite infections were beneficial to such patients in the age before antibiotics, because elevated temperatures
Box 1. Malaria parasites – species and life-cycle targets for vaccination

The major malaria parasites of humans are *P. falciparum* and *P. vivax*. Vaccine development is generally considered separately for each of these because they are distantly related and have only limited antigenic cross-reactivity. Other human malaria parasites *P. malariae*, *P. ovale* (comprising biological species *P. o. curtisi* and *P. o. wallikeri*), and *P. knowlesi* (a zoonosis from macaque monkeys) are less common globally and are not considered for vaccine development. Malaria parasites are haploid throughout the cycle except for a brief diploid zygote stage after fertilization in the midgut of the mosquito. Phases of the parasite life cycle that are candidate targets for vaccination are:

**Pre-erythrocytic stage.** Invasive sporozoites inoculated by mosquitoes and subsequent development to large schizonts within the liver (taking ~6 days in *P. falciparum*, and 10 days or longer for *P. vivax*) which do not cause disease but are a potential target to prevent blood stage infection.

**Asexual blood stage.** Causes all the symptoms of malaria as a result of significant parasitaemia achieved by repeated multiplication cycles (every ~48 h for *P. falciparum* and *P. vivax*), with merozoite invasion followed by growth and replication of the parasite within the erythrocyte (forming early trophozoite ‘ring’ stages, late trophozoites, and schizonts) before the release of more merozoites.

**Sexual stage.** Male and female gametocytes in the blood are ingested by mosquitoes and transform into gametes and ensuing motile zygotes (ookinetes) which might be targeted by antibodies taken with the parasites into the blood meal.

killed spirochetes more effectively than other available treatments, but many patients became immune to repeated malaria infections [18]. Studies on naturally acquired immunity in endemic populations then showed that malaria parasite levels in children or non-immune adults were reduced by passive transfer of serum Immunoglobulin G (IgG) from relatively immune adults, which has encouraged subsequent focus on the role of antibodies in acquired immunity to blood-stage parasites [19]. More detailed studies over the past 40 years have shown that immunization of human volunteers with attenuated parasites confers strong protection against experimental challenge infections [20,21].

Parasites may be attenuated at the infective sporozoite stage which invades hepatocytes [20], or at the subsequent blood stage in erythrocytes [21]. Intravenous vaccination using sporozoites isolated from irradiated mosquitoes requires a demanding regimen of five doses of >100,000 parasites over a 3 month period to elicit responses that prevent parasites developing beyond the pre-erythrocytic stage [22]. It is difficult to standardize the irradiation dose that each individual parasite receives, but alternative possibilities are chemical attenuation (e.g., alkylation of parasite DNA) [23], or genetic attenuation to remove the need for treatment of each batch. Multiple gene knockouts have been identified that, in combination, cause parasites to arrest development within the liver and elicit pre-erythrocytic stage immunity [24,25]. Engineering parasites to attenuate at the blood stage would require modification of metabolism to allow growth in selective laboratory media but not in normal human blood, as illustrated by parasites that have lost the apicoplast organelle under selection in culture [26,27].

