Rare Alleles and Signatures of Selection on the Immunodominant Domains of Pfs230 and Pfs48/45 in Malaria Parasites From Western Kenya

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Background: Malaria elimination and eradication efforts can be advanced by including transmission-blocking or reducing vaccines (TBVs) alongside existing interventions. Key transmission-blocking vaccine candidates, such as Pfs230 domain one and Pfs48/45 domain 3, should be genetically stable to avoid developing ineffective vaccines due to antigenic polymorphisms. We evaluated genetic polymorphism and temporal stability of Pfs230 domain one and Pfs48/45 domain three in Plasmodium falciparum parasites from western Kenya.

Methods: Dry blood spots on filter paper were collected from febrile malaria patients reporting to community health facilities in endemic areas of Homa Bay and Kisumu Counties and an epidemic-prone area of Kisii County in 2018 and 2019. Plasmodium speciation was performed using eluted DNA and real-time PCR. Amplification of the target domains of the two Pfs genes was performed on P. falciparum positive samples. We sequenced Pfs230 domain one on 156 clinical isolates and Pfs48/45 domain three on 118 clinical isolates to infer the levels of genetic variability, signatures of selection, genetic diversity indices and perform other evolutionary analyses.

Results: Pfs230 domain one had low nucleotide diversity (π = 0.15 × 10⁻²) with slight variation per study site. Six polymorphic sites with nonsynonymous mutations and eight haplotypes were discovered. I539T was a novel variant, whereas G605S was nearing fixation. Pfs48/45 domain three had a low π (0.063 × 10⁻²), high conservation index, and three segregating sites, resulting in nonsynonymous mutation and four haplotypes. Some loci of Pfs230 D1 were in positive or negative linkage disequilibrium, had negative or positive selection signatures, and...
INTRODUCTION

Genetic polymorphism of *Plasmodium falciparum* antigens has hampered efforts to develop an effective vaccine that is protective against pre-erythrocytic and asexual blood-stage parasites (Genton et al., 2002; Takala et al., 2007; Ogutu et al., 2009; Bergmann-Leitner et al., 2012; Neafsey et al., 2015; Ouattara et al., 2015). Recent efforts, however, have been made to develop vaccines that reduce and block *Plasmodium falciparum* transmission at the community level. Two of the existing transmission-blocking vaccine (TBV) candidates, *P. falciparum* surface protein 230 (Pfs230) (Sabeti et al., 2007; Lee et al., 2019, 2020; Singh et al., 2019, 2020; Tachibana et al., 2019; Huang et al., 2020; Healy et al., 2021) and *P. falciparum* surface protein 48/45 (Pfs48/45) (Singh et al., 2019, 2021; Lee et al., 2020) have been shown to elicit antibody responses in mice and people that block *P. falciparum* gametocyte fertilization in the mid-gut of the *Anopheles* vector.

Pfs230 is a cysteine-rich 230 kDa protein expressed by both male and female gametocytes (Rener et al., 1983; MacDonald et al., 2016). The antigen is thought to play a role in gamete fusion in the mosquito blood meal after forming a complex with another cysteine-rich protein, Pfs48/45 (Eksi et al., 2006). In comparison to antibodies elicited by immunization with other Pfs230 domains, Domain 1 (D1) has been shown to elicit transmission-blocking monoclonal antibodies with strong inhibitory activity against oocyst development in standard membrane feeding assays (Lee et al., 2019; Singh et al., 2019, 2020; Tachibana et al., 2019; Huang et al., 2020; Healy et al., 2021). Like Pfs230 D1, fusion with its counterpart Pfs48/45 D3 has good potential as a component of a TBV. The latter fused doublet antigen consists of three domains linked by disulphide bonds and contains 16 cysteine residues (Kocken et al., 1993; Lennartz et al., 2018). Unlike Pfs230, Pfs48/45 is anchored on the gamete surface membrane by glycoprophatidylinositol (Kocken et al., 1993; Dijk et al., 2001; Gilson et al., 2006; Lennartz et al., 2018) and is essential for male gamete fertility. Domain 3 has been shown to elicit antibodies in the host (Graves et al., 1988; Roeffen et al., 1994; Dijk et al., 2001; Bousema et al., 2010; Jones et al., 2015; Acquah et al., 2017; Singh et al., 2019; Baptista et al., 2022).

Pfs48/45 D3 is located at the C-terminus of the protein and contains binding sites for non-inhibitory and inhibitory human and mouse mAbs that reduce *P. falciparum* infection in mosquitoes (Vermeulen et al., 1986; Graves et al., 1988; Ouchtourov et al., 2007; Chowdhury et al., 2009; Singh et al., 2017, 2019; Kundu et al., 2018; Lennartz et al., 2018; Lee et al., 2020).

Antigenic polymorphism of Pfs230 D1 and Pfs48/45 D3 should be assessed in malaria endemic areas on a regular basis to support the successful development of these TBV candidates due to the fact that, if the targeted regions are genetically unstable, polymorphisms may cause critical codon changes within immunogenic epitopes, thereby reducing TBV efficacy. Several dimorphic sites on Pfs230 D1 and Pfs48/45 D3 have previously been identified (Kocken et al., 1995; Drakeley et al., 1996; Escalante et al., 1998; Conway et al., 2001; Jones et al., 2015; MacDonald et al., 2016; Kundu et al., 2018; Singh et al., 2020; Coelho et al., 2021); however, there is limited knowledge of the extent of genetic diversity, signatures of selection, and other evolutionary forces that may be shaping alleles in *P. falciparum* from different malaria transmission zones. We therefore performed an in-depth genetic analysis of *Pfs230 D1* and *Pfs48/45 D3* in parasites isolated from patients with uncomplicated falciparum malaria from three different areas in western Kenya.

