Molecular Ecology (2014) 23, 1979-1993

Patterns of selection on *Plasmodium falciparum* erythrocyte-binding antigens after the colonization of the New World

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Abstract

Pathogens, which have recently colonized a new host species or new populations of the same host, are interesting models for understanding how populations may evolve in response to novel environments. During its colonization of South America from Africa, *Plasmodium falciparum*, the main agent of malaria, has been exposed to new conditions in distinctive new human populations (Amerindian and populations of mixed origins) that likely exerted new selective pressures on the parasite's genome. Among the genes that might have experienced strong selective pressures in response to these environmental changes, the *eba* genes (erythrocyte-binding antigens genes), which are involved in the invasion of the human red blood cells, constitute good candidates. In this study, we analysed, in South America, the polymorphism of three *eba* genes (*eba-140*, *eba-175*, *eba-181*) and compared it to the polymorphism observed in African populations. The aim was to determine whether these genes faced selective pressures in South America distinct from what they experienced in Africa. Patterns of genetic variability of these genes were compared to the patterns observed at two

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housekeeping genes (*adsl* and *serca*) and 272 SNPs to separate adaptive effects from demographic effects. We show that, conversely to Africa, *eba-140* seemed to be under stronger diversifying selection in South America than *eba-175*. In contrast, *eba-181* did not show any sign of departure from neutrality. These changes in the patterns of selection on the *eba* genes could be the consequence of changes in the host immune response, the host receptor polymorphisms and/or the ability of the parasite to silence or express differentially its invasion proteins.

Keywords: adaptation, balancing selection, co-evolution, invasion, malaria

Received 8 August 2013; revision received 29 January 2014; accepted 7 February 2014

Introduction

One way to approach and analyse the adaptation of parasites to new environments is to analyse past events of emergence, for example the colonization of a new host species or a new population (Lee 2002; Pepin *et al.* 2010). During the evolution of human populations, emergence of infectious diseases has been highly recurrent, and there are now several examples of past emergences that might serve as models to analyse how pathogens adapted to new environmental conditions. The malaria agent *Plasmodium falciparum* is one of these.

P. falciparum is a protozoan parasite responsible for the most serious form of malaria, killing almost one million people every year, principally children under the age of five (Snow *et al.* 2005; WHO 2012). The number of clinical cases is estimated between 300 and 500 million per year in the world and is therefore a major public health concern (WHO 2012). During its life cycle, *P. falciparum* successively infects two hosts: a vector host of the genus *Anopheles* (mosquito) and the vertebrate host *Homo sapiens sapiens*.

During its evolutionary history, P. falciparum has emerged and colonized new environments several times (Hume et al. 2003; Tanabe et al. 2010; Yalcindag et al. 2012). One major event of P. falciparum emergence in a new area occurred in South America during the transatlantic slave trade that started 500 years ago from the African continent and during which millions of African slaves were forcedly transferred on the American continent (Conway et al. 2000; Yalcindag et al. 2012). During this colonization, the parasite encountered new environmental conditions: a new human host environment (Amerindian, European, and populations of mixed origins) and a new vector environment. The species complex Anopheles darlingi is the main vector in South America, while the major species of vectors in Africa are species of the complex An. gambiae and An. funestus (Mirabello & Conn 2006).

The environmental changes that occurred during the colonization of South America by *P. falciparum* likely resulted in powerful selective pressures that forced the

parasite to adapt to these new local environmental conditions and evolve towards different phenotypes and genotypes. Among the genes that might have experienced strong selective pressures in response to these environmental changes, the *eba* genes (erythrocyte-binding antigens genes), which are involved in the invasion of the human red blood cells, constitute good candidates.

Invasion of erythrocytes by the malaria parasite P. falciparum is a complex process involving a cascade of specific protein-protein interactions between the merozoite blood stage and the erythrocyte (Chitnis 2001). In the parasite, this involves two protein families: the erythrocyte-binding antigen (eba) (also called the erythrocyte-binding ligand ebl) and the Plasmodium falciparum-reticulocyte-binding Homolog (PfRH) gene families. We focused here on the eba gene family which consists of five divergent genes in P. falciparum of which three produce functional proteins: eba-140 (MAL13P1.60), eba-175 (MAL7P1.176) and eba-181 (PFA0125c) (Sim et al. 1990; Adams et al. 2001; Chitnis 2001). The EBA proteins have two cysteine-rich regions, one located near the N-terminal end of each protein and another near their C-terminal end (Adams et al. 1992, 2001). The N-terminal cysteine-rich region is termed 'Region II' (Adams et al. 1992) and consists of two tandem Duffy-binding-like (DBL) domains (called F1 and F2) that bind to specific receptors at the surface of the red blood cells. The receptors for eba-175 and eba-140 are glycophorin A and C, respectively (Camus & Hadley 1985; Joshua-Tor et al. 2005), while eba-181 binds to a still unknown receptor.

Eba genes are suspected to be under diversifying selection (Baum *et al.* 2003; Verra *et al.* 2006). This selection is likely of various origins. First, polymorphisms at these genes could be maintained by selection because they provide variable binding affinities to the polymorphic host glycoproteins (e.g. glycophorin A and B). Thus, it has been shown that non-synonymous polymorphisms in region II of *eba-140* and *eba-181* affected their receptor specificity (Mayer *et al.* 2002, 2004). Another source of selection could be the

immune system. EBA proteins are known to induce a protective immune response which could in turn select for polymorphic amino acids in the target regions (Tham *et al.* 2012).

