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Citation

Published Version
doi:10.1093/molbev/mss151

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SNP Genotyping Identifies New Signatures of Selection in a Deep Sample of West African *Plasmodium falciparum* Malaria Parasites

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Abstract

We used a high-density single-nucleotide polymorphism array to genotype 75 *Plasmodium falciparum* isolates recently collected from Senegal and The Gambia to search for signals of selection in this malaria endemic region. We found little geographic or temporal stratification of the genetic diversity among the sampled parasites. Through application of the iHS and REHH haplotype-based tests for positive selection, we found evidence of recent selective sweeps at a known drug resistance locus, at several known antigenic loci, and at several genomic regions not previously identified as sites of recent selection. We discuss the value of deep population-specific genomic analyses for identifying selection signals within sampled endemic populations of parasites, which may correspond to local selection pressures such as distinctive therapeutic regimes or mosquito vectors.

Key words: *Plasmodium falciparum*, natural selection, SNP, genome, West Africa.

*Plasmodium falciparum* is an obligate protozoan parasite that causes malaria worldwide, killing ~800,000 annually (WHO 2010). It faces strong selection pressures from host immune responses and from drugs and (potentially) vaccines. Genome-wide scans for natural selection can identify genetic loci responding to these pressures. Most previous scans for selection in *P. falciparum* have used global collections of parasites and were thus well positioned to identify loci undergoing selection worldwide (Mu et al. 2010; Van Tyne et al. 2011). However, much natural selection occurs at smaller geographic scales, both because selective forces such as drug regimes and insect vectors vary geographically and because selected phenotypes can appear independently in different regions. Detection of localized selection is difficult in global surveys but instead requires a deep survey from a single geographic region; the one study similar to this to date (Mu et al. 2007) examined primarily Asian parasites.

To search for selection in a single African parasite population, we collected 75 recent *P. falciparum* isolates (supplementary table S1, Supplementary Material online) from four locations in Senegal and The Gambia, a region spanning several hundred kilometers. All parasites were hybridized to a high-density Affymetrix single-nucleotide polymorphism (SNP) array with ~17,000 assays (Van Tyne et al. 2011); after filtering out mixed infections and closely related parasites (which are uninformative for haplotype-based tests of selection), we analyzed genotypes from 70 samples for selective sweeps.

We first checked for population structure within our sampled region. Principal components analysis (fig. 1) showed little structure, whereas $F_{ST}$ (which measures population differentiation) indicated a small, statistically significant difference between Senegal and The Gambia ($F_{ST} = 0.0072$, $P < 0.0001$). This proved to stem from subtle differences between culture-adapted parasites (present only in the Senegal set) and parasites isolated directly from patient blood (see supplementary note and supplementary fig. S1, Supplementary Material online). We found no significant structure in time across the ~10 years of sample collection. The lack of structure suggests enough gene flow within this region (on a scale of hundreds of kilometers) that sampling for genome-wide association studies need not be finer grained spatially, at least for this sample size. We also measured linkage disequilibrium throughout the genome and found it generally consistent with previous reports (see supplementary note and supplementary figs. S2 and S3 for details and caveats, Supplementary Material online).

We identified possible selective sweeps using two haplotype-based tests for positive selection: relative extended haplotype homozygosity (REHH) (Sabeti et al. 2002) and integrated haplotype score (iHS) (Voight et al. 2006). These tests identify alleles that lie on unusually long haplotypes for that

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region, indicative of a recent selective sweep. We detected 11 loci with genome-wide significance, including the well-characterized \textit{pfcr} locus (Fidock et al. 2000) and a large region on chromosome 6 (fig. 2, table 1; supplementary table S2 has detailed annotation, Supplementary Material online). Five of the loci overlap with previously reported signals (Mu et al. 2010; Van Tyne et al. 2011). The signals of selection appeared consistently in both our directly drawn and our culture-adapted parasite sets (supplementary fig. S4, Supplementary Material online) and were little changed when we restricted our analysis to a 2-year time period (supplementary fig. S5, Supplementary Material online) or used a uniform recombination map. After removing the 11 loci, the remainder of the genome roughly conforms to the null expectation for test scores (fig. 2c and d).

Senegal and The Gambia share a similar history of drug regimens, except for three years of amodiaquine use in Senegal but not in The Gambia (see supplementary note, Supplementary Material online). Given the role that loci such as \textit{pfcr}, with its large selective sweep, play in drug resistance (Fidock et al. 2000), it is reasonable to speculate that some of these novel sweeps also reflect adaptation to drug pressure. Although resistance alleles at loci such as \textit{pfcr}, \textit{pfmdr1}, and \textit{dhfr} contribute significantly to drug resistance in \textit{P. falciparum}, it is known that other genes also affect resistance to drugs (Patel et al. 2010) or have undergone compensatory changes to offset the fitness cost of resistance alleles (Jiang et al. 2008).