MATERIALS AND METHODS

Study Site and Sampling

Dry blood spots (DBS) were collected on filter paper from febrile malaria patients at health clinics in Homa Bay County (Kochia), Kisumu County (Chulaimbo), and Kisii County (Eamba) in 2018 and 2019 (Figure 1). The study site in Homa Bay is characterized by perennial transmission. Vector control consists of universal distribution of long-lasting insecticidal bed nets with annual indoor residual spraying of insecticides. The study site in Kisumu County also has perennial transmission. Vector control consists of long-lasting insecticidal bed nets (LLINs) alone. The site in Kisii County is malaria epidemic-prone with low transmission and residents use LLINs (Kapesa et al., 2018). In brief, four others (1813, 1955) and (1813, 1983) had a history of recombination. Mutated loci pairs in Pfs48/45 domain three had negative linkage disequilibrium, and some had negative and positive Tajima’s D values with no history of recombination events.

Conclusion: The two transmission blocking vaccine candidates have low nucleotide diversity, a small number of zone-specific variants, high nucleotide conservation index, and high frequency of rare alleles. With the near fixation a polymorphic site and the proximity of mutated codons to antibody binding epitopes, it will be necessary to continue monitoring sequence modifications of these domains when designing TBVs that include Pfs230 and Pfs48/45 antigens.

Keywords: Pfs230, Pfs48/45, transmission blocking vaccines, genetic diversity, evolutionary forces
drops of approximately 25 µL of blood from each patient were spotted on Whatman™ Blood Stain Cards (GE Healthcare WB100014) as previously described (Coombs and Fiscus, 2009). Each card was stored individually in silica gel-containing plastic bags before being transported to the joint International Centre of Excellence for Malaria Research (ICEMR) and Tom Mboya University College Laboratory in Homa Bay town for storage at -20 °C. 150 of the 372 DBS collected came from Homa Bay; 120 and 102 came from Kisumu and Kisii, respectively.

Amplification and Sequencing of Pfs230 Domain one and Pfs48/45 Domain three
Genomic DNA was extracted from filter paper using the modified Chelex resin (Chelex -100) method and stored at -20°C. As a positive control, DNA from the cultured laboratory strain NF54 was extracted and stored. Plasmodium species-specific real-time PCR targeting 18S ribosomal RNA gene was used to confirm P. falciparum positive DNA samples before amplification of specific target fragments of each gene (Ochwitz et al., 2021; Onyango et al., 2021). Primer sets were designed using Primer3 version 0.4.0 for Pfs230 D1 and Pfs48/45 D3 and in silico validation of each set was performed using the Sequence Manipulation Suite (Stothard, 2000). Among the 372 samples, 332 (89.3%) tested positive for P. falciparum DNA (n = 150, 120 and 62 from Homa Bay, Kisumu and Kisii, respectively) and were used to amplify Pfs230 D1 and Pfs48/45 D3 in a T100™ Thermal Cycler (Bio-Rad, Hercules, CA, United States). Briefly, 3 µL of sample DNA was added to a mixture of 11.5 µL of DreamTaq Green PCR Master Mix (2X), 0.5 µL of Pfs230 D1 forward (5′-TGGTGAAGCTGTCGAAGATG-3′) and reverse primers (5′-GTGTACCACAGGGGGAAGAG-3′) targeting 514 base pairs and 7.5 µL of double-distilled water. The thermal profile was set as follows 95 °C for 3 min, 34 cycles (94 °C for 30 s, 58 °C for 30 s and 72 °C for 45 s) and final extension at 72 °C for 6 min. For Pfs48/45 D3, similar reaction volume was prepared using forward (5′-TTTTCAAGAAGGAAAAGAAAAAGC-3′) and reverse primers (5′-GCCAAAAATCCATAATATGCTGA-3′) targeting 600bp. The PCR conditions were set as follows 95°C for 3 min, 34 cycles (94°C for 30 s, 55°C for 30 s and extension at 72°C for 45 s) final extension at 72 °C for 6 min. All the amplicons were assessed by gel electrophoresis in 1.5% w/v agarose gel before sequencing. For Pfs230 D1, 82, 39, and 35 samples from Homa Bay, Kisumu, and Kisii, respectively, were amplified. For Pfs48/45 D3, 36, 44, and 38 samples from Homa Bay, Kisumu, and Kisii, respectively, were amplified. All the PCR amplicons, together with positive controls, were purified using Exonuclease I and Shrimp Alkaline Phosphatase (ExoSAP-IT) and bi-directionally sequenced using 3730 BigDye® Terminator v3.1 Sequencing Standard kit on ABI PRISM® 3700 DNA Analyzer (Applied Biosystems, Foster City, CA, United States).
TABLE 1 | Polymorphic sites on Pfs230 domain one and Pfs48/45 domain two and three from Homa Bay, Kisumu and Kisii region in western Kenya; n: number of sequences harbouring mutations; *: reference (3D7 and NF54) allele only; D: Domain; Nsyn: Non-synonymous mutation; Syn: Synonymous mutation; A: Adenine; C: Cytosine; T: Thymine; G: Guanine.