**Figure 1.** Haploid *Plasmodium falciparum* genome containing 14 chromosomes. The whole genome of 23 Mb contains ~5500 protein-coding genes. Subtelomeric regions shaded orange are highly divergent in sequence and gene content among species; the core genome shaded blue has a high level of synteny among *Plasmodium* species, although some genes are species-specific. Loci encoding the 15 antigens incorporated in reported clinical vaccine trials are shown. Only six of these have clear orthologs in all *Plasmodium* species (shown in bold font). The full names of each of these antigens are given in **Box 2**. A region of chromosome 10 is enlarged on the bottom right of the figure to show a cluster of antigen-encoding genes including several that are highly polymorphic (most are members of the *msp3*-like gene family flanked by unrelated antigen genes at both ends of the cluster).
Although there is a precedent for use of live-parasite vaccines to prevent protozoan infections of domestic animals [28], subunit vaccines containing specific antigens are generally considered more feasible for broad delivery to human populations [4,5]. These may be constructed as recombinant proteins (that are either soluble, or form virus-like particles, or chemically conjugated nanoparticles), long synthetic peptides, recombinant plasmid DNA, or recombinant viral vectors, each potentially offering a product profile that could be incorporated into an Expanded Program for Immunization (EPI) schedule [29]. The most advanced candidate developed to date is the RTS,S vaccine, which includes a large portion of the primary sequence of P. falciparum CSP (circumsporozoite protein) – see Box 2 for abbreviations and names of all malaria antigens subsequently referred to) sequence co-expressed with the hepatitis B surface antigen as a virus-like particle, delivered with a potent proprietary adjuvant (AS01). This shows safety and short-term protective efficacy of 20–50% against malaria infection and disease in most clinical trial populations [30]. The experience of RTS,S and other recombinant protein vaccine development indicates the importance of adjuvant choice to elicit high-titer antibodies. Viral-vectorized vaccine delivery may be optimized through prime–boost schedules expressing malaria antigens by one recombinant virus (such as an adenovirus) in the first immunization and by an unrelated viral vector (such as a poxvirus) in boosting immunizations [31]. Some schedules incorporating particular candidate antigens show potent immunogenicity, although only a small minority of individuals have been protected from challenge infection and the determinants of efficacy need to be resolved [32]. However, for any subunit vaccine, inclusion of the most-effective target antigens is vital, and it is for their identification that parasite genomics is most important.

Box 2. Malaria parasite antigens
Abbreviated and full names of antigens with sequences incorporated in experimental vaccine trials already, and others mentioned in the article.

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Discovery of vaccine candidate antigens
Vaccine candidates based on 15 different antigens of P. falciparum have proceeded to clinical trials that have been published so far (Figure 1 and Box 2). These antigens are diverse proteins, expressed at various stages of the life cycle, and were first described during an early period of parasite molecular characterization; the sequences of these genes were published between 1984 and 2000 when most other genes of the parasite were still unknown (Figure 2). It is remarkable that there have been no reported vaccine trials incorporating any of the thousands of proteins encoded by genes first described in the reference parasite genome sequence published in 2002 [6]. It is very improbable that all promising candidate antigens were discovered ahead of the genome sequence, and some potential reasons for the lack of vaccine trials based on genomic findings are discussed below.

For the 15 vaccine candidate antigens that have undergone trials to date, there was a mean overall ‘lag period’ of 10 years from the primary publication of each antigen gene sequence until the first publication of a clinical vaccine trial incorporating a sequence of the respective antigen (Figure 2). There was a wide range in this time-window (from only 3 years for CSP, MSP1, and LSA-1, to 20 years for EBA-175), and for 11 out of the 15 antigens the period was less than 12 years. This shows that it is possible, as illustrated by most of these examples, to proceed from initial antigen gene discovery to a reported vaccine trial in less time than has already elapsed since the first malaria parasite genome publication in 2002. The absence of any published vaccine trial incorporating a new candidate antigen discovered in the genome sequence is therefore not a trivial observation. Another possible explanation is that most of the limited capacity and funding for malaria vaccine development may be committed to supporting previously identified candidates. Focus on a limited pre-existing portfolio might have unintentionally blocked new candidate development, even without proprietary or other conflicting interests which could exacerbate such an effect. A further possibility is that the increased scale of discovery promoted by genome sequence analysis has itself
inhibited the selection of candidates because there are now so many proteins to investigate that there may be increased uncertainty about the type and amount of data required for prioritization. Clearly, vaccine candidates should not continue to be prioritized for product development solely on the basis of the order in which they were discovered, and there is therefore a need for an evidence-based process to mark the candidacy of an antigen. Given the complexity of the parasite life cycle, and the diverse immune mechanisms that may be relevant, standardization will not be simple, but the following section outlines relevant evidence for inclusion.