Patterns of selection on eba genes were previously studied in Africa and Asia, and the strongest traces of selection were observed for eba-175, thus suggesting that among eba genes, this gene was the most submitted to diversifying selection (Baum et al. 2003; Verra et al. 2006; Amambua-Ngwa et al. 2012). In South America, no studies have been performed so far. Are eba genes submitted to the same type or same strength of selection in this area? May selection have changed following the colonization of the New World? We hypothesize that the differences in host environment experienced by the parasites during the colonization of South America resulted in changes in the strength and types of selection imposed on eba genes. On this continent, human populations are indeed different from those on the African continent (Zietkiewicz et al. 1997; Rosenberg et al. 2002).

To study selection on *eba* genes in South America, we examined the sequence diversity of the *eba* genes in several South American populations and compared it to the patterns of genetic diversity observed in several African populations. Because the human environment encountered by the parasite is heterogeneous on the South American continent, it is likely that *eba* genes may have experienced divergent selection (local adaptation). To take this possibility into account, we analysed the patterns of selection among *P. falciparum* populations living in different host environments. Thus, parasite populations specifically infecting Amerindians, Maroons (South American populations descending directly from African slaves) as well as populations of mixed origin were collected and studied. Patterns of

genetic variability of the *eba* genes were compared to the patterns observed at two housekeeping genes (*adsl* and *serca*) as well as SNPs (single nucleotide polymorphisms) distributed all over the genome to separate adaptive effects from pure demographic effects.

Material and methods

Sample collection

Our study was conducted on two types of sequence data: (i) sequences produced in this study and (ii) sequences retrieved from the literature. This concerned two kind of genes: the *eba* genes and two housekeeping genes considered to evolve mostly neutrally (Tanabe *et al.* 2010): the sarcoplasmic/endoplasmic Ca²⁺ -ATPase (*serca*) gene and the adenylosuccinate lyase (*adsl*).

Regarding the first kind of data, P. falciparum-infected human blood samples were collected from six different countries: two from Africa and four from South America (Fig. 1 and Table S1, Supporting information). In South America and in particular in French Guiana, two populations were collected in villages (Camopi and Trois Sauts) mainly inhabited by Amerindian persons (belonging to the Wayampi and Emerillon tribes), another in a village (Maripasoula) mainly inhabited by descendants of Maroon people (Boni ethny). Maroons were African slaves that escaped and formed independent settlements together. The other P. falciparum populations of South America were collected in areas inhabited by human populations of mixed origins. In Africa, samples were collected in Benin and Madagascar (See Table S1, Supporting information). Infected blood samples were collected either by venous puncture (~500 µL) or by finger-prick (~50 µL). All blood samples were collected after informed consent. Ethical clearance

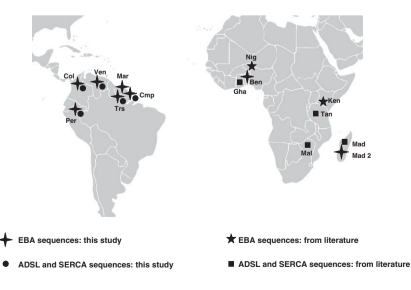


Fig. 1 Geographical location of study sites. In Africa: Ben, Benin; Gha, Ghana; Ken, Kenya; Mad, Madagascar; Mad 2, Madagascar2; Mal, Malawi; Nig, Nigeria; Tan, Tanzania. In South America: Cmp, Camopi (French Guiana); Col, Colombia; Mar, Maripasoula (French Guiana); Per, Peru; Trs, Trois Sauts (French Guiana); Ven, Venezuela. Indications regarding the places where *eba*, *adsl* and *serca* sequences were obtained are provided. For more information, please refer to Tables S1 and S2 (Supporting information).

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was obtained from local ethic committees in each country sampled.

For the sequences retrieved from the literature, this concerned only African populations. *Eba* sequences as well as *adsl* and *serca* sequences were retrieved. Several sources of sequences were used (Baum *et al.* 2003; Verra *et al.* 2006; Tanabe *et al.* 2010). Details regarding the location of the study sites are given in Fig. 1 and Table S2 (Supporting information).

DNA extraction, PCR conditions and sequencing

DNA was extracted using the DNeasy blood and tissue kit (Qiagen, France) according to the manufacturer's recommendations and eluted in 100 μ L of elution buffer per 200 μ L of whole blood or per filter blot.

From each isolate, the *eba-140*, *eba-175* and *eba-181* genes were amplified by a two-step semi-nested PCR using forward and reverse primers (Table S3, Supporting information). PCR amplification was performed on 20 μ L volumes containing 0.2 unit of Expand high fidelity GoTAQ enzyme (Promega, Madison, USA), 5 × Expand reaction buffer, 25 mM MgCl₂, 10 pmol/ μ L of each primer and 10 ng *P. falciparum* DNA. The PCR I cycles were: 94 °C (2 min), [94 °C (30 s), 48 °C for *eba-140*, 50 °C for *eba-175* and 52 °C for *eba-181* (30 s), 72 °C (2 min)] for 35 cycles and 5 min at 72 °C final extension. The PCR II cycles were 94 °C (2 min), [94 °C (30 s), 50 °C for *eba-140* and *eba-181*, 47 °C for *eba-175* (30 s), 72 °C (2 min)] for 35 cycles and 5 min at 72 °C final extension.

