Genetic variability and natural selection at the ligand domain of the Duffy binding protein in brazilian *Plasmodium vivax* populations

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Abstract

**Background:** *Plasmodium vivax* malaria is a major public health challenge in Latin America, Asia and Oceania, with 130-435 million clinical cases per year worldwide. Invasion of host blood cells by *P. vivax* mainly depends on a type I membrane protein called Duffy binding protein (PvDBP). The erythrocyte-binding motif of PvDBP is a 170 amino-acid stretch located in its cysteine-rich region II (PvDBP II), which is the most variable segment of the protein.

**Methods:** To test whether diversifying natural selection has shaped the nucleotide diversity of PvDBP II in Brazilian populations, this region was sequenced in 122 isolates from six different geographic areas. A Bayesian method was applied to test for the action of natural selection under a population genetic model that incorporates recombination. The analysis was integrated with a structural model of PvDBP II, and T- and B-cell epitopes were localized on the 3-D structure.

**Results:** The results suggest that: (i) recombination plays an important role in determining the haplotype structure of PvDBP II, and (ii) PvDBP II appears to contain neutrally evolving codons as well as codons evolving under natural selection. Diversifying selection preferentially acts on sites identified as epitopes, particularly on amino acid residues 417, 419, and 424, which show strong linkage disequilibrium.

**Conclusions:** This study shows that some polymorphisms of PvDBP II are present near the erythrocyte-binding domain and might serve to elude antibodies that inhibit cell invasion. Therefore, these polymorphisms should be taken into account when designing vaccines aimed at eliciting antibodies to inhibit erythrocyte invasion.
A series of competitive binding and directed-mutation strategies have demonstrated the importance of the DBP/DARC interaction [5,7-9]. Moreover, individuals lacking Duffy receptors on their erythrocytes are highly resistant to *P. vivax* invasion [10,11]. Because antibodies against PvDBP inhibit the DBP/DARC interaction *in vitro* and also block the invasion of human erythrocytes [12-14], this protein is a major candidate for vaccines. This observation is consistent with positive natural selection acting on PvDBPII and with allelic variation serving as a mechanism of immune evasion.

To formally test this hypothesis in a population genetics framework, several tests of neutrality have been applied to PvDBPII data (Tajima’s D; Fu and Li’s D and F; McDonald-Kreitman), and the rates of synonymous (\(d_s\)) and non-synonymous (\(d_N\)) substitutions have been compared based on the method of Nei and Gojobori (1986). However, these analyses have produced inconsistent results [29,30], and their assumptions are frequently inappropriate for the data analyzed. For instance, the Nei and Gojobori (1986) method of estimating \(d_N\) and \(d_S\) ignores either transition/transversion bias or unequal base/codon frequencies, which may lead to biased results [31]. On the other hand, phylogeny-dependent methods based on maximum likelihood estimates of \(d_N\) and \(d_S\) [31] assume no recombination and have been developed to be applied to sequences from independent, divergent species, rather than for intraspecific analysis [32]. In this study, these problems are overcome by a recently described probabilistic model called omegaMap [33] to infer whether diversifying natural selection has shaped the nucleotide diversity of PvDBPII. It uses a population genetics approximation to the coalescent with recombination (a natural model choice for population data), and it uses reversible-jump Markov Chain Monte Carlo (MCMC) to perform Bayesian inference on both the \(d_N/d_S\) ratio and the recombination rate, allowing each to vary along a DNA sequence. Being based on the Nielsen and Yang [34] model of codon evolution, omegaMap incorporates information about the transition/transversion ratio and codon frequencies from the data in a probabilistic framework. Specifically, the omegaMap approach was applied to PvDBPII to detect the action of diversifying natural selection on regions of PvDBPII containing B- and T-cell epitopes. This population genetics analysis was integrated with computational modelling of the 3-D structure of PvDBPII, which showed that several polymorphic residues (three of which are clearly under positive selection) map close to the erythrocyte-binding cluster of PvDBPII.