Evidence to prioritize candidate antigens

**Profiling naturally acquired immune responses**

Studies to identify specific human immune responses associated with protection from malaria are important, particularly when responses to multiple antigens are analyzed in parallel. These involve laboratory assays of antigen-specificity of naturally acquired serum antibodies, or peripheral blood lymphocytes in culture, and correlations with incidence of clinical malaria of the sampled individuals in cohort studies of endemic populations [19,33–35]. The incidence of clinical malaria is more relevant to measure than the incidence of any detected infection by the parasite because naturally acquired immunity mainly acts to reduce the density of parasites in the blood rather than entirely preventing infection [36,37]. A separate approach has been to analyze immune responses in individuals protected by experimental immunization with attenuated parasites, screening reactivity against various parasite proteins in an attempt to implicate particular specificities [38]. The difficulty in producing high-quality recombinant antigen reagents has limited the effectiveness of protein microarrays to scale up to the parasite proteome, and thus prioritization of antigens for analysis is ideally determined by predictions from genetic and functional studies.

**Genetic crosses and linkage group selection**

Experimental genetic crosses of parasites have been vital in identifying genes that encode targets of immunity [39], determinants of virulence [40], erythrocyte invasion [41], and drug resistance [42]. They have been most extensively applied in studies of rodent malaria parasites, particularly *P. yoelii* [40] and *P. chabaudi* [42]. Although some studies require cloning of progeny, it is also possible to analyze uncloned progeny to identify genomic regions influencing growth or survival under experimental conditions. This linkage group selection approach has identified loci encoding targets of strain-specific immunity in rodent malaria, confirming the importance of the merozoite surface protein MSP1 [39]. This approach may be widely applicable for genetic mapping of immune targets in other apicomplexan parasites, including *Eimeria* which causes coccidiosis in chickens [43].

Figure 2. 30-Year timeline of *P. falciparum* vaccine antigen sequence description and first clinical trial publication for each antigen. Dates of first description of gene sequences are shown by arrows above the timeline for each of the 15 *P. falciparum* antigens with sequences incorporated into vaccines reported in published clinical trials (arrows below the timeline indicate the first trial in each case). The color shading indicates the parasite life-cycle stage at which each of the antigens is mainly expressed: pre-erythrocytic (blue), asexual blood stage (red), sexual and zygote transmission stage (green). The nature of the vaccine constructs varied extensively, based on synthetic peptides, recombinant proteins, virus-like particles, or viral vectored systems. In some cases only a short sequence representing part of an antigen was included in the vaccine, combined with sequences from other antigens, whereas in other cases most of a primary sequence of a single antigen was incorporated. For several of the antigens, other vaccine constructs have also been designed and tested at later dates that are not shown on the scheme. The date of publication of the first parasite genome sequence in 2002 is marked by a broken arrow above the dateline.
It is notable that some important targets of immunity are encoded by genes occurring only in *P. falciparum* or *P. vivax* but not in rodent malaria parasites, including several of the vaccine candidates already developed (Figure 1). However, the linkage group selection approach has not yet been applied to identify targets of acquired immunity in human parasites directly. Very few genetic cross experiments have been published for *P. falciparum*, and none for *P. vivax*, because the procedure requires inoculating progeny parasites from mosquitoes to splenectomized chimpanzees to allow parasites to develop to blood stages. In one study, a key role of the *P. falciparum* merozoite ligand Rh5 (which does not have an ortholog in rodent parasites) in erythrocyte invasion was identified by mapping this locus for ability to invade *Aotus* monkey erythrocytes [41], and subsequent studies to characterize this potential target of immunity are noted below. Because experimental infections of chimpanzees may no longer be conducted, owing to ethical considerations, it would be relevant to consider studies of uncloned parasite progeny in human volunteers, under safety guidelines that have already been defined for experimental infections [44]. Genetic cross experiments on human malaria parasites may also utilize an immunodeficient mouse with adoptively transplanted human hepatocytes, allowing development of parasites from mosquitoes through the liver stage, after which they may be harvested into transfused human erythrocytes [45].