Pfs230 (N = 156)

| Segregating Sites | Domains | Allelic Frequency | Substituted Bases | Type of substitution | Codon Change | Type of Mutation |
|-------------------|---------|------------------|-------------------|---------------------|--------------|-----------------|
|                   |         | Homa Bay n (%)   | Kisumu n (%)      | Kisii n (%)         |              |                 |
| 1616              | D1      | 2 (5.1)          | -                 | -                   | *T/C         | Transition      |
| 1813              | D1      | 81 (98.8)        | 37 (94.9)         | 35 (100)            | *G/A         | Transition      |
| 1955              | D1      | 2 (2.4)          | 1 (2.6)           | 2 (5.7)             | *C/G         | Transversion    |
| 1964              | D1      | -                 | 1 (2.9)           | -                   | *A/T         | Transversion    |
| 1967              | D1      | -                 | 1 (2.6)           | -                   | *C/A         | Transversion    |
| 1983              | D1      | 35 (42.7)        | 19 (48.7)         | 16 (45.7)           | *A/C         | Transversion    |

Pfs48/45 (N = 118)

|                   |         | Allelic Frequency | Substituted Bases | Type of substitution | Codon Change | Type of Mutation |
|-------------------|---------|------------------|-------------------|---------------------|--------------|-----------------|
|                   |         | Homa Bay n (%)   | Kisumu n (%)      | Kisii n (%)         |              |                 |
| 753               | D2      | -                 | -                 | -                   | *T/C         | Transition      |
| 757               | D2      | -                 | 4 (9.1)           | 3 (7.9)             | *A/G         | Transition      |
| 762               | D2      | 1 (2.8)           | 3 (6.8)           | 3 (7.9)             | *C/G         | Transversion    |
| 911               | D3      | 1 (2.8)           | -                 | -                   | *T/A         | Transition      |
| 940               | D3      | 3 (6.3)           | 3 (6.8)           | 9 (23.7)            | *T/A         | Transversion    |
| 979               | D3      | -                 | -                 | 1 (2.6)             | *T/G         | Transversion    |

Data Analysis

All sequences were assembled using Geneious version 11.1.5 software, and multiple sequence alignment was performed using ClustalW. Polymorphic locus and codons were inferred after comparing each sequence to the respective sequence of positive control (NF54) as well as 3D7 (PF3D7_0209000 for Pfs230 and PF3D7_1346700 for Pfs48/45). DnaSP Version 6.12.03 (Rozas et al., 2017) and Arlequin version 3.5.2 (Excoffier and Lischer, 2010) were used to compute genetic diversity indices such as nucleotide diversity (π), haplotype diversity (Hd), number of haplotypes (h), number of segregating sites (S) and mean number of pairwise difference (k). Population Analysis with Reticulate Trees (Popart) version 1.7 software (Clement et al., 2000) was used to infer haplotype networks. Neutrality tests; Tajima’s D, Fu and Li’s D, Fu and Li’s Fs and Fu’s Fs statistics and test for the presence of Recombination events (Rm) and linkage disequilibrium (LD) were computed in DnaSP Version 6.12.03 and Arlequin version 3.5.2. Generated Tajima’s D values were plotted using GraphPad version 8.3.0. Both antigen structural delineation was done using Protein Homology/analogy Recognition Engine (PHYRE2) version 2.0 and generated models visualized and edited in UCSF Chimera version 1.15 (Pettersen et al., 2004).

RESULTS

Analysis of Mutations Detected in Pfs230 D1 and Pfs48/45 D3

Six loci (1,616, 1813, 1955, 1964, 1967, and 1983) in Pfs230 D1 were found to be polymorphic, resulting in nonsynonymous mutations (Table 1). The mutations were skewed toward transversion, with a transversion to transition ratio (Tv: Ts) greater than 0.5. Two polymorphic sites were singleton (1964 and 1967), whereas four dimorphic sites (1,616, 1813, 1955, and 1983) were parsimony informative. These polymorphisms resulted in I539T, G605S, T652R, E655V, T656N, and K661N codon changes. Nonsynonymous alterations T652R and K661N were on separate beta (β) pleated sheets connected by a loop containing mutated codons E655V and T656N (Supplementary Figure S1). G605S was also on the loop connecting different β pleated sheets. In general, western Kenya parasites had a high allelic frequency of G605S (98.08%), followed by progressively lower frequencies of K661N, T652R, and I539T. E655R and T656N were each observed at a frequency of <1%. The prevalence of various alleles was almost similar across the various study sites. For example, as shown in Table 1, G605S was the most common codon change in the three study sites. Only two P. falciparum isolates from Kisumu and Homa Bay County lacked this mutation.