**Methods**

**Parasite isolates and study sites**

One hundred twenty-two field isolates of *Plasmodium vivax* were obtained from six sites across the Brazilian Amazon: Cuiabá (Mato Grosso State [MT], *n* = 25), Acrélândia (Acre State [AC], *n* = 25), Manaus (Amazonas State [AM], *n* = 20), Augusto Correa (Pará State [PA], *n* = 18), Porto Velho (Rondónia State [RO], *n* = 20), and Macapá (Amapá State [AP], *n* = 14) (see additional file 1: Locations of blood collection in Brazil). Epidemiological data indicate that malaria transmission in these areas is generally hypo- to meso-endemic [35-37]. The samples were collected from the different regions as follows: AC, between March and April 2004; AM, May 2003; AP, November 2004; PA, October 2005; RO, from October to December 2004; and MT, from July 2003 to July 2004. At the time of blood sample collection, the annual parasite index (API: the number of positive blood smears per thousand inhabitants) of the contagion localities allowed these areas to be classified as high risk (API >50: Augusto Correa/PA and Acrélândia/AC), medium risk (API 10 to 50: Manaus/AM) or low risk for malaria transmission (API <10: Macapá/AP) [38]. In these regions, the individuals were locally infected. However, in Cuiabá/MT and Porto Velho/RO, individuals were infected elsewhere, and therefore the risk of malaria transmission reported there was calculated based on the average API from contagion localities. Thus, both localities were classified as high-risk areas for malaria transmission.

**PvDBPII amplification and sequencing**

Genomic DNA was immediately extracted from whole blood samples using the PUREGENE DNA isolation kit (Gentra Systems, Minneapolis, MN). The sequence encoding PvDBPII, corresponding to nucleotides 870 to 1545 (aa 290-515, reference sequence: Sal-I) [39], was amplified by PCR using high fidelity platinum Taq DNA polymerase (Invitrogen Corporation, Carlsbad, CA) and the primers forward: 5’ ATGTA GGAGAAAATCAGGAT 3’ and reverse: 5’ ACCTGCGGTCTGAACCTTTT 3’. The PCR products were purified using a GFX-96 PCR kit (GE
isolates. The PvDBPII gene polymorphisms identified sequence coverage was achieved for each of the analyzed multiple infections. All singletons were confirmed by infection or from the predominant variant of patients with multiple infections. It was inferred that these came from patients with a single infection or from the predominant variant of patients with multiple infections. All singletons were confirmed by sequencing from independent PCR amplifications. Six-fold sequence coverage was achieved for each of the analyzed isolates. The PvDBPII gene polymorphisms identified among the Brazilian Amazon isolates were compared to all DBP gene sequences of \textit{P. vivax} in GenBank (last update on 25th February 2010). The PvDBPII sequences identified here were deposited in GenBank with accession numbers EU812839 to EU812960. Sequence alignments and comparisons were performed using the Clustal W multiple alignment algorithm in BioEdit Sequence Alignment editor using manual editing [40].

Diversity analysis of PvDBPII
To estimate within-population diversity, the numbers of segregating sites and observed haplotypes were counted. Estimates of the nucleotide diversity (\(\pi\)), haplotype diversity (HD), and their corresponding standard deviations (SDs) were obtained using DnaSP 4.10 software [41]. Between-population differentiation was measured using the pairwise fixation index \(F_{ST}\) with Arlequin 3.01 software [42]. The \(F_{ST}\) index (between-population component of the genetic variance, Wright 1951) was calculated assuming the Kimura 2-parameter mutation model [43,44]. The correlation between pairwise population genetic distances (\(F_{ST}\)) and geographical distances was estimated by nonparametric Spearman’s rank correlation.

Analysis of recombination and linkage disequilibrium (LD)
The recombination parameter \(\rho\) among contiguous sites across PvDBPII was estimated by the method described by Li and Stephens [45] using PHASE 2.1 software. The parameter \(\rho\), which can be estimated by different statistical methods, depends both on the recombination rate and on the effective population size, and therefore, it is proportional to the number of recombination events per generation that occurred in a population. The algorithm was run twice with 500,000 iterations, with a 20,000 iteration burn-in and a thinning interval of 1,000 iterations. The pairwise linkage disequilibrium (LD) across PvDBPII was analyzed using the \(r^2\) statistic [46], and its significance was assessed by calculating LOD (base-10 logarithm of odds) scores. LD was calculated only for SNPs with a minor allele frequency (MAF) higher than 0.1 in the total Brazilian sample. These calculations were performed and visualized using the program Haploview [47].