Full-length *serca* and *adsl* were amplified by PCR using Takara LA taq polymerase (Takara Bio, Japan) in a reaction mixture, as previously described (Tanabe *et al.* 2010). PCR products were purified by PCR Spin Column Kits (Qiagen) and sequenced by Eurofins (MWG, Ebersberg, Germany).

All of the sequences were manually checked and assembled using CodonCode Aligner software (www. codoncode.com). Sequences were aligned using CLUSTAL w with BIOEDIT software version 7.0.9.0.

Additional genetic information

For the same samples as the ones used in this study (not those obtained from literature), we previously had obtained some genetic information on a set of 272 SNP and 12 microsatellite markers (please see Yalcindag *et al.* (2012) for more information on these markers). This information, especially that regarding the SNPs, will be further used to disentangle the effects of demography from those of selection in shaping the patterns of genetic variability at *eba* genes.

Multiplicity of infections

Blood samples are frequently infected with two or more haploid clones of *P. falciparum*, resulting in the detection of two or more alleles at polymorphic loci. Isolates with more than one allele at any of the 12 microsatellite loci were removed from the analyses [see Table S1 (Supporting information) for more details and Yalcindag *et al.* (2012)]. For Africa, we discarded between 55 to 80% of the samples. For South America, between 20 and 47% of the samples were discarded in each population (Table S1 and S2, Supporting information). A subset of these mono-infected samples was sequenced for each population (Table 1).

Data analyses

The five genes were studied separately. Hence, the analyses described hereafter were performed for each gene. Because we could expect changes of selection patterns between Africa and South America, comparisons between the populations from the two continents were systematically made.

Genetic diversity and population differentiation

For each population, sequence diversity was estimated by π , the average nucleotide diversity, using the software DNASP V5.10. (Rozas *et al.* 2003). For each pair of populations, levels of genetic differentiation were measured by the Wright fixation index, F_{ST} , and departure from 0 was tested using ARLEQUIN ver.3.1 (Excoffier *et al.* 1992). Average values were subsequently computed for all pairs of populations belonging to the same continent (F_{ST} intra).

Tests of neutrality

Several statistics and tests were used to explore the patterns of selection occurring on *eba* genes in the South American populations.

Frequency-spectrum-based tests of selection. First of all, Tajima's *D* statistics (Tajima 1989) were computed for each population and compared to the values observed in the African populations. This statistic detects departures from neutrality in allele frequency distributions by comparing the number of polymorphic sites and the pairwise nucleotide diversity, two measures of within population sequence diversity (Tajima 1989). Under neutrality, these two measures are similar and the Tajima's *D* displays a null value. Under balancing selection, a positive value is expected, while under positive selection, a negative one is observed. Departure from 0 **Table 1** Summary statistics on polymorphism for Region II of Plasmodium falciparum eba genes, serca and adsl genes in six SouthAmerican and eight African populations