In contrast to the six nonsynonomous mutated sites observed in Pfs230 D1, Pfs48/45 D3 had three segregating sites (Table 1). Singleton sites were found at loci 911 and 979 in parasites isolated from patients residing in Homa Bay County and Kisii County, respectively. A low frequency polymorphism at locus 940 was observed across parasite populations in all three counties, and was parsimony-informative. These transition bias mutations at loci 911, 940, and 979 resulted in nonsynonymous mutations V304D, L314I, and C327G, respectively. The variants were in the Pfs48/45 D3 antigen loop connecting different β pleated sheets (Supplementary Figure S1). Codon change C327G in D3 was found in only one sequence in parasites isolated from a patient in Kisii County (Table 1). The ability of the designed primer set to cover Pfs48/45 D3 also allowed for the discovery of a singleton site 753 (Y251Y) and parsimony-informative sites 757 and 762 (K253E and N254K) (Table 1; Supplementary Figure S1). Except for G605S, which was near dimorphic codons on
Pfs48/45 domain 2, the superimposed structure revealed dimorphic codons of Pfs48/45 D3 antigen close to those of Pfs230 D1 (Supplementary Figure S2).

**Genetic Diversity of Pfs230 and Pfs48/45 Genes in Western Kenya**

Pfs230 D1 from the three sites had \( \pi \) of 0.15 × 10\(^{-2} \), \( k \); 0.68 and haplotype diversity (Hd) of 0.57 (Table 2). The domain had a nucleotide conservation index of 98.7% with a total of eight haplotypes circulating in western Kenya (Figure 2). Kisumu had the highest \( \pi \) (0.18 × 10\(^{-2} \)) followed by Kisii (0.15 × 10\(^{-2} \)) and Homa Bay (0.12 × 10\(^{-2} \)). The site also had the most haplotypes 6) and the highest Hd (0.63) when compared to Kisii and Homa Bay, which had four haplotypes each and Hd of 0.60 and 0.52, respectively (Figure 2). Haplotype 2 (Hap_2) with the mutated codon G605S was the most common in western Kenya and at each study site. This was followed by Hap_4 (mutated codons G605S and K661N), Hap_7 (G605S, T652R, and K661N), and Hap_8 (T652R and K661N) (Supplementary Table S1). The remaining haplotypes (Hap_1, Hap_3, Hap_6, and Hap_5) were observed at a lower frequency. Only one sequence (from Chulaimbo) in western Kenya lacked a mutated site and had 100% sequence identity to the laboratory strain PF3D7_0209000 or NF54 sequence used as a positive control (Figure 2).

Pfs48/45 D3 from western Kenya had low \( \pi \) (0.06 × 10\(^{-2} \)) and Hd (0.25) (Table 2). The domain had a conservation index of 99.3%, with four haplotypes circulating in the study area (Figure 2). Kisii parasites had the highest \( \pi \) (0.10 × 10\(^{-2} \)) followed by Homa Bay (0.05 × 10\(^{-2} \)) and Kisumu (0.03 × 10\(^{-2} \)) (Table 2). Kisii and Homa Bay study sites each had three haplotypes in circulation, while Kisumu had four (Figure 2). The majority of haplotypes lacked a mutation (Hp_1) or had 100% sequence identity to the laboratory strain PF3D7_1346700 or the NF54 sequence used as a positive control (85.6%). This was followed by Hp_3 (mutated codon L314I), which had an overall frequency of 12.7%.

**TABLE 2 | Summary of genetic diversity indices for Pfs230 domain one and Pfs48/45 domain three from parasites in western Kenya N: Sample size; C: Conservation index; S: Segregating sites; \( \pi \): nucleotide diversity; Vars: Variants; Hd: Haplotype diversity.**

| Pfs230 D1 Region | N  | C (%) | S   | \( \pi \) (×10\(^{-2} \)) | h  | Tajima’s \( D \) | Fu’s \( D \) | FLD* | FLF* |
|------------------|----|-------|-----|--------------------------|----|---------------|------------|------|------|
| Homa Bay         | 82 | 99.40 | 3   | 0.12                     | 4  | -0.11         | -0.21      | -0.54 | -0.45 |
| Kisumu           | 39 | 98.90 | 5   | 0.18                     | 6  | 0.60          | -0.80      | -0.73 | -0.82 |
| Kisii            | 35 | 99.40 | 3   | 0.15                     | 4  | 0.60          | 0.80       | -0.94 | -0.29 |
| W. Kenya         | 156| 98.70 | 6   | 0.14                     | 8  | 0.56          | -0.82      | -3.33 | -1.03 |

| Pfs48/45 D3      |    |       |     |                         |    |               |            |      |      |
|------------------|----|-------|-----|-------------------------|----|---------------|------------|------|------|
| Homa Bay         | 36 | 99.50 | 2   | 0.05                     | 3  | 0.21          | -1.09      | -1.42 | -0.80 |
| Kisumu           | 44 | 99.80 | 1   | 0.03                     | 2  | 0.13          | -0.60      | -0.30 | 0.55  |
| Kisii            | 38 | 99.50 | 2   | 0.10                     | 3  | 0.41          | -0.21      | -0.12 | -0.81 |
| W. Kenya         | 118| 99.30 | 3   | 0.06                     | 4  | 0.25          | -0.94      | -1.87 | -2.06 |

**FIGURE 2 | TCS-network analysis of the relationship of haplotypes based on the the Pfs230 domain one and Pfs48/45 domain three from parasites in malaria-endemic and epidemic-prone regions of western Kenya. (A) represents haplotypes based on observed sequence variation in Pfs230 D1 whereas (B) represents haplotypes in Pfs48/45 D3. The network shows the distribution of haplotype within malaria-endemic Homa Bay (red) and Kisumu (Chulaimbo) (green) as well as epidemic Kisii highland (purple). The hatch marks represent the number of mutations resulting in a specific haplotype whereas the size of the circle equates to the frequency of the observed haplotypes. Haplotypes labelled Hap_1 (NF54) for Pfs230 D1 and Hap_1 (NF54) for Pfs48/45 D3 lacked mutated locus and have 100% sequence identity to laboratory strain NF54 or 3D7. Nucleotide sequences of all haplotypes were submitted to GenBank (accession: MW624857- MW625101).**
while the rest (Hp_2 and Hp_4) had a frequency <1% (Supplementary Table S2).