Functional studies
Parasite genetic cross experiments map loci down to a few cM, typically in the order of approximately 50 kB which may contain upwards of 10 genes, after which targeted gene knockout and allelic replacement may identify causal genes. Large libraries of gene knockouts are now available for rodent malaria parasites that are most amenable to genetic manipulation [46], while targeted modification of *P. falciparum* has also undergone scale-up [47] and methodological innovation [48,49]. In conjunction with these approaches, protein–protein interaction studies have revealed functions of particular gene products, as illustrated by defining the role of Rh5 in merozoite invasion, after genetic mapping [41].

Experimental knockout of the Rh5 gene in *P. falciparum* was not achieved despite repeated attempts, suggesting that it may be essential to parasite growth at the cultured asexual blood stage [50]. Prey–bait screening of protein interactions then revealed that Basigin is the erythrocyte surface receptor for Rh5 [51], a crucial finding that led to studies on antibody inhibition of the binding [52,53] and inhibition of erythrocyte invasion [54], as well as structural analysis of the molecular interactions [55]. The relative abundance of *Rh5* transcript at the merozoite-containing schizont stage appears to vary among parasite isolates from clinical cases [56], but anti-Rh5 antibodies significantly inhibit merozoite invasion by all isolates tested so far [52]. Naturally occurring amino acid polymorphisms in Rh5 do not affect human Basigin receptor binding or antibody inhibition [52], but several mutations that alter Rh5 binding to *Aotus* monkey erythrocytes have occurred in a parasite line adapted to these hosts [57]. Another merozoite protein, termed the Rh5-interacting protein (Ripr), has been subsequently identified and is being analyzed as a potential target of immunity [58]. The prey–bait system that identified the parasite–host Rh5–basigin interaction allows broad screening, and has already identified merozoite protein MTRAP as a ligand binding to semaphorin-7A on erythrocytes, although antibodies were unable to block this particular interaction [59]. Apart from Rh5–Ripr, other key interactions between parasite proteins include a functional complex between the candidate antigen AMA1 and rhoptry neck protein 2 (RON2) at the apical tip of merozoites during erythrocyte invasion, and immunization of animals with both components has elicited more inhibitory antibodies to AMA1 than was achieved by immunization with AMA1 alone [60].

Characterizing developmental and epigenetic variation
Expression of *P. falciparum* genes at particular life-cycle stages has been described by transcriptome profiling, with particular detail for the cultured 48 h cycle of the asexual blood stages [61,62]. Apart from stage-specificity, which is largely orchestrated by a family of AP2 transcription factors [63], many genes show clonally variant expression among individual parasites at a particular stage [64] (Box 3). A survey of asexual blood stages of several laboratory clones and subclones of *P. falciparum* estimated that approximately 10% of all genes exhibit clonally variant transcription [65]. Detection of expressed protein in individual parasites also reveals clonal variation that can be hard to resolve by analysis of transcript levels in bulk-cultured populations [66].

Malaria parasite genes in subtelomeric chromosomal regions show extreme patterns of sequence diversity and clonal expression variation, particularly the var gene family with approximately 60 copies in each haploid *P. falciparum* genome encoding variants of the parasite-infected erythrocyte membrane protein 1 (PfEMP1) [6]. The antigenic repertoire of PfEMP1 is too large to allow development of a multivalent vaccine to cover the existing diversity, but two lines of investigation support the concept of a vaccine based on a small subset of antigenic types. First, parasites which adhere to the receptor chondroitin sulfate A (CSA) on placental capillary endothelia in pregnant women express a particular variant of PfEMP1, termed VAR2CSA, that is encoded by only one or a few similar var gene types in each parasite genome [67,68], although there is considerable antigenic diversity due to allelic polymorphism within populations. A vaccine which either covered VAR2CSA polymorphism within a multi-allelic formulation or which focused responses on conserved epitopes might protect against pregnancy-associated malaria, and this prospect has led to structural studies to design proteins based on particular domains eliciting antibodies that inhibit receptor binding [69,70]. A second aspect of var gene specialization is that the expression of particular genes is associated with severe clinical forms of malaria in children [71–73], probably due to variant PfEMP1 products causing infected erythrocyte cytoadhesion to particular capillary endothelial surfaces [74,75] or enhanced rosetting of uninfected erythrocytes around infected erythrocytes [76]. Antigenic
Box 3. Antigen gene polymorphism and expression variation

Some parasite antigens are encoded by a single gene with allelic polymorphism, and others are encoded by different paralogous genes that can switch transcriptional expression (Figure 1).