Contrary to intuition, several of the loci with evidence of directional selection are known antigenic loci, which are thought to be subject primarily to balancing selection; these include \textit{ama1}, \textit{trap}, \textit{ssp2}, \textit{clag2}, and possibly \textit{PF13_0074} and \textit{PF14_0726}. Similar findings have been noted previously (Mu et al. 2010). Genes under balancing selection typically exhibit short haplotypes, as their SNPs segregate for unusually long periods of time during which recombination breaks down haplotypes. If the selective sweeps do originate in these genes (and not at nearby variants absent from the array), they may reflect nonimmune pressures. Many highly polymorphic antigenic loci in \textit{P. falciparum} have roles unrelated to immune evasion and presumably fix adaptive mutations related to these functions. For example, \textit{clag3} genes, in the same gene family as one of our sweep candidates (\textit{clag2}), exhibit the high polymorphism typical of genes encoding surface-expressed cytoadherence proteins. Nevertheless, two \textit{clag3} genes were recently found to be associated with resistance to antimalarial drugs, suggesting that their products also mediate entry of drugs (Ndungu et al. 2011).

Alternatively, we may be detecting positive selection at these loci as a variant unfamiliar to the host immune system rises from very low frequency. Previous studies of malaria antigenic genes (\textit{ama1} in particular) have shown complex patterns of selection (Cortes et al. 2003; Polley et al. 2003; Gunasekera et al. 2007), with evidence for both positive and balancing selection (Mu et al. 2010); a long-haplotype signal at \textit{ama1} has been detected in Asia (Mu et al. 2010). Sequencing-based study of these regions, preferably in multiple geographic regions, should provide better insight into the selective processes at work.

Another unexpected candidate locus codes for thrombospondin-related anonymous protein (TRAP), a conserved protein expressed in \textit{P. falciparum} sporozoites. It contains extracellular domains associated with hepatocyte invasion (Muller et al. 1993; Wang et al. 2005) and has previously shown evidence of balancing selection (Weedall et al. 2007). TRAP has also been found to mediate the invasion of salivary glands in mosquitoes (Ghosh et al. 2009). It is possible that maintenance of long haplotypes reflects adaptation to divergent ligands in different mosquito species.

\textbf{Materials and Methods}

\textit{Plasmodium falciparum} isolates were obtained in 2008 from three sites near Banjul in The Gambia and during the period 2001–2009 from three clinics in Senegal—in Thies (100 km from Dakar), in Pikine (10 km from Dakar), and in Vellingara (500 km from Dakar and on the other side of The Gambia). Gambian parasites were taken from clinical samples, whereas Senegal samples included both directly drawn and culture-adapted parasites; see supplementary note for details on DNA extraction, Supplementary Material online. Culture-adapted samples with evidence of multiple infection, based on polymorphism typing at the \textit{msp} loci (Viriakosol et al. 1995), were subcloned to isolate a single strain.

SNP calling was as described in Van Tyne et al. (2011): parasite DNA was hybridized to a \textit{P. falciparum} Affymetrix array containing 74,656 SNP markers and genotypes called using the BRLMM-P algorithm. SNPs were validated by comparing array genotypes with Sanger sequencing genotypes for 17 reference strains (Van Tyne et al. 2011); perfect concordance was required, as was a minimum 80% call rate. Genomic positions and translations are based on the PlasmoDB v5.0 assembly and annotation (PlasmoDB.org). SNPs within \textit{var}, \textit{rifin}, or \textit{stevor} genes were dropped to reduce artifacts from duplicate sequence. After these filters,
**Fig. 2.** Signals of selective sweeps. Significance $-\log_{10}(P\text{-value})$ for all SNPs for (a) REHH and (b) iHS; QQ plots (observed vs. expected $P$ values) for (c) REHH and (d) iHS. Panels c and d also show the $P$ value distribution with sweep loci removed (teal). Dashed lines: Bonferroni significance (0.05 level).

