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Population genomics studies identify signatures of global dispersal and drug resistance in *Plasmodium vivax*

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Urls. Malaria Research and Reagent Resource Repository (MR4), http://www.malaria.mr4.org/; PlasmoDB, http://www.plasmodb.org/; Broad *Plasmodium* 100 Genomes project, https://olive.broadinstitute.org/projects/plasmodium_100_genomes.

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Abstract

Plasmodium vivax is a major public health burden, responsible for the majority of malaria infections outside Africa. We explored the impact of demographic history and selective pressures on the *P. vivax* genome by sequencing 182 clinical isolates sampled from 11 countries across the globe, using hybrid selection to overcome human DNA contamination. We confirmed previous reports of high genomic diversity in *P. vivax* relative to the more virulent *Plasmodium falciparum* species; regional populations of *P. vivax* exhibited greater diversity than the global *P. falciparum* population, indicating a large and/or stable population. Signals of natural selection suggest that *P. vivax* is evolving in response to antimalarial drugs and is adapting to regional differences in the human host and the mosquito vector. These findings underline the variable epidemiology of this parasite species and highlight the breadth of approaches that may be required to eliminate *P. vivax* globally.

P. vivax causes an estimated 15.8 million clinical malaria cases each year¹ but remains understudied in comparison to *P. falciparum* because of its lower mortality and lack of a continuous *in vitro* culture system. Small-scale surveys of samples collected from human patients ('isolates') in the last century and adapted for propagation in splenectomized monkeys have indicated higher genetic diversity for *P. vivax* relative to *P. falciparum* and discordant phylogeographical histories for the two species², likely caused, in part, by the unique ability of *P. vivax* to persist in a dormant state in the liver for months or years. This capacity for dormancy has enabled *P. vivax* to historically cause infections in Europe and North America, where the temperate climate disfavors year-round transmission by mosquitoes³.

We used hybrid selection⁴ to enrich parasite DNA in blood samples heavily contaminated with human DNA obtained from patients infected with *P. vivax*, collected as part of the International Centers of Excellence for Malaria Research program⁵. We sequenced 182 isolates from 11 countries across the globe (Fig. 1) and obtained high sequencing depth and quality for the majority of samples. Hybrid selection using synthetic baits (Online Methods and Supplementary Fig. 1) increased the fraction of parasite DNA in all but four samples, with the greatest enrichment in samples obtained starting with <1% parasite DNA (Fig. 2). We supplemented these new data with published sequences from patient isolates and monkey-adapted laboratory lines for a final data set of 195 *P. vivax* genomes

(Supplementary Table 1). We aligned reads to the Salvador I reference genome assembly⁶ and 'called' SNP variants as described in Supplementary Figure 2. Because *P. vivax* is haploid in the human host, we used rates of heterozygous variant calls to classify 122 of the newly sequenced isolates as single-genotype infections and inferred the remaining 41 samples with sufficient coverage as multigenotype ('complex') infections (Online Methods and Supplementary Fig. 3). We restricted our population genetic analyses of diversity and divergence to single-genotype isolates.

We performed a principal-component analysis (PCA; Fig. 3a and Supplementary Fig. 4) on the SNP variants to explore the global population structure of P. vivax. Similar to a recent microsatellite study⁷, the major axis of differentiation divides the New World from Old World samples. Given recent evidence for an African origin of *P. vivax*⁸, it has been proposed that New World *P. vivax* populations had been founded during the colonial era by a now-extirpated European parasite population⁹. The second and third principal components define a distinct Papua New Guinea (PNG) cluster and a cluster of isolates from Mexico, respectively. We estimated the fraction of the genome exhibiting identity by descent among pairs of samples from geographical regions harboring at least 12 single-genotype isolates (Online Methods) and predicted that isolates sampled from Mexico have recent common ancestry (Supplementary Fig. 5), explaining their unusual clustering. Malaria in Mexico has steadily declined over the last decade to <10,000 cases¹, and the circulation of genetically similar P. vivax may improve prospects for disease control. In contrast to the P. vivax population in Mexico, the PNG population is very diverse from other *P. vivax* populations. Our data support a set of SNPs that differentiate P. vivax populations from those in the Americas, Asia and PNG, with applications for extending the ability to identify the source of imported infections ¹⁰ via SNP barcoding ¹¹ or other genotyping methodologies.