Population	Gene	size (bp)	Ν	S	Si	Κ	Sp	NSp	Hd	π	Tajima's D*	Fu & Li's D*	Fay & Wu's H
Africa													
Benin	Eba-140	1620	8	5	3	4	0	5	0.643	0.001	-0.25	-0.5	0.28
	Eba-175	1457	8	9	1	7	0	9	0.964	0.003	1.4	1.1	-0.57
	Eba-181	1714	9	4	2	4	0	4	0.75	0.001	0.08	-0.26	-1
Madagascar	Eba-140	1620	11	4	2	4	0	4	0.691	0.0007	-0.44	-0.4	-0.76
	Eba-175	1457	7	12	3	7	0	13	1	0.004	1.06	0.81	-0.14
	Eba-181	1714	11	4	1	6	0	4	0.855	0.001	0.33	0.41	0.78
Kenya	Eba-140	1620	45	7	5	7	0	8	0.8	0.0009	-0.45	-2.18*	-0.39
	Eba-175	1457	39	15	1	21	0	16	0.9	0.004	1.21	1.13	0.91
	Eba-181	1714	24	2	1	3	0	2	0.5	0.0003	-0.087	-0.66	0.33
Nigeria	Eba-140	1620	24	7	5	8	1	7	0.8	0.001	-0.88	-1.64	-0.06
	Eba-175	1457	30	14	3	14	0	15	0.8	0.003	0.96	0.36	0.21
Ghana	Adsl	1416	37	7	6	8	6	1	0.7	0.0006	-1.42	-3.13**	0.1
	Serca	3687	37	13	9	13	6	7	0.8	0.0004	-1.59*	-2.72**	-2.96
Madagascar 2	Adsl	NA											
	Serca	3687	18	10	2	12	4	6	0.9	0.0008	-0.09	0.49	1.91
Malawi	Adsl	NA											
	Serca	3687	38	14	7	19	4	10	0.9	0.0006	-1.22	-1.6	1.63
Tanzania	Adsl	1416	72	7	5	8	5	2	0.5	0.0005	-1.33	-2.94**	-0.36
	Serca	3687	62	29	19	33	11	19	0.9	0.0007	-1.98**	-3.6***	-2.11
South America													
Camopi	Eba-140	1620	29	3	0	3	0	3	0.431	0.0007	1.37	0.95	0.56
	Eba-175	1457	23	13	6	4	0	13	0.498	0.002	-0.97	-0.93	-3.31
	Eba-181	1714	27	3	0	3	0	3	0.655	0.0008	1.92*	0.96	0.25
	Adsl	1416	18	1	0	2	1	0	0.523	0.0004	1.5	0.67	-0.13
	Serca	3687	13	4	1	5	2	2	0.808	0.0004	0.11	0.33	0.83
Trois Sauts	Eba-140	1620	26	3	0	4	0	3	0.612	0.0009	1.94*	0.97	-0.16
	Eba-175	1457	24	0	0	NA	0	0	NA	0	NA	NA	NA
	Eba-181	1714	24	3	0	3	0	3	0.304	0.0004	-0.2	0.98	-0.78
	Adsl	1416	11	0	0	NA	0	0	NA	0	NA	NA	NA
	Serca	3687	6	2	1	3	1	1	0.733	0.0002	0.12	0.06	-0.26
Maripasoula	Eba-140	1620	13	3	0	2	0	3	0.462	0.0009	1.36	1.08	0.32
	Eba-175	1457	15	11	0	3	0	11	0.6	0.003	1.22	1.44**	1.3
	Eba-181	1714	9	4	0	4	0	4	0.778	0.001	1.15	1.27	-0.7
	Adsl	NA											
	Serca	NA											
Colombia	Eba-140	1620	20	3	0	3	0	3	0.647	0.0008	1.41	1	0.4
	Eba-175	1457	20	12	0	3	0	12	0.568	0.003	1.17	1.46**	-5.64
	Eba-181	1714	20	0	0	NA	0	0	NA	0	NA	NA	NA
	Adsl	1416	20	1	0	2	1	0	0.479	0.0003	1.26	0.65	-0.41
	Serca	3687	20	4	0	2	1	3	0.189	0.0002	-0.95	1.11	-2.69
Peru	Eba-140	1620	29	3	0	3	0	3	0.586	0.0002	2.12**	0.95	0.1
	Eba-175	1457	30	10	0	4	0	10	0.595	0.003	1.75*	1.4**	-0.07
	Eba-181	1714	29	3	1	3	0	3	0.549	0.0006	1.05	-0.25	0.06
	Adsl	1416	32	1	0	2	1	0	0.417	0.0003	1.04	0.59	0.25
	Serca	3687	10	3	0	3	2	1	0.622	0.0003	0.02	1.15	0.8
Venezuela	Eba-140	1620	10	3	0	3	0	3	0.733	0.0009	1.6	1.15	0.44
	Eba-175	1457	9	14	3	6	0	14	0.889	0.004	0.58	0.73	-0.72
	Eba-181	1714	9	1	0	2	0	1	0.5	0.0003	0.98	0.84	0.25
	Adsl	1714	6	1	0	2	1	0	0.6	0.0003	1.44	1.05	0
	1 10000	1110	0	1	0	4	1	0	0.0	0.000 f		1.00	0

N, Number of sequences obtained; S, number of polymorphic sites; Si, number of singleton nucleotide alleles; K, number of haplotypes; Hd, Haplotype diversity; Π, observed average pairwise nucleotide diversity; Sp, Synonymous substitution; NSp, Nonsynonymous substitution; NA, sequence not available. All values calculated using DNASP 5.10 (Rozas *et al.* 2003). *0.05 < *P-value* < 0.1; ***P-value* <0.05; ****P-value* <0.01.

was tested using a permutation procedure implemented by DNASP v.5.10. (Rozas *et al.* 2003).

Second, we computed the Fu and Li's D^* statistics (Fu & Li 1993). Similarly to Tajima's D, this test detects a departure from neutrality by comparing two estimates of sequence diversity: one based on the number of singleton nucleotides and one derived from the number of segregating sites. Under neutrality, Fu and Li's D^* displays a value of 0. A negative value of Fu and Li's D^* indicates an excess of singletons, suggestive of positive selection. At the opposite, a positive value of this D^* parameter suggests a lack of singletons and of rare alleles, which is indicative of balancing selection.

We finally computed the Fay and Wu's H statistic. Contrarily to Tajima's D and Fu and Li's D*, this test was specifically designed to detect recent events of selective sweep. This test compares the nucleotide diversity estimated from expected heterozygosity against nucleotide diversity estimated from the allele frequency of the derived allele at each position (Fay & Wu 2000). In this context, a negative value of Fay and Wu's His indicative of an excess of high-frequency-derived alleles, consistent with recent positive selection. The determination of the identity of the derived alleles was done using the orthologous sequences of the five genes retrieved from the genome of Plasmodium reichenowi, a close relative of P. falciparum infecting chimpanzees (Accession nos: eba-140 (AY572433), eba-175 (AJ251848), eba-181 (AY572435), serca (AB122148) and adsl (AB519183)).

Population genetic differentiation-based tests of selection. Selection may affect the degree of differentiation between populations (Nielsen *et al.* 2005). When a locus shows higher levels of genetic differentiation compared with other loci, this may then be interpreted as evidence for positive selection. In contrast, balancing selection, which tends to maintain more genetic diversity within populations, generates lower $F_{\rm ST}$ values than expected under neutrality (Nielsen *et al.* 2005).