Tests of neutrality
The method developed by Wilson and McVean [33], implemented in the omegaMap software, was used to estimate the parameter \(\omega = d_N/d_S\), where \(d_N\) is the rate of non-synonymous substitutions and \(d_S\) is the rate of synonymous substitutions. Estimates of the parameter \(\omega\) are indicators of the action of natural selection on coding sequences. Under no selection \(\omega\) is expected to be equal to 1, under negative selection the value is expected to be significantly less than 1, and under positive selection the value is expected to be significantly greater than 1. This method employs a Bayesian approach to parameter estimation that is independent of phylogeny, and it is therefore less prone to falsely detecting diversifying selection. The model uses a population genetic approximation to the coalescent with recombination, and it uses reversible-jump Markov Chain Monte Carlo (MCMC) to perform Bayesian inferences of both \(\omega\) and the recombination parameter \(\rho\), allowing both parameters to vary along the sequence. In this study, analyses were conducted using 10 randomly chosen orderings of the haplotypes and the following priors: \(\mu\) (synonymous transversion) = Improper inverse, \(\kappa\) (transition/transversion rate) = Improper inverse, \(\varphi\) (insertion/deletion rate) = Improper inverse. Since \(\omega\) requires a proper prior, a log-normal distribution centered on log (\(\omega\)) = 0 was used with a wide standard deviation of 2. An average block length of 20 codons was chosen to be around 10% of the sequence length. Estimates of the recombination parameter \(\rho\) were obtained by omegaMap among contiguous sites across PvDBPII. For \(\rho\), there is no natural “reference” prior, so it was used an informative prior based on Gunasekera \textit{et al.} [48]: a log-normal distribution with parameters log (0.1) and 1. As it was expected that there would be less information regarding \(\rho\) than \(\omega\), an average block length of 40 codons was assumed for \(\rho\) to impose greater smoothing. Sixteen independent MCMC chains were run for 90,000 iterations with a 10,000 iteration burn-in. The chains were compared for convergence and merged to obtain the posterior distributions. The results showed the geometric mean for the posterior distribution of \(\omega\) and \(\rho\) and the 95% HPD (highest posterior density) interval. To determine the influence of the choice of priors on the posteriors, the analyses were repeated with an alternative set of priors (\(\mu = \text{Exponential mean 0.02}; \kappa = \text{Exponential mean 2.0}; \varphi = \text{Exponential mean 0.1}; \omega = \text{Exponential mean 1.0}; \rho = \text{Exponential Mean 0.14}\)). The means for priors were chosen using the posterior distributions generated with the first prior set. A mean value of 1 was used for the \(\omega\) prior based on the null hypothesis of selective neutrality.

To integrate the evolutionary analysis with structural information, the posterior probability of selection at
each codon was mapped onto the 3-dimensional structure of PvDBP II (see below) to visualize the spatial distribution of selected sites. The correlation between $F_{ST}$ for each polymorphism and its $\omega$ estimators was calculated using Pearson’s correlation (Minitab statistical software, release 13.20).

**Prediction of promiscuous T-cell epitopes**

Because only a few epitopes have been described for PvDBP [30,49], an *in silico* analysis was used to infer new and promiscuous PvDBP II T-cell epitopes able to bind to several alleles of host MHC molecules. To identify the MHC Class-I binding regions in PvDBP II, the ProPred-I program was used [50]. This program uses matrices for each of the 47 MHC Class-I alleles the elements of which denote the affinity of antigen amino acids for specific MHC alleles, thereby allowing peptides in PvDBP II with antigenic properties to be identified. Quantitative matrices derived from the published literature [51] and implemented in ProPred were also used to predict MHC Class-II binding regions in PvDBP II [52].

**3-D structural model of PvDBP II**

Homology modelling was performed with Modeller 8v2 software using the X-ray structure of the *Plasmodium knowlesi* DBL domain as a template (Pka-DBL) (PDB file: 2C6j, chain A) [53]. This template shares 71% sequence identity with PvDBP II. The 3-D structure of PvDBP II was modelled between amino acids 261 and 553. The quality of these models was evaluated with PROCHECK v.3.5.4 software [54]. Superpositioning and renderings were carried out with PyMol v1.0 software [55].

**Results**

**Polymorphisms in PvDBP II among Brazilian Amazon isolates**

By sequencing 675 bp of the PvDBP region II of 122 isolates from six Brazilian Amazon regions, 19 polymorphic sites (three synonymous substitutions and 16 non-synonymous substitutions) were identified with an overall nucleotide diversity of 0.0081 (Table 1 and Figure 1A). Of 16 sites with non-synonymous polymorphisms, one site at position 386 was not dimorphic. At this site, lysine could be substituted by glutamine or asparagine (K386Q/N). A positive correlation was found between the risk of malaria transmission in each area (estimated based on the annual parasitic index [API]) and both the nucleotide diversity and the haplotype diversity of PvDBP II (Spearman correlation coefficients: $r_s = 0.811$, $p = 0.049$; $r_s = 0.828$, $p = 0.041$, respectively). Most of the non-synonymous polymorphisms were non-conservative (75%) with changes of the physico-chemical family of the respective amino acid. In a comparison of the polymorphisms identified in isolates studied here to the 401 PvDBP II sequences available in GenBank, only one was recognized as exclusive to Brazilian isolates: N305N (a synonymous substitution of Asparagine at position 305), and this was present in 10 isolates from Amapá (see additional file 2: Table with description of PvDBP II polymorphisms identified in Brazilian isolates).