**Signatures of Selection, Linkage Disequilibrium and Recombination Events**

*Pfs230* D1 from *P. falciparum* isolates deviated from a standard neutral model. Tajima’s D (-0.8), FLD* (-0.9) and FLF* (-1.0) tests were all negative and non-significant (*p > 0.05*) (Table 2). However, Fu’s Fs result (-3.3) was significant (*p = 0.023*). Tajima’s D test results were also non-significant (*p > 0.05*) in each site (Table 2). Despite the overall negative Tajima’s D results, there was a slight variation among individual mutated loci on *Pfs230* D1. Locus 1983 (codon change K661N) had a significant (*p < 0.05*) positive (1.9) Tajima’s D result, whereas the rest had negative results (Figure 3). Apart from deviating from a standard neutral model, some loci pairs (1813, 1955) had positive LD (D’) results with highly significant (*p < 0.001*) \( \chi^2 \) values of 19.7 and 12.7 among Homa Bay and Kisumu sequences, respectively (Supplementary Table S1). Among Kisii sequences, loci pairs (1813 and 1983) had positive D’ results, despite the fact that the \( \chi^2 \) values was non-significant and the \( r^2 \) value was low (0.03). Other loci with positive D’ results, low \( r^2 \) values and non-significant (*p > 0.05*) \( \chi^2 \) values in Homa Bay and Kisumu parasites include loci pairs (1813,1983) (1955,1983) and (1967,1983). Some loci pairs on Kisu and Kisii sequences had negative D’ results (Supplementary Table S1). On *Pfs230* D1 from parasites in Homa Bay and Kisumu, recombination events were detected across loci pairs (1813, 1955) and (1967, 1983) (Supplementary Table S1).

All of the *Pfs48/45* D3 sequences had non-significant (*p > 0.05*) negative Tajima’s D -1.9, FLD*: 2.1 and FLF*: 2.0 results. The Fu’s Fs (-1.9) result was, however, significant (*p = 0.096*). Locus 940 (13141) among Kisii sequences had a significant (*p > 0.05*) positive Tajima’s D (0.8) results (Figure 3). There was no evidence of Rm or positive D’ results at any of the dimorphic loci within *Pfs48/45* D3. However, some loci pairs from Homa Bay and Kisii had negative D’ results (Supplementary Table S2).

**DISCUSSION**

The *Pfs230* D1 and *Pfs48/45* D3 antigens are important candidate antigens in the development of an effective TBV. Despite having a high frequency of rare alleles in western Kenya, both targets had low nucleotide diversity. Two variants, each on *Pfs230* D1 and *Pfs48/45* D3, were novel and private to western Kenya. The study validated five previously described polymorphic sites on *Pfs230* D1 (Singh et al., 2020). In this study, G605S, one of the five mutated codons, was fixed in some study areas but not in others. *Pfs230* D1 had the most mutations, while the *Pfs48/45* D3 was the most conserved. Mutated loci from both domains were either under purifying or balancing selections. Other genetic forces revealed to have shaped alleles on the two genes included inbreeding and genetic drift with recombination being discovered only *Pfs230* D1.

*Pfs230* D1 from western Kenya had low nucleotide diversity, with significant Fu’s Fs results indicating a high frequency of rare alleles. In addition to the previously reported 15 polymorphisms on *Pfs230* D1 from parasites in Asian (Bangladesh, Cambodia, Laos, Myanmar, Thailand, and Vietnam) and African (Democratic Republic of the Congo, Ghana, Guinea, Malawi, Mali, Nigeria, Senegal, and Gambia) countries (MacDonald et al., 2016; Singh et al., 2020), this study discovered one additional mutation (I539T). This novel variant was identified only at the Kisumu study site along with five other polymorphisms (G605S, T652R, E655V, T656N, and K661N). These findings validate five previously described polymorphisms reported by (Singh et al., 2020). We speculate that the relatively higher nucleotide diversity index and number of haplotypes in Kisumu compared to other sites in western Kenya is related to the region’s slightly higher malaria transmission and absence of IRS activities (Oduma et al., 2021; Ochwedo et al., 2022).