<table>
<thead>
<tr>
<th>Chromosome</th>
<th>Gene</th>
<th>Gene annotation</th>
<th>Test</th>
</tr>
</thead>
<tbody>
<tr>
<td>2</td>
<td>PFB0935w</td>
<td>Cytoadherence-linked asexual protein 2</td>
<td>iHS</td>
</tr>
<tr>
<td>4</td>
<td>PFD0260c</td>
<td>Sequestrin</td>
<td>iHS</td>
</tr>
<tr>
<td>4</td>
<td>PFD0735c*</td>
<td>Conserved <em>Plasmodium</em> protein, unknown function</td>
<td>iHS</td>
</tr>
<tr>
<td>4</td>
<td>Intergenic</td>
<td></td>
<td>REHH</td>
</tr>
<tr>
<td>6</td>
<td>PFF1335c</td>
<td>4-methyl-5-(B-hydroxyethyl)-thiazol monophosphate biosynthesis enzyme</td>
<td>Both</td>
</tr>
<tr>
<td></td>
<td>PFF1350c</td>
<td>Acetyl-coenzyme a synthetase</td>
<td></td>
</tr>
<tr>
<td></td>
<td>PFF1365c</td>
<td>HECT-domain (ubiquitin transferase), putative</td>
<td></td>
</tr>
<tr>
<td></td>
<td>PDD1460c</td>
<td>Conserved <em>Plasmodium</em> protein, unknown function</td>
<td></td>
</tr>
<tr>
<td></td>
<td>PFF1470c</td>
<td>DNA polymerase epsilon, catalytic subunit a, putative</td>
<td></td>
</tr>
<tr>
<td>7</td>
<td>PF07_0035</td>
<td>Cg1 protein</td>
<td>Both</td>
</tr>
<tr>
<td>7</td>
<td>PF07_0036</td>
<td>Cg6 protein</td>
<td></td>
</tr>
<tr>
<td>7</td>
<td>PF07_0038*</td>
<td>Cg7 protein</td>
<td></td>
</tr>
<tr>
<td></td>
<td>MAL7P1_28*</td>
<td>pfcr; chloroquine resistance transporter</td>
<td></td>
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<tr>
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<td>PF07_0042</td>
<td>Conserved <em>Plasmodium</em> protein, unknown function</td>
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<tr>
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<td>PF07_0053</td>
<td>Conserved <em>Plasmodium</em> protein, unknown function</td>
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</tr>
<tr>
<td>11</td>
<td>PF11_0344</td>
<td>Apical membrane antigen 1 precursor, AMA1</td>
<td>iHS</td>
</tr>
<tr>
<td>13</td>
<td>PF13_0074</td>
<td>Surface-associated interspersed gene 13.1 (SURFIN13.1)</td>
<td>iHS</td>
</tr>
<tr>
<td>13</td>
<td>PF13_0201*</td>
<td>Sporozoite surface protein 2</td>
<td>iHS</td>
</tr>
<tr>
<td>14</td>
<td>PF14_0726</td>
<td>Conserved <em>Plasmodium</em> protein, unknown function</td>
<td>iHS</td>
</tr>
</tbody>
</table>

*aGenes that are no longer significant if double Bonferroni correction applied.*
missing SNP genotypes were imputed for use in the selective sweep search, with 12,885 SNPs successfully imputed.

$F_{ST}$ was calculated using the method of Weir and Cockerham (1984); statistical significance was calculated by permuting population labels. iHS was computed according to the method of Voight et al. (2006) and REHH according to the method of Sabeti et al. (2002). Recombination maps for iHS were generated with the software package LDhat (McVean et al. 2002), using the program interval with a block penalty of 5.0. As both of these tests do not tolerate missing data, SNPs were imputed with PHASE 2.1.1 (Stephens et al. 2001; Stephens and Scheet 2005). A total of 7,486 fully imputed SNPs were polymorphic among the 70 individuals.

REHH and iHS scores were normalized in 20 equal-sized bins of derived allele frequency, with ancestral alleles inferred from a $P$. reichenowi hybridization (van Tyne et al. 2011). Standardized iHS and REHH scores were converted to two-tailed $P$ values using a normal distribution. Genome-wide significance was calculated 1) separately without correction for the two tests and 2) jointly across both tests. As the two tests are not independent, a Bonferroni correction for multiple testing is quite conservative. To test whether long haplotype signals could come from differences in sample preparation, iHS scores were calculated separately for direct and culture-adapted samples.

**Supplementary Material**

Supplementary methods, notes, figures S1–S5, and tables S1 and S2 are available at Molecular Biology and Evolution online (http://www.mbe.oxfordjournals.org/).

**Acknowledgments**

The authors thank colleagues at the MRC Unit in The Gambia including Michael Walther and Omar Janha for support in sample collection and processing. They thank colleagues at the University Cheikh Anta Diop and L’Hopital Aristide le Dantec for support in sample collection and processing. This study was supported by the Bill and Melinda Gates Foundation, the Ellison Medical Foundation, the ExxonMobil Foundation, NIH Fogarty, the Broad Institute, and NIAID. A.A.-N. and D.J.C. were supported by funding from the UK Medical Research Council. D.J.P. was supported by an NSF Graduate Research Fellowship.

**References**