The polymorphisms observed in Brazil were arranged in 34 haplotypes (Figure 1B), corresponding to a mean haplotype diversity of 0.934 (Table 1). The modal haplotype Hap19 was observed in all six sampled regions. The $F_{ST}$ values of the populations ranged from 2.33 to 30.29%, with the highest values corresponding to the Amapá population (Table 1). Moreover, the $F_{ST}$ values did not correlate with the geographic distance between populations (Spearman correlation coefficient $r_s = 0.0750$, $p = 0.7905$).

**Intragenic recombination and linkage disequilibrium in PvDBP II**

In the absence of recombination and recurrent mutation, it would be expected that the 19 polymorphic sites observed would be arranged into a maximum of 20 haplotypes. Because 34 haplotypes were detected, it was inferred that recombination (and possibly recurrent mutation) has shaped the pattern of PvDBP II variation in Brazilian populations. Figure 2A shows the variation in the recombination parameter ($\rho$) (calculated using the software omegaMap) along the PvDBP II sequence, indicating that at least one region was clearly affected by recombination (aa 381-390, lower 95% HPD > 0.1). These results were further confirmed by estimating the recombination parameter with PHASE 2.1 software (Figure 2B) and also by identifying regions with low LD across PvDBP II using the statistic $r^2$ measured with Haploview (Figure 3A). Consistent with the observed haplotype structure and $\rho$ estimates, linkage disequilibrium (LD) across the PvDBP II sequence is low in the Brazilian isolates (Figure 3A). In fact, an analysis of the 12 SNPs in the whole sample with a minor allele frequency higher than 0.1 showed that only 8 of 66 pairwise combinations showed $r^2 > 0.50$. The highest $r^2$ values were observed for several spatially close polymorphisms, encompassing the SNPs delimited by N417K and I464I (LOD scores always > 6.64). In particular, three residues within this set were spatially close in the 3-D model (417, 419 and 424). Because the LD analysis could be affected by population structure, the data analysis was repeated without the Amapá population (the most differentiated population according to $F_{ST}$ value), and an almost identical LD pattern was observed (Figure 3B).
Evidence of natural selection on PvDBP\textsubscript{II} and its epitopes

To gain insight into the action of natural selection, the omegaMap approach was used [33], which is suitable for genomic regions (such as the PvDBP\textsubscript{II} gene) where recombination prevents the reliable reconstruction of a phylogeny. This analysis, which estimates \( \omega \) and its associated HPD interval, also revealed that natural selection does not act uniformly within the gene (Figure 4A). The sequence contains regions where purifying selection seems to predominate, regions with \( \omega \) values close to 1 that appear to be neutral or less affected by selection and regions under positive natural selection (\( \omega > 1 \)), which are mainly in a central region of the protein (aa 368-439). To determine the influence of the choice of priors on the posteriors, the analyses were repeated with an alternative set of priors used (see additional file 3: Recombination and selection parameters in PvDBP\textsubscript{II} with an alternative set of priors).

Pathogen epitopes are responsible for host immune recognition. Thus, the association between positive natural selection and the presence of T- and B-cell epitopes in PvDBP\textsubscript{II} was tested (Figure 4A). The results indicated that \( \omega \) was higher for sites belonging to B-cell and T-cell epitopes (experimentally identified and predicted) than for non-epitope sites (Mann Whitney U = 2587.00, \( p < 0.0001 \)) [30,49]. The present study therefore showed that positive natural selection preferentially acts on epitopes in PvDBP\textsubscript{II}. In addition, the epitopes in the region showing the highest \( \omega \) values also have greater nucleotide diversity (0.011 - 0.046) than the value for all of PvDBP\textsubscript{II} (0.0081) (Figure 4B). In particular, nucleotide diversity was the highest in \textit{in silico} predicted epitope Ia. Ia contains three spatially proximate polymorphic residues at positions 417, 419 and 424 and shows a clear signature of positive natural selection, with \( \omega \) values of 3.72, 3.72 and 3.31, respectively.

Spatial distribution of polymorphisms across PvDBP\textsubscript{II}

The crystallographic structure of the Pk\textalpha-DBL binding-domain was used to produce a computational model of PvDBP\textsubscript{II}. In Figure 5A, this model of PvDBP\textsubscript{II} is presented in space-filling form, and it shows the previously identified essential, invariant and surface-exposed contact residues required for the recognition of DARC on human erythrocytes [53], as well as the residues that are polymorphic in the Brazilian Amazon samples. Although polymorphic residues were widely distributed throughout the sequence, the polymorphic residues at positions 417, 419, and 424, which are part of a block of high LD, are close to the erythrocyte-binding cluster of PvDBP\textsubscript{II} (Figure 5A). Both K417 and I424 are prevalent in Brazil as well as in other studied areas (Additional file 2).