Maximum-likelihood phylogenetic analysis yielded a tree concordant with the PCA (Fig. 3b), highlighting the distinctiveness of the PNG population (alternative reference genome maximum-likelihood trees with bootstrap values are available in Supplementary Fig. 6). We also observed a cluster of African (Mauritania and Madagascar) isolates that grouped closely with Indian isolates midway between the clades from Asia and the Americas. Previous studies of *P. vivax* mitochondrial diversity have identified shared or similar haplotypes among small numbers of sequenced African and Indian/Pakistani samples^{9,12}, suggesting that P vivax was potentially reintroduced to Africa from South Asia by colonial seafarers after its virtual extirpation in Africa owing to the spread of the infection-deterring Duffynegative mutation in human populations¹³. Given that the nuclear genome undergoes sexual recombination, the present data sets potentially offer a more sensitive history of gene flow between populations than mitochondrial data. Admixture analysis of the nuclear SNP data (Fig. 3c and Supplementary Fig. 7) suggests that the isolates from India and Africa exhibit a highly heterogeneous ancestry relative to isolates from other populations, and we inferred that they may derive from both New and Old World lineages. This finding adds support to the hypothesis that contemporary African and South Asian P. vivax populations are genetically similar and suggests that South Asian P. vivax populations may have genetically mingled with European ('New World') lineages during the colonial era or could reflect ancient connections between human populations in the eastern Mediterranean, the Middle East and the Indian subcontinent inferred from admixture signals in human genetic data 14.

The transatlantic slave trade that occurred between the sixteenth and nineteenth centuries continues to shape the contemporary epidemiology of malaria in South America. Genotyping the human Duffy mutation identified no homozygous samples in our collection (Supplementary Table 1) but identified ten samples from Colombian individuals that were heterozygous, five of whom self-identified as having African ancestry. In this setting of non-African parasites infecting people of partial African descent, we observed the dominance of disease susceptibility conferred by the wild-type Duffy allele. Further, we found no evidence of increased copy number of the *P. vivax* Duffy-binding protein gene hypothesized to overcome the barrier to infection presented by Duffy negativity in Madagascar¹⁵ (Supplementary Table 1). South American *P. vivax* epidemiology reflects recent contact between humans and parasites introduced to this region from disparate geographical locations.

Our previous studies of P vivax have highlighted its high diversity in comparison to P falciparum, suggesting a larger and/or more stable effective population size, but were limited by sample size². In this expanded data set, we found that even the least genetically diverse P vivax subpopulation in our study was more diverse than a sampling of diverse P falciparum isolates (Fig. 4). All of our sampled P vivax populations were significantly more diverse than P falciparum (P< 0.001, Wilcoxon signed-rank test), with Old World populations showing nearly twice as much nucleotide diversity. The population samples from Myanmar and Thailand were as diverse as the PNG samples, suggesting that the high divergence of the PNG population is due to protracted genetic isolation and not an elevated mutation rate. The most diverse genes are antigenic, such as members of the merozoite surface protein 3 (MSP3)¹⁶ and MSP7 multigene families and the serine-repeat antigen family (SERA)¹⁷, genes important for immune evasion (Supplementary Fig. 8).

To explore the genomic profile of divergence and selection among P. vivax populations, we calculated the fixation index (F_{ST}) across each chromosome for pairs of well-sampled populations separated by differing geographical distances. The divergence profile we observed in a comparison of Old World versus New World single-genotype samples (Fig. 5a) was not generally similar to the profile that we observed in a comparison of samples from PNG and Thailand. Indeed, there were some notable differences suggesting population-specific selection pressures (Fig. 5b and Supplementary Table 2). Among the genes exhibiting similar signals of divergence for both the proximal and distal geographical comparisons were two loci associated with antifolate drug resistance in *P. falciparum*: *DHPS* (dihydropteroate synthase) and *DHFR-TS* (dihydrofolate reductase–thymidylate synthase). DHPS and DHFR-TS genes also reside in regions of heightened linkage disequilibrium (Fig. 5c,d), suggesting that they have been subjected to selective sweeps. These results indicate that sulfadoxine, pyrimethamine and other antifolate drugs constitute the most important recent evolutionary pressure on P. vivax parasite populations. Application of a McDonald-Kreitman test to P. vivax genes with orthologs in the sister taxon Plasmodium cynomolgi¹⁸ of Southeast Asian macaques identified strong signals of selection on other genes, including several associated with red blood cell invasion, for example, MAEBL (merozoite-adhesive erythrocytic binding protein; Supplementary Fig. 9 and Supplementary Table 3). Other notable genes identified as potentially adapting to regional differences in the human host or

the mosquito vector, or due to the influence of other environmental variables, are indicated in Table 1 and Supplementary Figures 10–15.