![Antigenic polymorphism and variation](image)

- One locus
- Allelic polymorphism
- Expression fixed

- Multiple loci
- Minimal allelic polymorphism
- Expression switching

Figure 1. Parasites are depicted here as oval-shaped cells expressing a single antigen type on the surface, encoded by a gene at a particular chromosomal locus marked by a similar shape which is enlarged for visibility. Transcriptionally active loci are indicated with an asterisk. For absolute schematic simplicity, no cellular details (e.g., nucleus) are shown. (A) Allelic polymorphism of a single locus with fixed expression (as for many malaria vaccine candidates such as CSP, TRAP, MSP1, and AMA1). (B) Variation due to alternative expression of different loci with minimal allelic polymorphism (as for malaria parasite antigens including some of the EBA and Rh ligands using alternative erythrocyte receptors). (C) Variable expression of highly polymorphic loci contributing to a very extensive potential repertoire of diversity (as for parasite genes encoding PfEMP1, and for some of the CLAG/RhopH1 and MSP3-like proteins).

cross-reactivity among candidate PfEMP1 sequences is being studied towards identifying a subset of candidate products for a multivalent vaccine against severe malaria [77].

Other large subtelomeric gene families in *P. falciparum* termed rifin and stevor encode proteins that are variably expressed on infected erythrocytes and merozoites, although whether antibodies to these are protective is not

![MSPDBL2 expression](image)

Figure 3. Example of variation in expression of a malaria parasite antigen. The *P. falciparum* MSPDBL2 protein is encoded by an msp2-like gene expressed on merozoites contained in mature blood-stage schizonts. The expression is variable among individual parasites because most schizonts are negative whereas a minority are positive. The figure shows example data from one parasite clone (HB3) (reproduced with modification from [66]). (A) A single microscopy field showing fixed parasite-infected erythrocytes. The left panel shows immunofluorescence of schizonts that react with an anti-MSPDBL2 antibody; the right panel shows DAPI (*4,6-diamidino-2-phenylindole*)-stained parasite DNA in the cells (schizonts having multiple blue clustered nuclei). (B) Proportions of mature schizonts positive for MSPDBL2 vary among different subclones of the HB3 parasite. The eight subclones shown were cultured separately for several weeks, and then analyzed by microscopy counting of hundreds of mature schizonts from each to determine the percentage positive (with 95% confidence intervals).
as well established as for PfEMP1 [78,79]. Merozoite proteins encoded by smaller gene families are also variably expressed, including several encoded by msp3-like genes clustered in a non-subtelomeric position of the genome [66] (Figure 1). For one of these proteins, MSPDDBL2, only a minority of mature merozoite-containing schizonts within any parasite line are positive, and Figure 3 illustrates the variability among cultured subclones of a single parasite clone. Among members of the EBA and Rh families of merozoite proteins that bind alternative erythrocyte receptors for invasion, some reciprocal patterns of expression and function have been shown. In particular parasite clones, experimental knockout of the gene encoding the ligand EBA175 (which binds to glycoporphin A on the erythrocyte surface) leads to increased expression of the alternative ligand Rh4 (which binds to complement receptor 1 on the erythrocyte surface), supporting a concept of vaccine design to incorporate both of the alternative ligands [80]. These candidates might be combined along with other alternatives, such as EBL-1 that is expressed in some parasites to bind to glycoporphin B, or EBA140 that binds to glycoporphin C [81], or alongside more broadly required merozoite proteins such as Rh5 [54] or its adjunct binding partner Ripr [80,82]. Several of these parasite ligands also contain allelic sequence polymorphisms that have potential adaptive significance (Box 3).