The three-dimensional model of PvDBP\textsubscript{II} also shows the residues according to the posterior probability of their estimated \( \omega \) (Figure 5B). Residues that seem to be less affected by positive selection (shown in white) were distributed throughout the protein, and interestingly, residues associated with high \( \omega \) (likely to have evolved under positive selection, shown in green) were close to (and on the same face of) the proposed DARC-recognition site. Moreover, these residues tend to be in epitope regions, which were surface exposed (Figure 5C).

Discussion

The data obtained in this study regarding the nucleotide diversity of PvDBP\textsubscript{II} among 122 \textit{P. vivax} isolates from Brazil provide evidence that (i) recombination plays an important role in determining the haplotype structure of PvDBP\textsubscript{II} and (ii) natural selection acts differentially across the PvDBP\textsubscript{II} sequence. Indeed, the gene appears to contain neutrally evolving codons as well as codons evolving under natural selection, and these latter codons are associated with high \( \omega \) values (diversifying selection). The most striking conclusion of the current study is that diversifying natural selection acts on one PvDBP\textsubscript{II} region containing B- and T-cell epitopes previously identified and predicted \textit{in silico} [30,49]. In particular, the amino acid residues at positions 417, 419, and 424 are associated with high \( \omega \) values and map close to the erythrocyte-binding cluster of the molecule.

| Table 1 Nucleotide and haplotype diversity of PvDBP\textsubscript{II} gene and FST among Brazilian populations |

| Population\textsuperscript{a} | \( S \textsuperscript{b} \) | \( n \textsuperscript{c} (SD) \) | \( H \textsuperscript{d} \) | \( Hd \textsuperscript{e} (SD) \) | \( FST (\times100)\textsuperscript{f} \) |
|-----------------|------|-----------------|------|-----------------|-----------------|
| AP (N = 14)     | 11   | 0.0050 (0.0013) | 6    | 0.747 (0.111)  | -               |
| AM (N = 20)     | 11   | 0.0057 (0.0009) | 7    | 0.779 (0.082)  | 21.02*          |
| PA (N = 18)     | 11   | 0.0070 (0.0004) | 8    | 0.869 (0.049)  | 18.68*          |
| RO (N = 20)     | 17   | 0.0087 (0.0008) | 16   | 0.979 (0.021)  | 30.29*          |
| MT (N = 25)     | 16   | 0.0080 (0.0009) | 14   | 0.897 (0.043)  | 14.51*          |
| AC (N = 25)     | 13   | 0.0079 (0.0005) | 11   | 0.917 (0.028)  | 26.59*          |
| Total (N = 122) | 19   | 0.0081 (0.0003) | 34   | 0.934 (0.012)  |                |

Populations from Amapa (AP), Amazonas (AM), Para (PA), Rondonia (RO), Mato Grosso (MT) and Acre (AC). N: number of isolates; S: Number of segregating (polymorphic/variable) sites; \( K \): Average number of pairwise nucleotide differences; \( H \): Number of haplotypes; \( Hd \): Haplotype diversity; \( \pi \): Observed average pairwise nucleotide diversity, \( FST \): Fixation index, a measure of genetic differentiation between populations. * \( p < 0.05 \).
Diversifying selection has been observed in multiple proteins involved in parasite-host interactions, and it is generally attributed to the selective pressure of the host’s immune response [56]. This study inferred that the evolution of the ligand domain of PvDBP, particularly of the codons for epitopes 45, 48, 66, Ia, IIa and IIb, is driven by diversifying selection (as indicated by high $\omega$ values). These epitopes are surface exposed and have significantly higher nucleotide diversity than the other residues in this region. However, inferential methods can lack the statistical power to reject neutrality and detect the action of diversifying selection, particularly when the analysis is restricted to a few codons such as epitope sequences. In the present study, this may be the case for epitopes 5, 16, 18, 20, Ib and Ic. In this context, the use of appropriate methods for inferring natural selection is important, and new methods based on more realistic assumptions about molecular evolution (such as that of Wilson-McVean) will help to assess the role of diversifying selection in parasite diversity.