We also observed a divergence signal on chromosome 12 enhanced in the New World versus Old World comparison relative to the PNG versus Thailand comparison. The divergence signal was centered on a surface-expressed gamete gene, Pvs47 (Fig. 5e), orthologous to a locus in P. falciparum (Pfs47) that is a strong determinant of successful evasion of the mosquito JNK immune response¹⁹ and that was likely subjected to selection as *P. falciparum* adapted to New World vectors during the colonial era²⁰. New World anopheline vectors belong to the subgenus Nyssorhynchus, which diverged from the lineage leading to Old World anopheline mosquitoes approximately 100 million years ago²¹, and likely presented comparable evolutionary challenges to both P. falciparum and P. vivax after human-mediated introduction of malaria parasites to the New World in the last 500 years. Concordant with the hypothesis that Pvs47 was subject to strong selection upon the arrival of P. vivax in the New World, we observed considerably reduced haplotype diversity in Mexican, Colombian, Brazilian and Peruvian P. vivax populations in comparison with Old World populations (Fig. 5f), a pattern that was not exhibited by *DHFR-TS* or *DHPS* (Supplementary Fig. 16) but which has been observed previously in *P. falciparum* for *Pfs47* (ref. 22). Despite reduced haplotype diversity and strong divergence between populations, Pvs47 did not exhibit heightened linkage disequilibrium (Fig. 5e and Supplementary Fig. 14), suggesting that it was subject to selection less recently than the *DHFR-TS* and *DHPS* drug resistance loci.

Our global survey of genomic diversity in *P. vivax* provides insights into the population structure, demographic history and selective pressures acting upon this species, contextualizing and extending previous, smaller-scale observations². Just as malaria has influenced the human genome²³, this *P. vivax* global genomic data set reflects ongoing evolutionary interactions with the parasite's human and mosquito hosts, interactions that are likely to intensify as disease elimination efforts escalate.

Online Methods

Sample collection, hybrid selection and DNA sequencing

We collected venous blood samples from *P. vivax*-infected patients after obtaining informed consent and following ethical approval at the local institutional review board of each participating site. Details of the sites are available in the Supplementary Note. The monkey-adapted *P. vivax strains* Panama (MRA-343), Vietnam ONG (MRA-341) and Nicaragua I (MRA-340) were obtained from the Malaria Research and Reference Reagents Resource (see URLs). We extracted genomic DNA from each sample and used a modified nested PCR method targeting the multicopy small-subunit ribosomal gene of *P. vivax* for species confirmation³³. We constructed 200-bp-insert Illumina libraries for each isolate and performed hybrid selection on each sequencing library using genome-wide baits to enrich parasite DNA, with synthetic oligonucleotide baits to test maximum possible enrichment, as described⁴. We sequenced the hybrid-selected libraries with 101-bp paired-end reads in indexed batches of 12 on Illumina HiSeq 2000 or 2500 lanes and evaluated parasite DNA enrichment by comparing the fraction of HiSeq reads mapping to the *P. vivax* reference

genome assembly to the mapping fraction observed in low sequencing coverage generated on the MiSeq platform.

Human Duffy antigen/chemokine receptor and P. vivax Duffy-binding protein genotyping

We determined the genotype of the human Duffy antigen/chemokine receptor (DARC) of *P. vivax*-infected patients by using Burrows-Wheeler aligner (BWA)³⁴ to align fastq files of 177 sequenced isolates against the DARC nucleotide sequence (human genome Build 38) and visualizing read alignments. These were classified as either Duffy homozygous positive, homozygous negative or heterozygous according to the presence of a point mutation in the promoter region (a T>C mutation at position –33 in the GATA box). The *in silico* results were validated using a modified wet-lab assay³⁵ (Supplementary Table 4 and Supplementary Note). We mapped all paired-end reads for each isolate to the *P. vivax* Salvador I reference genome and used the indel detection software Pindel³⁶ to detect *P. vivax* Duffy-binding protein (*DBP*) gene duplications. We validated these results by PCR amplification of flanking regions of *DBP*³⁷.