Two approaches based on the analysis of genetic differentiation were used to analyse selection in the South American populations. First, we used the outlier detection approach implemented by Beaumont and Nichols (Beaumont & Nichols 1996) and implemented in the program LOSITAN (Antao *et al.* 2008). This method was implemented using the set of 272 SNP markers previously genotyped in the same South American populations (Yalcindag *et al.* 2012) in addition to the SNPs detected in *adsl, serca* and the three *eba* genes. Outlier values of genetic differentiation at specific loci were detected by running 10 000 coalescent simulations to generate the expected distribution of Wright's coefficient F_{ST} as a function of expected heterozygosity (*He*) which corresponds to the average probability of choosing two different alleles at random from two different demes. The distribution of $F_{\rm ST}$ was obtained by simulating an island model with a distribution centred on the empirical estimates averaged over loci. This average $F_{\rm ST}$ cannot be assumed to be neutral because (initially unknown) selected loci may be included in the computation. We thus first ran LOSITAN to determine a subset of candidate selected loci and then removed them for the computation of the neutral $F_{\rm ST}$. The value obtained is likely to be a better approximation of the neutral $F_{\rm ST}$. The approach is expected to be robust with respect to variation in mutation rate among loci, sample size and departure from mutation/drift equilibrium (Beaumont & Nichols 1996).

Second, using the set of SNPs detected to be neutral in South America with the Beaumont and Nichols (1996)'s method, we performed another analysis to study the patterns of selection on eba genes. Under neutrality, FST measured between several pairs of populations using two independent sets of markers is expected to be correlated and equal considering the two sets of markers display the same mutation rate. Patterns of selection in genes can thus be searched by (i) estimating the slope of the linear regression of the F_{ST} measured using the SNPs of the target genes between pairs of populations with those obtained between the same populations using the set of SNPs identified as evolving neutrally in South America and (ii) comparing the slope to the distribution of slopes observed under neutral expectations. A resampling test was used to test the departure of the slopes obtained with the candidate genes with the expectation under neutrality. Namely, assuming k segregating sites (SNP) were recorded in the target gene, k SNP were extracted randomly from the neutral data set, F_{ST} were computed for each pair of populations from these k SNP and then regressed against the F_{ST} computed from the whole SNP data set. This process was repeated 1000 times to yield an empirical distribution of slopes under the neutrality hypothesis.

Results

Sequence diversity and genetic differentiation

For *eba* genes, sequences obtained and analysed in our study covered 1620 bp of *eba-140*, 1457 bp of *eba-175* and 1714 bp of *eba-181*. For the housekeeping genes, *adsl* and *serca*, sequences covered 1416 and 3687 bp, respectively. The total number of sequences obtained and analysed for each gene and each population is given in Table 1.

Figure 2 shows the summary statistics computed for each gene and averaged over all populations in South

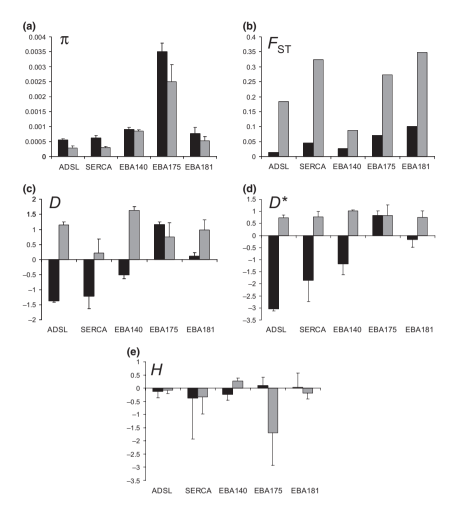


Fig. 2 Comparison between Africa (black bars) and South America (grey bars) of the average values of (a) π , the average nucleotide diversity, (b) F_{ST} between populations, (c) Tajima's *D* index,(d) Fu and Li's *D** index and (e). Fay and Wu's *H* index. Values are given for each gene. For (a), (c), (d), (e), error bars: standard error of the mean.

America and Africa (see also Table 1 for more details on the same summary statistics but computed for each gene and each population). Figure 2a represents the average nucleotide diversity index (π). Overall genes and populations, a significant lower genetic diversity was observed in South America compared with Africa (GLM, Gaussian structure, *P-value* = 0.01642). This decrease in genetic diversity in South America is observed for all genes (Fig. 2a), but the proportion that was lost compared with Africa varied from one gene to another. It was the lowest for *eba-140* followed by *eba-175, eba-181* and finally *adsl* and *serca*, the two housekeeping genes.

Regarding genetic differentiation between populations (Fig. 2b), average $F_{\rm ST}$ values computed between populations in South America were significantly higher than the average $F_{\rm ST}$ observed in Africa (GLM, *P*value = 8.8×10^{-5}). The lowest mean $F_{\rm ST}$ value observed in South America was obtained for *eba-140*. Pairwise $F_{\rm ST}$ values as well as *P*-values of the test of differentiation are given in Table S4 (Supporting information), for each gene.

Tests of neutrality

Frequency-spectrum-based tests of selection. An average positive value of Tajima's D was observed for each gene in South America, while in Africa, negative mean values were observed for all genes except eba-175 and eba-181. Tajima's D statistics observed for each population and each gene are given in Fig. 2c and Fig. 3a as well as in Table 1. Tajima's D values were significantly higher in South America than in Africa for eba-140 (Wilcoxon rank sum test, P-value = 0.009524) and serca (P-value = 0.03175), but not for *adsl* (P-value = 0.333), eba-175 (P-value = 0.9048) and eba-181 (P-value = 0.25). Very similar trends were observed for Fu and Li's D* as shown in Fig. 2d, 3b and Table 1. Note that among South American populations, only one population displayed negative values for both Tajima's and Fu and Li's indices: the one from Camopi for eba-175.