Missense mutation G605S was found in parasites at a slightly higher allelic frequency (AF = 0.98) than in other geographical regions, as described by (Singh et al., 2020) (AF = 0.94) (MacDonald et al., 2016), (AF = 0.11), and (Coelho et al., 2021) (AF = 0.91). With only two clinical isolates in western Kenya lacking this mutation, G605S is almost completely fixed.
This indicates the presence of selection pressure from either host antibodies, vector immune response or genetic drift (decreased variation and increasing homozygosity), may be stronger on \( P. falciparum \) populations from Kisii (low parasite population size) or Homa Bay (endemic site with declining parasite population size) compared to Kisumu (Chulaimbo) (Hancock and Di Rienzo, 2008; Honnay, 2013; Oduma et al., 2021). In contrast, the second most common polymorphism, K661N, was found in Kisumu at a higher frequency than in Kisii and Homa Bay. This reversal in the observed G605S and K661N frequencies could be attributed to factors such as recombination events (Rm), which are known to interfere with linked loci and could be effective on linked dimorphic loci pair 1813 and 1983 (responsible for G605S and K661N mutations respectively), thus increasing diversity in Kisumu (Chulaimbo) (Mejia, 2012) as opposed to Kisii and Homa Bay parasites, which also lack Rm between the two sites. Since immunogenic epitope binding light chain of \( mAb \) is close to the dimorphic codon G605, selection pressure from host antibodies on the epitope may be affecting the surrounding codons (MacDonald et al., 2016; Singh et al., 2020). This codon is located within a disulphide loop (from 593 to 611) that is thought to be stabilizing the epitope binding of TB 4F12 mAb (Singh et al., 2020). With near-complete fixation, the mutation may be beneficial to parasites but have a negative effect on the epitope binding affinity of TB 4F12 mAb. This needs to be looked into further by immunoadsays of haplotypes with this polymorphism. Polymorphism I539T was found near codons 542–592 that contain 3G2 and 5G3 mAb binding epitopes which were previously shown to have no detectable oocyst reduction activity (Singh et al., 2020). Other polymorphisms, T652R, K661N on different \( \beta \) pleated sheets, and E655V, T656N on disulphide loops linking the two loops, were distally located from the epitope that binds TB 4F12 mAb (Singh et al., 2020). When the two fusion proteins were superimposed, these four polymorphic codons were closer to mutated codon V304D, L314I, and C327G on \( Pfs48/45 \) D3, supporting the hypothesis that antibodies could be sterically interfering with protein-protein interaction (Singh et al., 2020). \( Plasmodium falciparum \) may induce these mutations in response to antibody-induced pressure in order to circumvent the blockade of fusion between \( Pfs230 \) D1 and \( Pfs48/45 \) D3, resulting in an uninterrupted gametocyte fertilization process.

The novel missense polymorphism C327G on \( Pfs48/45 \) D3 has the potential to be very important because it can interfere with one of the six cysteine residue pairings (pairing between codon C298 and C327) on the 85RF45.1 mAb epitope (Kundu et al., 2018). Other polymorphisms (Y251Y, K253E, N254K on \( Pfs48/45 \) D2 and V304D, L314I in \( Pfs48/45 \) D3) have been observed in \( P. falciparum \) populations in other malaria endemic regions (Conway et al., 2001; Jones et al., 2015; Kundu et al., 2018). However, none of these polymorphisms had been previously reported by a study conducted in the Asembo Bay area of western Kenya (Escalante et al., 1998). Though not the focus of this study, polymorphisms on codon 254 is thought to influence the type of host antibody that binds to the epitope bearing this mutation on \( Pfs48/45 \) antigen (Köcken et al., 1995).

The three polymorphic codons Y251Y, K253E, and N254K on \( Pfs48/45 \) D2, are close to the disulphide loop that stabilizes the epitope binding TB 4F12 mAb on \( Pfs230 \) D1, thus suggesting steric interference from the antibodies. \( Pfs48/45 \) domain three is highly conserved, with low nucleotide and haplotype diversity when compared to \( Pfs230 \) D1. The key polymorphism based on this domain was L314I, which has a higher allelic frequency in Kisii highlands than in Homa Bay and Kisumu. Despite the presence of a high frequency of rare alleles, the majority of parasites lacked polymorphic loci on \( Pfs48/45 \) D3.

Inbreeding, recombination, and natural selection were identified as major drivers of the observed mutations in \( Pfs230 \) D1 and \( Pfs48/45 \) D3. The presence of linkage disequilibrium confirmed the history of selection pressure and inbreeding across various loci in \( Pfs230 \) D1 and \( Pfs48/45 \) D3 (Larrañaga et al., 2013). Some polymorphisms were considered intermediary because they had negative linkage disequilibrium (\( D' \)) values (Silvela et al., 1999). The negative \( D' \) values also confirmed a history of random drift, which is decreasing the number of variants while increasing homozygosity that may play a role in the parasite’s loss of favourable mutations if it persists (Barton, 2010).

The presence of natural selection was confirmed by the Tajima’s \( D \) values. Overall negative Tajima’s \( D \) results revealed that purifying selection was affecting the majority of loci within \( Pfs230 \) D1 and \( Pfs48/45 \) D3, reducing genetic diversity (Cvijović et al., 2018). The aforementioned selection was, however, weak because the computed negative Tajima’s \( D \) values in both antigens were not significant. Individual Tajima’s \( D \) results for each codon revealed all other dimorphic codons to be under purifying selection, with the exception of K661N on \( Pfs230 \) D1 from all study sites and V304D on \( Pfs48/45 \) D3 from Kisii, which are under strong and weak balancing selection, respectively. The two mutated loci under balancing selection may play an important role within the \( Pfs230 \) D1 and \( Pfs48/45 \) D3 fusion proteins, which may explain why they are maintained in the \( P. falciparum \) population from western Kenya (Escalante et al., 1998). These findings support previous postulation (Jones et al., 2015) that selection pressure is acting on immunogenic domains of \( Pfs48/45 \).