**Detecting signatures of natural selection on antigens**

Pathogen antigens are likely to be subject to frequency-dependent selection from acquired immune responses, and particular patterns of allelic polymorphism may reveal departures from neutral expectations as a consequence. Several antigens already considered to be malaria vaccine candidates first showed such evidence [83], encouraging prospects that the identification of genes with similar patterns may reveal previously unidentified targets of immunity. Genome-wide scans of allele-frequency distributions in endemic West African populations have implicated particular genes as being under balancing selection, with alleles having more-intermediate frequencies than expected in the absence of selection, and in contrast with most of the genome [66,84] (Figure 4). This was particularly apparent for many genes with peak expression at the invasive merozoite stage, including members of the eba, clag/RhopH1, surfin, and msp3-like gene families, supporting the idea that this extracellular stage is particularly susceptible to acquired immunity [66,84]. Interestingly, a few of the antigen genes that are apparently under balancing selection may also have been under recent directional selection in local populations [85–87], suggesting that new alleles are occasionally introduced into the repertoire by mutation or immigration, and rapidly increase in frequency before being checked by allele-specific acquired immunity.

Genome sequences for the small number of *P. vivax* strains published to date indicate that this species is more polymorphic than *P. falciparum*, with nucleotide diversity being approximately twice as high, suggesting that this parasite has been widespread in humans for longer [88]. Analyses of *P. vivax* candidate antigen gene sequences in population surveys implicate ama1, trap, and dbp as being under balancing selection [89], consistent with results for their *P. falciparum* homologs [83]. To allow a provisional genome-wide scan for evidence of selection in parasite species for which there are only a few whole-genome sequences, information on polymorphism may be analyzed alongside comparative data on divergence between species. Contingency tests on polymorphism-versus-divergence ratios include the McDonald–Kreitman (MK) test for skew in the ratios of nonsynonymous and synonymous single-nucleotide polymorphisms (SNPs) and interspecies fixed differences, and the Hudson–Kreftman–Aguade (HKA) test of the ratio of pairwise nucleotide polymorphism versus interspecies divergence, which may be compared across genes. This is illustrated for data on *P. falciparum* polymorphisms among a small number of laboratory lines compared with fixed differences between *P. falciparum* and the chimpanzee parasite *P. reichenowi* [10] (Figure 5). Genes with outlying values for MK and HKA ratios were largely independent, and known antigen genes more commonly had only high HKA ratios.

Genome-wide scans for loci under selection have not yet incorporated detailed analyses of repeat sequences, subtelomeric chromosome regions, or other loci with highly divergent alleles. Such sequences are not reliably characterized by short-read mapping to reference genomes and are therefore omitted from most analyses. However, advances in longer-read sequencing and methods of *de novo* assembly may soon capture the substantial
Polymorphism that exists in these regions [90], highlighting the need for suitable population genetic analytical methods for such loci, where processes of mutation and recombination do not fit simple models.

The predictive power of analyses to identify patterns of selection on antigen genes should be evaluated by epidemiological and experimental validation of the putative targets. In epidemiological studies, large cohorts of individuals need to be followed in order for levels of naturally-acquired antibodies against an array of different recombinant proteins to be tested for correlations with reduced prospective risk of clinical malaria [33–35]. To test for activity against blood-stage parasites in culture, available assays include antibody inhibition of erythrocyte invasion by merozoites [82], antibody opsonization enabling phagocytosis of merozoites [91], and antibody-dependent cellular inhibition (ADCI) of growth in the presence of monocytes [92]. Allele-specific inhibition has been demonstrated for antibodies to P. falciparum AMA1, an N-terminal part of MSP1, and P. vivax DBP [93–96], supporting the prediction that signatures of balancing selection on these merozoite antigens are driven by naturally acquired immune responses.