Fay and Wu's H values cannot be interpreted the same way as Tajima's D and Fu and Li's D^* as the parameter H was specifically designed to detect recent selective sweeps. Table 1 reports the values of H for

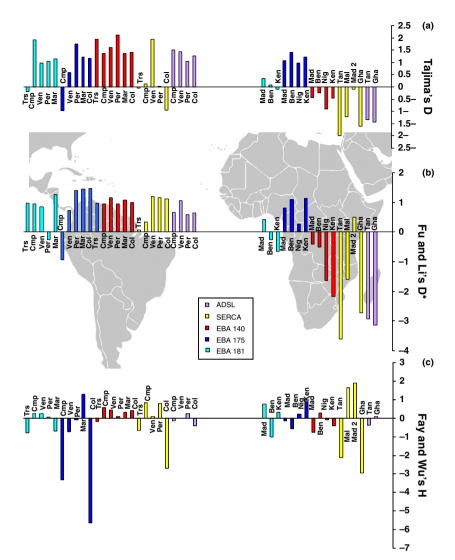


Fig. 3 Comparison between Africa and South America of (a) Tajima's *D* values obtained for each gene and each population, (b) Fu and Li's *D**, and (c) Fay and Wu's *H*. In Africa: Ben, Benin; Gha, Ghana; Ken, Kenya; Mad, Madagascar; Mad 2, Madagascar 2; Mal, Malawi; Nig, Nigeria; Tan, Tanzania. In South America: Cmp: Camopi (French Guiana); Col, Colombia; Mar, Maripasoula (French Guiana); Per, Peru; Trs, Trois Sauts (French Guiana); Ven, Venezuela.

each gene in each population. Although none of them significantly depart from 0, it has to be noted that, in agreement with Tajima's and Fu and Li's indices, a strong negative value was obtained in South America for *eba-175* in the population of Camopi (see Figs 2e and 3c).

 F_{ST} -based tests of selection. Two distinct tests of selection based on the analysis of genetic differentiation between populations were performed. The first one used the outlier approach developed by Beaumont & Nichols (1996) and implemented using a set of 272 SNP markers previously genotyped in the same South American populations (Yalcindag *et al.* 2012) in addition to the SNPs detected in *adsl, serca* and the three *eba* genes. The test (implemented with the software LOSITAN) identified five SNPs that fell above the upper 95% boundary and 12 SNPs that displayed lower estimates of genetic differentiation than expected under neutrality (data not shown). Regarding the SNPs belonging to the *eba* genes, two of three fell below the boundary for *eba-140* (the remaining was very close to the boundary, *P-value* = 0.0365), one of 14 for *eba-175* and none for *eba-181* (of four). For *adsl* and *serca*, all SNPs fell inside the range of $F_{\rm ST}$ values expected under neutrality (Fig. 4).

The second test of selection compared the F_{ST} measured between pairs of populations in South America using the neutral set of SNPs (identified in the former test) and the F_{ST} measured using the SNPs from the studied genes (see Material and Methods for details). Figure 5 shows the correlations obtained between pairwise F_{ST} values computed using the neutral set of SNPs with the F_{ST} computed with the housekeeping (*adsl* and *serca*) or the *eba* genes. The relationships obtained with the F_{ST} computed from the neutral set of SNPs and a

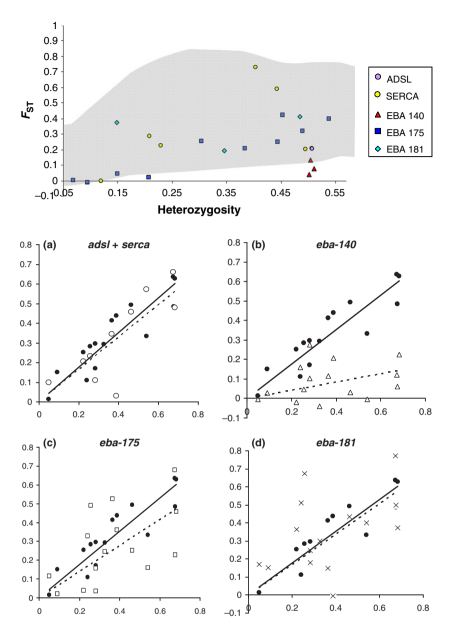


Fig. 4 F_{ST} values plotted against expected heterozygosity (*He*) for the SNPs observed within each gene (*adsl, serca, eba-140, eba-175* and *eba-181*). Grey area: area including 95% of the neutral F_{ST} estimated under an island model with the program LOSITAN and using the total number of SNPs: 272 from Yalcindag *et al.* (2012) + the SNPs obtained in each sampled gene (*adsl, serca, eba-140, eba-175* and *eba-181*).

Fig. 5 Relationships between $F_{\rm ST}$ values estimated between each pair of South American populations using a set of neutral SNP markers and those obtained using: a) *adsl* + *serca* (open circles/dotted regression line), b) *eba-140* (open triangles/dotted regression line), c) *eba-175* (open squares/dotted regression line), d) *eba-181* (crosses/dotted regression line). The relationships observed between the neutral SNPs and neutral microsatellite makers genotyped within the same populations are given for comparison within each panel (black circles/continuous regression line).

set of neutral microsatellite markers genotyped within the same populations (Yalcindag *et al.* 2012) are provided for comparison. As shown, the slopes of the regression line obtained between microsatellites and SNPs are similar to the slopes of the regression observed between neutral SNPs and the candidate genes except for *eba-140* and *eba-175* (Fig. 5) for which resampling tests showed that they display lower slopes than expected under neutrality (Fig. 6 and Material and Methods for more details).