Sequence processing and alignment

We aligned reads from all isolates to the Salvador I reference genome assembly using BWA³⁴ with default parameters and processed the resulting alignments using SAMtools³⁸ and Picard v1.66. We obtained sequence data for 12 previously published *P. vivax* isolates from the Sequence Read Archive, converted them into fastq format and aligned them to Salvador I. We used REALPHY³⁹ software to assess whether our choice of reference genome assembly biased variant calls used for downstream analyses, using two alternative *P. vivax* reference genomes, North Korean and Mauritania I (ref. 2) (Supplementary Fig. 6).

SNP discovery and quality filtering

We used the Genome Analysis Toolkit (GATK) UnifiedGenotyper⁴⁰ to identify SNPs in each isolate according to GATK best practices⁴¹ as part of a SNP identification pipeline (Supplementary Fig. 2). We used two parameters to exclude low-quality sites: GQ 40 and QUAL 30. We filtered SNP variants in single and complex infections according to separate criteria and removed all heterozygous calls from the single-genotype data set (Supplementary Fig. 2). After merging variant calls from all isolates, we removed calls from sites exhibiting a low call rate across the sample set and sites occurring in members of the highly repetitive *P. vivax* variant interspersed repeat (*vir*) multigene family.

Single-genotype versus multigenotype infection classification

To distinguish legitimate heterozygous variants in infections containing multiple genotypes from spurious heterozygous calls, we identified a trusted set of variants exhibiting high-quality homozygous calls for two or more segregating alleles in individual samples. This curated homozygous reference variant set was then used to measure the prevalence of heterozygous SNPs for each isolate, such that only in mixed infections should multiple alternative genotypes be observed. We classified samples as containing more than one genotype if they exhibited greater than 1,236 heterozygous variants (Supplementary Fig. 3).

Population structure

We used a set of filtered variant calls without heterozygous polymorphisms to visualize relationships between *P. vivax* populations using PCA implemented in the SNPRelate R library⁴². We used a combination of two and three eigenvectors to represent the structure of the polymorphism in the population. A more conservative PCA, limited to high-quality (at least 1 million callable sites) single-genotype isolates, was used to confirm the topology (Supplementary Fig. 4). We constructed maximum-likelihood trees using the filtered variant call set limited to homozygous polymorphisms in high-quality *P. vivax* isolates using RAxML⁴³ and performed admixture analysis with the Admixture software package⁴⁴ using the full set of homozygous filtered variant SNPs.

Genetic diversity

We calculated genome-wide nucleotide diversity (π) and Tajima's D from high-quality, single-genotype isolates (Supplementary Table 1) using R and Python custom scripts to tabulate all classes of segregating sites using filtered homozygous variant calls. In addition, we calculated π for 574 one-to-one high-quality orthologs in P falciparum and P vivax. We calculated F_{ST} using the VCFTools package⁴⁵. We identified identical-by-descent genomic regions in pairwise comparisons of isolates using a hidden Markov model analysis of SNP variants⁴⁶.

Tests of neutrality/selection

We used data for 81 high-coverage, single-genotype infection isolates to search for signals of natural selection using a McDonald-Kreitman test and the *P. cynolmolgi* B strain reference genome as an outgroup⁴⁷. We created coding-sequence alignments using *P. vivax* population sequence and the single-copy orthologs between species for 2,254 genes using the program MUSCLE⁴⁸. We performed a McDonald-Kreitman test on the observed polymorphic and divergent synonymous and nonsynonymous mutations in each alignment and evaluated significance using a two-tailed χ^2 test to obtain a *P* value. A *q*-value error correction was applied to the set of 2,254 gene tests to correct for multiple sampling⁴⁹. We also calculated *a* for each gene, a summary statistic that estimates the proportion of substitutions fixed by natural selection⁵⁰, using a custom script.

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

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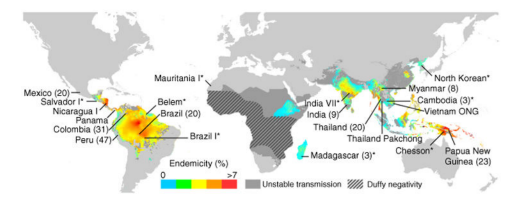


Figure 1.
Country of origin for 195 *P. vivax* patient isolates indicated on a map of *P. vivax* malaria endemicity. Asterisks mark the five published monkey-adapted reference genomes Brazil I, Salvador I, Mauritania I, North Korean and India VII and the published Belem and Chesson strains. The four additional monkey-adapted strains, Vietnam ONG, Nicaragua I, Panama and Thailand Pakchong, sequenced as part of this study are also indicated. Numbers of clinical isolates are given in parentheses. Lines do not represent accurate within-country locations; locations are listed in supplementary table 1. The map was adapted from the Malaria MAP Project²⁴