The presence of weak purifying selection acting on dimorphic sites may impact not only host mAb binding and functional activity but also be affected by selective pressure in the mosquito vector (Lombardo and Christophides, 2016). This pressure could be exerted on individual antigens before or after complex formation. Findings in the present study support future investigations that examine functional antibody responses such as the ability of \( Pfs230 \) and \( Pfs48/45 \) antibodies that activate human plasma complement and reduce mosquito infectivity in membrane feeding assays.

**CONCLUSION**

The \( Pfs230 \) D1 and \( Pfs48/45 \) D3 in \( P. falciparum \) from western Kenya have low nucleotide diversity and a high conservation index with high frequency of rare alleles. Among the observed polymorphisms in \( Pfs230 \) D1, G605S is nearly fixed in the population. Natural selection, inbreeding, and, to some extent, recombination are important driving forces in shaping these alleles in the two antigens. With the discovery of novel polymorphic sites, the two domains of the \( Pfs230 \) and \( Pfs48/45 \)
45 from different malaria-prone regions, including areas where clinical trials have been conducted, should be monitored indefinitely. This will help track the genetic stability of the two TBV candidates.

DATA AVAILABILITY STATEMENT

The datasets presented in this study can be found in online repositories. The names of the repository/repositories and accession number(s) can be found in the article/Supplementary Material.

ETHICS STATEMENT

The studies involving human participants were reviewed and approved by the Maseno University Ethics Review Committee (MUERC protocol No. 00456) and the University of California, Irvine Institutional Review Board (HS#2017 – MUERC protocol No. 00456) and the University of California, approved by the Maseno University Ethics Review Committee. The studies involving human participants were reviewed and approved by the Maseno University Ethics Review Committee. The datasets presented in this study can be found in online repositories. The names of the repository/repositories and accession number(s) can be found in the article/Supplementary Material.

AUTHOR CONTRIBUTIONS

KO conceptualised and designed the study, collected data, carried out the experiments, curated and analyzed the data, and drafted, edited, and revised the final manuscript. FA carried out the experiments and revised the final manuscript. WO collected data and revised the final manuscript. EM carried out the experiment and revised the final manuscript. ID revised the final manuscript. SO collected data, PO collected data, HA administration, SO administration and revised the final manuscript, AO revised the final manuscript. WM revised the final manuscript. AG revised the final manuscript. M-CL drew the map. GY developed the study’s concept and revised the final manuscript. DZ developed the concept, carried out the experiment, curated the data, and revised the final manuscript. JK developed the study’s concept and revised the final manuscript.

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SUPPLEMENTARY MATERIAL

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fgene.2022.867906/full#supplementary-material

REFERENCES

Acquah, F. K., Obboh, E. K., Asare, K., Boampong, J. N., Nuvor, S. V., Singh, S. K., et al. (2017). Antibody Responses to Two New Lactococcus lactis-Produced Recombinant Pf64/45 and Pf630 Proteins Increase with Age in Malaria Patients Living in the Central Region of Ghana. Malar. J. 16 (1 16), 1–11. doi:10.1186/s12936-017-1955-0

Baptista, B. O., de Souza, A. B. L., Riccio, E. K. P., Bianco-Junior, C., Totino, P. R. R., Martins da Silva, J. H., et al. (2022). Naturally Acquired Antibody Response to a Plasmodium falciparum Chimeric Vaccine Candidate GM22.6c and its Components (MSP-3, GLURP, and Pf64/45) in Individuals Living in Brazilian Malaria-Endemic Areas. Malar. J. 21, 6. doi:10.1186/s12936-021-04020-6

Barton, N. H. (2010). Mutation and the Evolution of Recombination. Phil. Trans. R. Soc. B 365, 1281–1294. doi:10.1098/rstb.2009.0320

Bergmann-Leitner, E. S., Duncan, E. H., Mease, R. M., and Angov, E. (2012). Impact of Pre-existing MSP142-Allele Specific Immunity on Potency of an Erythrocryptic Plasmodium falciparum Vaccine. Malar. J. 11, 315. doi:10.1186/1475-2875-11-315

Boussena, T., Roeffen, W., Meijerinck, H., Mwerinde, H., Mwakalinga, S., van Gemert, G.-J., et al. (2010). The Dynamics of Naturally Acquired Immune Responses to Plasmodium falciparum Sexual Stage Antigens Pf6230 & Pf64/45 in a Low Endemic Area in Tanzania. PLoS ONE 5, e14114. doi:10.1371/journal. pone.0014114

Chowdhury, D. R., Angov, E., Kariuki, T., and Kumar, N. (2009). A Potent Malaria Transmission Blocking Vaccine Based on Codon Harmonized Full Length Pf64/45 Expressed in Escherichia coli. PLOs ONE 4, e6352. doi:10.1371/journal.pone.0006352

Clement, M., Posada, D., and Crandall, K. A. (2000). TCS: a Computer Program to Estimate Gene Genealogies. Mol. Ecol. 9, 1657–1659. doi:10.1046/j.1365-294X.2000.00100.X