Discussion

The distribution of polymorphism of three genes involved in the invasion of erythrocytes (*eba-140*,

eba-175, eba-181) was studied in several populations of South America to determine how these genes responded to the environmental changes that followed the colonization of this continent by *P. falciparum*. Indeed, during this colonization, *P. falciparum* encountered new environments (in particular new human populations: European, Amerindian and populations of mixed origins), which likely exerted new selective pressures on the parasite.

We focused our analysis on Region II of each gene because this region encodes the erythrocyte-binding domains that bind to specific proteins at the surface of the human red cells. Region II is also known to be antigenic that is to be recognized by the immune system (e.g. Persson *et al.* 2013).

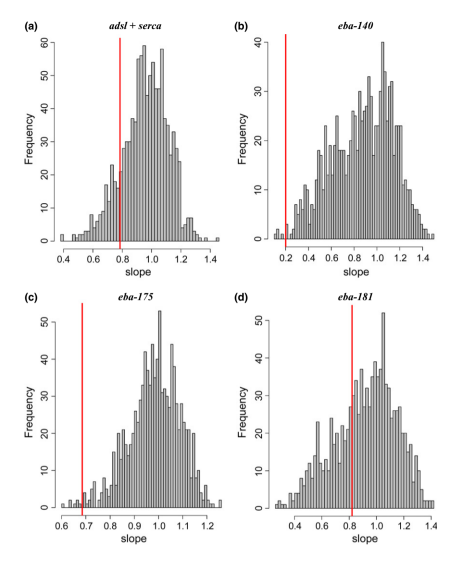


Fig. 6 Distribution of slope values obtained using a randomization procedure of the SNPs to create two independent neutral data sets of the size of the data set used in our study. The slope observed for the observed data set is reported as a red bar. a) The slope observed for *adsl* + *serca*, b) The slope observed for *eba-140*, c) The slope observed for *eba-175* and d) The slope observed for *eba-181*.

Our results extend earlier findings suggesting that the different *eba* genes might be under different selective pressures in Africa but also and, more importantly, show that these selective pressures might have changed in nature and/or intensity following the arrival of *P. falciparum* in South America.

Patterns of selection in Africa

The analysis of polymorphism at genes expected to have evolved neutrally is a way to determine how demography has shaped patterns of genetic variability overall genome. Departures from such patterns at particular loci would then result from selection. In our study, *adsl* and *serca* genes, two genes expected to have evolved neutrally (Tanabe *et al.* 2010), display negative values of Tajima's *D* and Fu and Li's *D** in Africa. This result is consistent with results previously reported showing that low frequency haplotypes are more frequent in the African populations than expected under neutral evolution (Baum *et al.* 2003; Joy *et al.* 2003; Chang *et al.* 2012). Such pattern of genetic variability could be interpreted as the consequence of a recent expansion of the *P. falciparum* population in Africa. According to Chang *et al.* (2012) that analysed the polymorphism of Senegalese strains at a genomic scale, this expansion could have taken place between 20 000 and 40 000 years ago, while Joy *et al.* (2003) estimated that this expansion might have taken place a little more recently in the history of *P. falciparum*, between 200 and 14 500 years ago.

Because demographic history affects polymorphism in the genome the same way, any departure from this pattern could be interpreted as the consequence of selection (Nielsen *et al.* 2005). In our study, *eba-175* displayed a positive value of Tajima's *D* index in all studied African populations. Although none of them were significantly different from 0, the fact that all populations showed similar patterns is consistent with the hypothesis that this gene is submitted to balancing (diversifying) selection, as previously suggested (Baum *et al.* 2003; Verra *et al.* 2006; Amambua-Ngwa *et al.* 2012). This selection was attributed to acquired immunity. Because *eba-175* proteins are the target of immune response and that this response can inhibit invasion, new alleles of this gene would constantly be selected to avoid immune detection thus favouring the maintenance of a high diversity of variants (Baum *et al.* 2003; Verra *et al.* 2006).

Eba-140 showed negative values and eba-181 values close to 0, positive or negative. In the case of eba-140, these negative values were previously regarded as suggestive of the absence of selection on this gene in Africa (Baum et al. 2003; Verra et al. 2006). Although this might indeed be the case, a lower level of diversifying selection compared with eba-175 is another possibility. Indeed, despite negative values, it has to be noted that the Tajima's and the Fu and Li's values computed in the different populations are, on average, higher than for the housekeeping genes in Africa (Wilcoxon rank sum test, *P-value* = 0.057). This could thus be the consequence of balancing selection acting on this gene but at a lower level than for eba-175. Such a hypothesis would be more consistent with observations made in natura: the expression of eba-140 seems to be correlated with that of eba-175 in African populations (Jennings et al. 2007; Gomez-Escobar et al. 2010) and eba-140 is also known to induce a protective immune response and thus to be exposed to the immune system (Tham et al. 2012; Persson et al. 2013). A similar remark can be made for eba-181. The values close to 0 are not expected under neutral evolution and could thus be interpreted as a consequence of immune-mediated selection. Again, the fact that eba-181 is known to induce an immune response in the African populations supports this hypothesis (Ford et al. 2007).