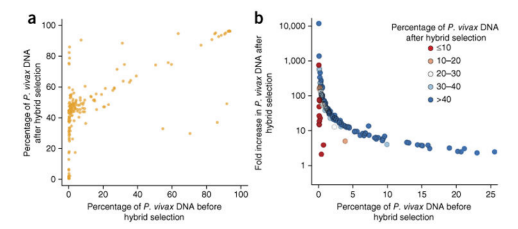


Figure 2. Enrichment of *P. vivax* DNA extracted from clinical samples using the hybrid selection method. (a) Enrichment of parasite DNA measured as the percentage of total sequence reads alignable to the *P. vivax* reference genome before versus after hybrid selection. Enrichment was variable but nearly universal. (b) Fold enrichment of parasite DNA, as compared to the percentage of alignable reads before hybrid selection. Hybrid selection resulted in the greatest fold enrichment for samples containing the lowest amount of parasite DNA before hybrid selection.

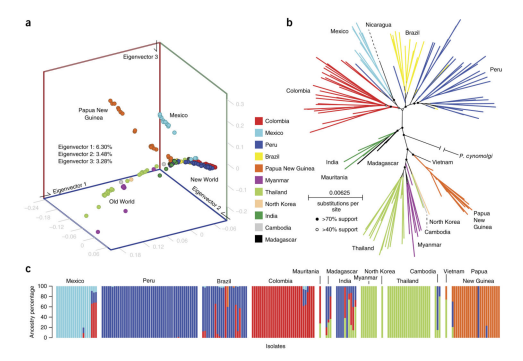


Figure 3. Global population structure of P vivax. (a) Three-eigenvector PCA using variation data from all P vivax isolates. Each isolate is colored according to its origin, and the percentage contribution of each eigenvector is indicated. (b) A maximum-likelihood phylogenetic tree with 50 bootstraps computed through alignment to the Salvador 1 reference genome. Colors and labels correspond to those in $\bf a$, and the sister taxon to P vivax, P cynomolgi, was included as an outgroup. Select internal nodes are highlighted with open and closed circles, representing >40% and >70% bootstrap support, respectively. (c) Admixture analysis of the full set of variation data from 195 isolates using 100 bootstraps. Ancestry for each isolate was assessed according to an optimized cluster value of K = 5 (see supplementary Fig. 7 for admixture plots under six different K cluster values). Colors correspond to the subpopulations seen in $\bf a$ and $\bf b$ and were assigned to the five most deeply sampled subpopulations of P vivax.

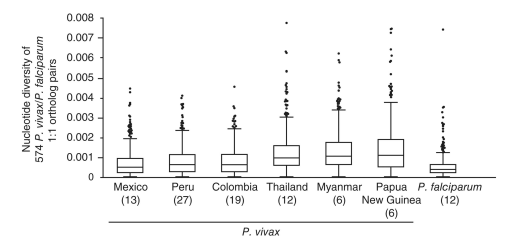
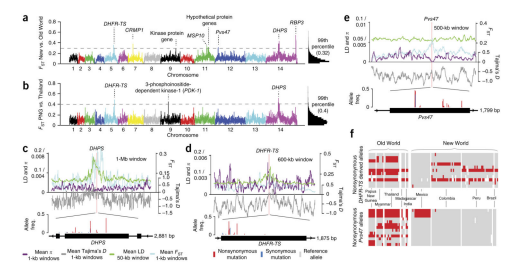


Figure 4. Diversity in orthologs across six P. vivax subpopulations and P. falciparum. Box-and-whisker plots of nucleotide diversity (π) calculated from 574 one-to-one orthologs in single-infection, high-quality isolates from Mexico, Colombia, Peru, Thailand, Myanmar and Papua New Guinea subpopulations are shown. The center and ends of the boxes represent the 25th, 50th and 75th percentiles of π values for each subpopulation, and the whiskers on each plot represent the range of π within 1.5 times the interquartile range of the lower and upper quartiles. Data points beyond the range of the whiskers are plotted as black dots. For P. falciparum, π was calculated from orthologs in a population of 12 African isolates previously sequenced at the Broad Institute (see URLs). P. falciparum was significantly less diverse than all plotted P. vivax populations (P< 0.001) using both a paired Student's t test and a Wilcoxon signed-rank test.

Figure 5.