Comparisons with previous analyses

The low linkage disequilibrium (LD) and high recombination parameter across the PvDBPII gene suggest that meiotic recombination partially accounts for the observed haplotype diversity. This result corroborates the findings of Martinez et al. [29], who also found high recombination rates in PvDBPII among isolates from Papua New Guinea, Colombia and Korea. Using omegaMap with PNG sequences, similar results were obtained (see additional file 4: Spatial variation in recombination ($r$) and omega parameter ($\omega$) calculated using Figure 2).
omegaMap software among Papuan New Guinea PvDBP\textsubscript{II} sequences). However, Cole-Tobian and King [30] found a low recombination rate in PvDBP\textsubscript{II}, but the test used (Sawyer’s permutation test) is not appropriate for estimating the recombination rate in relatively short regions like this [57]. The results presented here are supported by three different analyses: Haploview; PHASE and omegaMap. These results reinforce the importance of recombination as well as natural selection in the generation and maintenance of the genetic diversity of PvDBP\textsubscript{II}. Multiple recombination events have also been described in other \textit{P. vivax} antigens, such as the Apical Membrane Antigen 1 (PvAMA1), Merozoite Surface Protein-3\textalpha{} (MSP) and MSP-1 [48,58,59].

Previous attempts to infer the action of natural selection in PvDBP\textsubscript{II} with different neutrality tests yielded inconclusive results. Martinez \textit{et al.} [29] used Tajima’s [60] and Fu-Li’s [61] neutrality tests and found an excess of rare alleles for PvDBP\textsubscript{II}, which is consistent with either the action of a selective sweep or the purifying natural selection (as suggested by the authors). A limitation of these tests is that the allelic spectrum is influenced both by natural selection and by the demographic history of the population, and it is not trivial to discriminate between the influence of these two variables without significant knowledge of the demographic history of \textit{P. vivax} populations, which is not the case. A second type of neutrality test applied to the PvDBP\textsubscript{II} data compares the pattern of synonymous (assumed to be neutral) and non-synonymous polymorphisms (for which natural selection is possible). Because the demographic history of a population affects synonymous and non-synonymous sites equally, these tests are less sensitive to the potential confounding effects of the demographic history of populations. Applications of these analyses to PvDBP\textsubscript{II} data include the McDonald-Kreitman test [62] and comparisons between \(d_S\) and \(d_K\) estimated using the Nei-Gojobori method [63], and these suggest the action of positive natural selection acting on PvDBP\textsubscript{II} [29,30]. However, the new method of Wilson-McVean [33] has enabled us to map how the action of natural selection varies across PvDBP\textsubscript{II} and to integrate this evolutionary information with an epitope analysis and structural modelling. Indeed, the Wilson-McVean method has four features that improve its performance in correctly identifying the effect of diversifying selection. First, it assumes a more realistic mutational model [34] that allows for different rates of transitions and transversions and for differences in codon frequencies. Second, its fully Bayesian approach allows \(\omega\) parameters and their associated HPD intervals to be estimated, providing a measurement of uncertainty. Third (and more importantly), the Wilson-McVean method uses a reversible-jump MCMC to explore the posterior distribution of \(\omega\), allowing it to vary along the length of the sequence. Although incorporating these more realistic assumptions implies a computational cost, the analyses have demonstrated their value. The Wilson-McVean method allowed us to map how diversifying selection has acted across PvDBP\textsubscript{II} and to test the association between the presence of epitopes and the action of natural selection. Last (but not least), recombination, which has been important in shaping PvDBP\textsubscript{II} diversity, does not affect the performance of Wilson-McVean estimations. These features, the fact that a highly polymorphic and functionally important region of PvDBP was examined for which results of functional studies are available, and the complementary structural and epitopes analyses partially overcome one of the limitations of this study: the small size of the studied region. A thorough examination of the entire \textit{PvDBP} sequence in different populations will provide a better understanding of how natural selection has shaped the different regions of the gene across different populations, including Brazil.