Towards vaccine formulation

Design of malaria vaccines should incorporate discoveries from the above areas of research, although there remains no standardized means or structure for capturing these. Different alternative approaches will remain potentially viable. In particular, vaccines designed to prime immune responses in infants to specific targets of naturally-acquired immunity need to either focus immune responses on conserved parts of the antigens or to elicit polyvalent responses that cover the natural antigenic diversity. If polymorphisms are clustered in regions of a primary sequence, it may be possible to use a relatively conserved part of the sequence, an approach that has been followed to focus on a C-terminal sequence of MSP3 of P. falciparum as a vaccine component [92]. When polymorphisms are broadly interspersed with conserved parts of sequences, for example within AMA1 or the P. vivax DBP, such dissection to remove polymorphic sites may not be possible, although particular formulations may still induce strong antibody responses to conserved epitopes [95,97]. Alternatively, multiple allele-specific antibodies may be generated to give protection against the range of allelic types circulating within populations, a ‘diversity-covering’ approach used for several vaccine formulations that have reached clinical trials, based on MSP1 [98], MSP2 [99], and AMA1 [100], as well as allelic combinations of both MSP1 and AMA1 together [101,102].

Antigenic diversity in relation to vaccination has been intensively studied for AMA1, in which the particular importance of a highly polymorphic amino acid position has been shown in several studies, including a clinical trial demonstrating allele-specific protection [103,104]. Experimental animal antibody inhibition of erythrocyte invasion in culture indicates that a multivalent vaccine containing between three and five different allelic types of AMA1 would cover most of the epitope polymorphism in this antigen [93,94]. For the polymorphic N-terminal region of MSP1, polyvalent hybrid proteins have been constructed that induce a wide range of antibody specificities which recognize the diverse natural allelic sequences [105,106]. In the cases of other antigens (the C-terminal region of MSP1 and the near-full length of MSP2 separately), for which known allelic sequences group into dimorphic allelic classes, two allelic types have been incorporated into formulations for clinical trial [98,99,101].

There remains no consensus template for an effective malaria vaccine, but the success of multivalent vaccines against viral and bacterial infections supports the rationale to characterize individual targets of immunity in detail to assist the design of subunit vaccines. Engineering of more effective immunogens is vital, and should involve modeled protein design to focus responses on important epitopes [107], as well as the use of the most-effective delivery systems [108]. Application of these principles, combined with lessons from empirical studies of a small number of malaria candidate antigens to date, should enable genome-scale studies on the parasite to be more systematically translated into the development of malaria vaccines.

Figure 5. Genome-wide scan of gene polymorphism in P. falciparum versus divergence with the chimpanzee parasite P. reichenowi. Each point represents comparative data for two different indices (HKA ratio and MK skew) for a gene, with sequence data from five P. falciparum laboratory lines compared with a single P. reichenowi genome (reproduced with modification from reference [10]). Several known antigen genes are labeled for illustration. The HKA ratio for each gene (on the x axis) is the average pairwise nucleotide polymorphism in P. falciparum divided by the average interspecies nucleotide divergence (K), with a broken line shown at an arbitrary ratio of 0.15 to visually indicate genes with highest relative levels of polymorphism on the right of the plot. The MK skew for each gene (on the y axis) is the log-transformed odds ratio from the 2 x 2 table of numbers of nonsynonymous and synonymous polymorphisms within P. falciparum over the numbers of nonsynonymous and synonymous interspecies fixed differences. An MK skew value of zero represents an odds ratio of 1.0 reflecting where the intraspecific and interspecific ratios do not differ. Positive MK skew values indicate ratios for genes reflecting an excess of nonsynonymous polymorphism or deficit of nonsynonymous fixed differences compared with neutral expectations, marking in red those that are significant with Fisher’s exact tests. Three of these genes that also have high HKA ratios are shown in bold outline (including AMA1). Some genes are not plotted here as they have infinite MK skew values (although significant in a number of cases by Fisher’s exact test), mostly due to having no synonymous polymorphisms. Abbreviations: HKA, Hudson-Kreitman-Aguade test; MK, McDonald–Kreitman test.

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