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Signals of selection in P vivax subpopulations. (a) Genome-wide view of the divergence (F_{ST}) between Old World and New World populations calculated from 1-kb windows. Plotted is a moving average across each of the 14 chromosomes, with a histogram of F_{ST} values on the right. The 99th percentile of the distribution of 1-kb windows forms a cutoff for outliers, which are labeled with the nearest gene annotation to the peak F_{ST} . (b) F_{ST} comparison as in **a** between the Thailand and Papua New Guinea populations. (c-e) Detailed plots for DHPS (c), DHFR-TS (d) and Pvs47 (e), overlaying F_{ST} values from **a** and **b**, the collapsed estimate of mean linkage disequilibrium (LD) calculated across a 50-kb window for each nucleotide, a moving average of π in 1-kb windows for isolates from Southeast Asia and Tajima's D in 1-kb windows. The location of each gene is shown (pink bar) above a schematic of the gene. Homozygous variants within coding sequences are indicated, including nonsynonymous changes, synonymous changes and deletions. Polymorphism

columns are scaled by their allele frequency in the global population. (f) Haplotype map for the *P. vivax* genes *DHFR-TS* and *Pvs47*. Red blocks denote nonsynonymous changes, gray blocks indicate the presence of the reference allele and white blocks show regions of missing

data. Nonsynonymous sites in *DHFR-TS* are limited to derived alleles.

Table 1
Notable *P. vivax* genes with their associated population genetic statistics

Gene name (identifier)	MK test z score	New World/Old World F _{ST} z score	Diversity (π)	Function
Mosquito transmission				
TRAP-like protein (<i>TLP</i> , PVX_113965)	-0.723	6.82	0.001	Important for cell traversal of invasive sporozoite stage ²⁵
Cysteine-repeat modular protein-1 (<i>CRMPI</i> ; PVX_099005)	-0.164	7.27	0.0016	Expressed in both vertebrate and mosquito hosts for host tissue targeting and invasion 26
Red blood cell invasion				
Merozoite-adhesive erythrocytic binding protein (<i>MAEBL</i> ; PVX_092975)	-3.958	4.04	0.0011	Erythrocyte-binding-like protein involved in host receptor recognition in <i>P. falciparum</i>
Merozoite surface protein 10 (<i>MSP10</i> ; PVX_114145)	1.178	6.10	0.0019	Single-copy gene with high human antibody reactivity ²⁷ ; possible vaccine candidate ²⁸
Drug resistance				
Phosphoinositide-dependent protein kinase 1 (<i>PDK1</i> ; PVX_091715)	0.0518	-0.109	0.0009	Associated with artemisinin resistance pathway in <i>P. falciparum</i> ²⁹ ; exhibited strong divergence between PNG and Old World <i>P. vivax</i> in this study
Phosphatidylinositol 3 kinase (<i>PI3K</i> ; PVX_080480)	0.899	0.537	0.0009	Associated with artemisinin resistance pathway in <i>P. falciparum</i> ²⁹ ; showed high polymorphism in our global set of <i>P. vivax</i> isolates
Multidrug resistance 1 gene (<i>MDR1</i> ; PVX_080100)	-3.150	3.41	0.001	Implicated in <i>P. vivax</i> drug resistance ³⁰ ; exhibited high divergence in this study
Chloroquine resistance transporter (<i>CRT</i> ; PVX_087980)	-0.101	1.61	0.0009	Ortholog in <i>P. falciparum</i> implicated in chloroquine resistance ³¹ ; exhibited high polymorphism in some <i>P. vivax</i> Thailand samples in this study
Kelch12 propeller domain gene (<i>K12</i> ; PVX_083080)	-1.486	1.65	<0.0001	Ortholog in <i>P. falciparum</i> implicated in artemisinin resistance ³² ; highly conserved in our global set of <i>P. vivax</i> isolates

Some of the genes listed were identified as potentially adapting to regional differences in the human host or mosquito vector or due to the influence of other environmental variables because of a significant McDonald–Kreitman (MK) test, P_{ST} score or both. McDonald–Kreitman test results and the New World/Old World P_{ST} values for each gene are shown as z scores, and putative functions are given for the genes in P_{ST} values for each gene are shown as z scores, and putative functions are given for the genes in P_{ST} vivax where known. The P_{ST} vivax kelch12 propeller domain gene, the ortholog of an important drug resistance locus in P_{ST} falciparum, did not appear to be under selective pressure in our global set of P_{ST} vivax isolates and was also included.