\begin{figure}[h]
\centering
\includegraphics[width=\textwidth]{figure3.png}
\caption{Linkage disequilibrium in PvDBP\textsubscript{II} Patterns of pairwise linkage disequilibrium (\(r^2\)) in the PvDBP\textsubscript{II}-encoding gene from Brazilian isolates. (A) Linkage disequilibrium was calculated for SNPs for which the frequency of the minor allele was at least 0.1 in the entire sample. White: \(r^2 = 0\); black: \(r^2 = 1\); shades of gray: \(0 < r^2 < 1\). The bracket indicates the cluster of amino acids with the highest \(r^2\). (B) LD was calculated as previously described but not including the Amapá population.}
\end{figure}
Figure 4 Spatial variation in selection parameters across PvDBPα. Omega parameter estimates (ω) calculated using omegaMap. ω > 1 is evidence for diversifying natural selection, whereas ω < 1 is evidence for purifying natural selection. The sitewise mean (solid line) and 95% HPD intervals (dotted lines) are shown (A). The short lines above the solid line represent previously identified epitopes: 5 [299-VNNTDNHFR/S/DIFTR-313], 16 [321-LIYDAVEGD(L/F)/KLNNYR-335], 18 [325-AVEGD(L/F)/KLNNYR-339], 20 [329-GDLL(L/F)/KLNNYR-343], 45 [379-SIFGT(D/G)(E/K)(K/N) AQQ(R/H)RKQ-393], 48 [385-(E/K)(K/N)AQQ(R/H)RKQWWNESK-399] and 66 [421-ICK(L/I)NV(A/P)VNIEPQIY-435] [30, 49] and in silico predicted promiscuous epitopes: Ia [416-G(N/K)F(I/M)WICK(L/I)-424], Ib [482-KSYD(Q/E)WITR-490], Ic [497-VLSNKF(I/K)SVKNAEK-510], IIa [408-YSVKKRKG(N/K)-417] and IIb [417-8(F/R)/WCK/L/N-426]. (B) Nucleotide diversity (π) and standard deviation (SD) were calculated for each epitope sequence. Epitope nucleotide diversity values significantly higher than the global nucleotide diversity of PvDBPα (π = 0.0081) are shown in bold.
However, on the basis of the results presented here, it is anticipated that, if the observed level of recombination affects the entire gene, signatures of natural selection might not propagate across large genomic regions. Therefore, population genetics methods that are able to examine signatures of selection at localized regions may be more appropriate for an evolutionary analysis of PvDBP.

**Effects of population structure**

In the present study a high level of population structure was observed, as indicated by high $F_{ST}$ values (from 2.33 to 30.29%), and this could affect the estimates of both linkage disequilibrium and $\omega$. However, when the most differentiated population (Amapá) was excluded from the analyses, the same LD pattern was still observed, which indicates that these results are not an artifact of population structure. In particular, the residues at positions 417, 419, and 424, which are associated with high $\omega$ values and map near the DARC binding domain in the structural model, are in LD even when the Amapá population is excluded from the analysis.

The possibility that the high $\omega$ values inferred could be an artifact of population structure was also investigated. Two features of the data suggest that this is not the case. First, if there is more differentiation ($F_{ST}$) of non-synonymous polymorphisms than synonymous polymorphisms, high $\omega$ values may emerge when $\omega$ is estimated with sequences from different clustered populations, but this is not the case in the data presented here (non-synonymous $F_{ST}$: 14.7%, synonymous $F_{ST}$: 24.0%). Second, if population structure creates high $\omega$ values, a correlation between the $F_{ST}$ for non-synonymous sites and the $\omega$ values of the codons containing these sites is expected, but this was not observed (Pearson correlation 0.2647, $p = 0.303$). Therefore, there is no evidence that the observed pattern of LD and inferences about positive natural selection are artifacts of the population structure of our Brazilian samples.

**Immunological implications**

*Plasmodium vivax* immune evasion could be explained by a model based on a "just-in-time" release mechanism of PvDBP [53,64]. This model proposes that DBP is released from micronemes immediately before binding to DARCs, thereby avoiding immune inhibition by direct binding of antibodies to PvDBP. Therefore, residues on the surface of the protein that is opposite to that of the erythrocyte-binding domain remain exposed and are immune system targets. Supporting this model is a previous observation that polymorphic residues occur on the opposite side from the erythrocyte-binding domain [53]. Therefore, from the perspective of population genetics, it may be expected that positive natural selection imposed by the host’s immune response would preferentially act on the protein surface opposite from the erythrocyte-binding domain, and values of $\omega > 1$ would be concentrated on this side of the protein. Because the results presented here do not match this expectation (Figure 5), the analysis of the Brazilian isolates does not support the "just-in-time" release model. Notably, the results presented here show that stretches with high genetic diversity and residues under positive selection are present near the erythrocyte-binding domain. These results would be expected under a model of immune escape, in which polymorphic residues near the binding site elude recognition by inhibitory antibodies, implying that positive selection acts preferentially on residues near the erythrocyte binding site. This model is different from the "just-in-time" model but is not mutually exclusive.

In fact, the polymorphic residues at positions 417, 419 and 424, which form a unit of LD and are associated
with high ω values, map to near the erythrocyte-binding domain. The residue at position 417 was also reported to be part of a linked haplotype in PNG (N417K, W437R, I503K) [9]. This model of immune evasion has been proposed for PvDBP [24,65-67] and also for PfAMA1 [68]. Recent functional results from Grimberg et al. [13] support this mechanism and show that antibodies involved in invasion-inhibitory activity also block PvDBP-DARC interaction. Recently, it was demonstrated that some epitopes described here (epitopes la and lb) were also recognized by inhibitory antibodies in immune sera of individuals from Papua New Guinea as linear B-cell epitopes [69]. These epitopes include residues 417, 419 and 424, suggesting their involvement as targets of inhibitory antibodies.