Coelho, C. H., Tang, W. K., Burkhardt, M., Galson, J. D., Muratova, O., Salinas, N. D., et al. (20212021). A Human Monoclonal Antibody Blocks Malaria Transmission and Defines a Highly Conserved Neutralizing Epitope on Gametes. Nat. Commun. 12 (1 12), 1–12. doi:10.1038/s41467-021-21955-1

Conway, D., Machado, R. L. D., Singh, B., Dessert, P., Mikes, Z. S., Povoa, M. M., et al. (2001). Extreme Geographical Fixation of Variation in the Plasmodium falciparum Gamete Surface Protein Gene Pf64/45 Compared with Microsatellite Loci. 115, 145–156. doi:10.1016/S0166-6851(01)00278-x

Coombs, R. W., and Fiscus, S. (2009). Processing of Dried Blood Spots Standard Operating Procedure/Procedure Title: Processing of Dried Blood Spots Standard Operating Procedure Approved by (Network): Network Name, Title Signature Date ACTG Reviewed by (Laboratory): Name, Title Signature Date Revision History Version Effective Date (Dd/mm/yyyy) Comments.

Cvijovic, I., Good, B. H., and Desai, M. M. (2018). The Effect of Strong Purifying Selection on Genetic Diversity. Genetics 209, 1235–1278. doi:10.1534/GENETICS.118.301058

Drakeley, C. J., Durras Singh, M. T., Povo, M., Conway, D. J., Targett, G. A. T., and Baker, D. A. (1996). Geographical Distribution of a Variant Epitope of Pf64/45, a Plasmodium falciparum Transmission-Blocking Vaccine Candidate. Mol. Biochem. Parasitol. 81, 253–257. doi:10.1016/0168-6851(96)02718-1

Ekis, S., Czesny, B., van Gemert, G.-J., Sauerwein, R. W., Eling, W., and Williamson, K. C. (2006). Malaria Transmission-Blocking Antigen, Pf6230, Mediates Human Red Blood Cell Binding to Exflagellating Male Parasites and Oocyst Production. Mol. Microbiol. 61, 991–998. doi:10.1111/j.1365-2958.2006.05284.x

Escalante, A. A., Lal, A. A., and Ayala, F. J. (1998). Genetic Polymorphism and Natural Selection in the Malaria Parasite Plasmodium falciparum. Genetics 149, 189–202. doi:10.1093/genetics/149.1.189

Excoffier, L., and Lischer, H. E. L. (2010). Arlequin Suite Ver 3.5: A New Series of Programs to Perform Population Genetics Analyses under Linux and Windows. Mol. Ecol. Resour. 10, 564–567. doi:10.1111/j.1755-0998.2010.02847.x

Genton, B., Betuela, I., Felger, I., Al-Yaman, F., Anders, R. F., Saul, A., et al. (2002). A Reombinant Blood-Stage Malaria Vaccine Reduces Plasmodium falciparum Density and Exerts Selective Pressure on Parasite Populations in a Phase 1-2b Trial in Papua New Guinea. J. Infect. Dis. 185, 820–827. doi:10.1086/339342
Blocking Antibody Responses against *Plasmodium falciparum*. Front. Immunol. 10, 1256. doi:10.3389/fimmu.2019.01256

Singh, S. K., Thrane, S., Janitzek, C. M., Nielsen, M. A., Theander, T. G., Theisen, M., et al. (2017). Improving the Malaria Transmission-Blocking Activity of a *Plasmodium falciparum* 48/45 Based Vaccine Antigen by SpyTag/SpyCatcher Mediated Virus-like Display. Vaccine 35, 3726–3732. doi:10.1016/J.VACCINE.2017.05.054

Stothard, P. (2000). The Sequence Manipulation Suite: JavaScript Programs for Analyzing and Formatting Protein and DNA Sequences. BioTechniques 28, 1102–1104. doi:10.2144/00286IR01

Tachibana, M., Miura, K., Takashima, E., Morita, M., Nagaoka, H., Zhou, L., et al. (2019). Identification of Domains within Pfs230 that Elicit Transmission Blocking Antibody Responses. Vaccine 37, 1799–1806. doi:10.1016/j.vaccine.2019.02.021

Takala, S. L., Coulibaly, D., Theria, M. A., Dicko, A., Smith, D. L., Guido, A. B., et al. (2007). Dynamics of Polymorphism in a Malaria Vaccine Antigen at a Vaccine-Testing Site in Mali. PLoS Med. 4, e93. doi:10.1371/journal.pmed.0040093

van Dijk, M. R., Janse, C. J., Thompson, J., Waters, A. P., Braks, J. A. M., Dodemont, H. J., et al. (2001). A Central Role for P48/45 in Malaria Parasite Male Gamete Fertility. Cell. 104, 153–164. doi:10.1016/S0092-8674(01)00199-4

Vermeulen, A. N., van Deursen, J., Brakenhoff, R. H., Lensen, T. H. W., Ponnudurai, T., and Meuwissen, J. H. E. T. (1986). Characterization of *Plasmodium falciparum* Sexual Stage Antigens and Their Biosynthesis in Synchronised Gametocyte Cultures. Mol. Biochem. Parasitol. 20, 155–163. doi:10.1016/0166-6851(86)90027-7

**Conflict of Interest:** The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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