Recent data from experimental and natural infections support our findings that PvDBP polymorphisms might have immunological relevance. Using rabbit immune sera, VanBuskirk et al. [9] demonstrated that polymorphisms change the antigenic character of DBP and can compromise immune inhibition. Moreover, evidence was found that in the Amazon area, long-term exposure to malaria is required to allow the production of anti-DBP antibodies able to inhibit the binding of different PvDBP∈ variants to DARC [12]. These results support the hypothesis that immune responses against PvDBP are haplotype-dependent, as has also been proposed for the apical membrane antigen 1 of P. falciparum (PFAMA1), another micronemal protein vaccine candidate [70]. However, neutrality tests simply test the hypothesis that the observed pattern of diversity cannot be explained merely by the neutral model of evolution, and may suggest which type of natural selection has acted. Inferences about natural selection need to be complemented with experimental tests of the effects of the observed variations on parasite invasion and immune escape.

Conclusions
The low LD and the high recombination rates observed in PvDBP∈ haplotypes suggest that recombination and diversifying selection are major factors influencing PvDBP∈ haplotype diversity in the Brazilian Amazon region. The results presented here stress the importance of using appropriate tests (such as the omegaMap approach [33]) to infer the relative roles of natural selection and recombination when determining the pattern of diversity of parasite genes. These genes are often involved in interactions with host immune systems, and they are strong candidates for involvement in co-evolutionary processes. However, all methods of inferring the action of positive selection have limitations [31,33,71]. Therefore, as pointed out by Nozawa et al. [71], it is important to complement (as in this study) pure population genetics inferences with information from structural modelling, in-silico and experimental inferences about epitopes, and functional studies. Taken together, these results suggest that vaccine strategies based on inhibitory anti-DBP antibodies that take into account PvDBP∈ haplotype variability can overcome the problem of antigen variability and generate a strain-transcending and long-lived vaccine. Moreover, the recent finding of Duffy-negative individuals infected with P. vivax emphasizes the importance of a multivalent vaccine strategy that, in addition to eliciting inhibitory antibodies, may also reduce the ability of parasite strains to escape immunological control.

Additional material

Additional file 1: Locations of blood collection in Brazil. The cities are indicated with their respective states: Amazonas (AM), Pará (PA), Amapá (AP), Acre (AC), Rondônia (RO) and Mato Grosso (MT). The areas where malaria is endemic are shown in gray.

Additional file 2: PvDBP∈ polymorphisms of Brazilian isolates. This file provides descriptions of the PvDBP∈ polymorphisms identified in the Brazilian isolates by geographic region as well as polymorphisms that were previously described in other localities.

Additional file 3: Recombination and selection parameters in PvDBP∈ with an alternative set of priors. The spatial variation in recombination and selection parameters across PvDBP∈ was obtained using omegaMap software. The analysis was performed using an alternative set of priors: μ = Exponential mean 0.02; α = Exponential mean 2.0, φ = Exponential mean 0.1, ω = Exponential mean 1.0, p = Exponential Mean 0.14. The means for the priors for μ, α and p were chosen using the posterior distributions generated with the first prior set. (A) Recombination parameter estimates (φ). The sitewise mean (solid line) and 95% HPD (highest posterior density) interval (dotted lines) are shown. (B) Omega parameter estimates (ω). The sitewise mean (solid line) and 95% HPD intervals (dotted lines) are shown.

Additional file 4: Spatial variation in recombination (φ) and omega (ω) among Papuan New Guinea PvDBP∈ sequences calculated using omegaMap (A) Evidence of recombination is φ > 0.1. (B) Evidence of diversifying natural selection is ω > 1 and for purifying natural selection it is ω < 1. The sitewise mean (solid line) and 95% HPD intervals (dotted lines) are shown.

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Authors’ contributions
TNS performed all the experiments and analyzed the data. APM supervised the statistical analysis. DJW managed the omegamAP analysis. PRK supervised the modelling experiments. CIPT, JMS, AARDAC, MUF and UHSG organized and supervised the field work, LHC and CFAB designed the study. CFAB and ETS supervised the data analysis. TNS, CFAB and ETS wrote the paper with critical suggestions from LHC and MUF. All authors read and approved the final manuscript.

Competing interests
The authors declare that they have no competing interests.

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