Contrasting patterns of selection in South America

In South America, the patterns of genetic variability observed among populations are highly different from what is observed in Africa: on average, genetic diversity is lower, genetic differentiation between populations is much larger, and an excess of intermediate frequency mutations associated with positive Tajima's and Fu and Li's values is observed.

Part of these differences with the African populations can be attributed to the demographic history of the South American *P. falciparum* populations. It is thought that *P. falciparum* was first introduced into South America 500 years ago with the arrival of Europeans, and its introduction from Africa occurred during the course of the transatlantic slave trade. During this introduction, P. falciparum populations likely experienced severe bottlenecks due to founding events leading to a loss of genetic diversity compared with the African source populations (Yalcindag et al. 2012). This loss is known to concern most preferentially the rare frequency mutations which tend to generate positive Tajima's and Fu and Li's values (Fay & Wu 1999). Regarding genetic differentiation, lower effective population size as well as multiple colonization events could explain the stronger geographical structuring observed between South American populations (Yalcindag et al. 2012). Indeed, it has been shown that the colonization of the New World occurred at least twice independently leading to the establishment of two genetically distinct populations of P. falciparum in different regions of the continent (Yalcindag et al. 2012).

Neutrality tests based on the analysis of the frequency spectrum or genetic differentiation realized on eba genes and their comparison with observations made on housekeeping genes or a neutral set of SNP markers indicate that changes in the nature and/or intensity of selection could also explain some of the patterns of genetic variability observed in the South American populations. In contrast to Africa where eba-175 seems to be under the strongest selective pressures, in South America, evidences of selection are the strongest for eba-140. This gene indeed displays the highest average values of Tajima's and Fu and Li's indices and significantly lower levels of genetic differentiation between populations compared with what was observed at neutral markers. In addition, the loss of genetic variability was the lowest for eba-140 in South America. All these elements suggest that, contrary to what happens in Africa, eba-140 could be under stronger balancing selection in South America than eba-175 or eba-181.

Several nonexclusive hypotheses could be proposed to explain these variations in the relative levels of selection acting on *eba* genes between South America and Africa. First of all, this could be due to changes in the relative levels of expression of these different genes in South America compared with Africa. The most expressed genes are the most submitted to the immuemediated selection.

Variability in the expression of the proteins involved in erythrocyte invasion has now been well documented (see Cortes 2008 and Tham *et al.* 2012 for review). Several experiments with culture adapted parasites or with field isolates have thus revealed extensive variability in the level of *eba* and *PfRH* gene expression. These variations were shown to be correlated with variations in the pathways utilized by *P. falciparum* to invade the red blood cells (Bei *et al.* 2007), and silencing of some genes was shown to be transmitted epigenetically (Cortes *et al.* 2007). An obvious role of this ability to differentially express the proteins involved in erythrocyte invasion proteins is that it may confer flexibility to the parasite to invade different erythrocyte lines including mutant erythrocytes with alterations in the composition of the surface proteins (Cortes 2008), which may vary from one geographical location to another (see e.g. Zimmerman *et al.* 2003). This ability to silence or express different ligands could also play a role in immune evasion. Because merozoite surface proteins are the target of invasion-inhibitory antibodies, switching invasion pathways could mediate escape from the immune system (Cortes 2008). Merozoite ligand expression could thus be under frequency-dependent selection by the immune response (Ford *et al.* 2007).

Several elements support the hypothesis that the relative levels of expression of EBA proteins as well as the invasion pathways used by *P. falciparum* in South America might indeed have changed following its arrival on the continent. Lobo et al. (2004) and more recently Lopez-Perez et al. (2012) have characterized the invasion pathways used by field isolates in different regions of South America and both studies found that, contrarily to what was observed in several African populations (Tham et al. 2012), eba-175-mediated invasion was not the dominant pathway of invasion for the majority of the strains. This could thus explain why eba-175 seems to be less subject to balancing selection in South America than in Africa. In these experiments, nothing was carried out regarding eba-140. For eba-181, Lopez-Perez et al. (2012) have shown that the protein could be a major pathway for certain isolates, but overall was less expressed than eba-175. Some immunological studies also provide elements in favour of a change in the use of the different EBA proteins in South America compared with Africa. Thus, Ford et al. (2007) compared the humoral responses against full-length region II of eba-175 and eba-140 of two human Brazilian populations and a Cameroonian one (nothing was carried to assess the immunoreactivity of the individuals against eba-181 region II). They demonstrated that the Brazilian populations had variable ability to recognize P. falciparum invasion ligands and that these responses were distinct from those of the African population. Thus, while most African individuals reacted strongly and similarly to all region II recombinants, a far lower proportion of individuals reacted against eba-140 and eba-175 recombinants in South America, and the response was more prevalent and more intense for eba-140 than for eba-175 (Ford et al. 2007). These observations suggest therefore that eba-140 could be more highly expressed than eba-175 in South America and that, as a consequence, the immune-mediated selection could be stronger in the American isolates against eba-140 than against eba-175.

Far from us the idea to say that eba-140 is the main pathway of invasion in South America. It does only suggest that, most likely, it could be more expressed than eba-175 and eba-181 and as a consequence could be more subjected to the immune-mediated selection. Our results do not mean either that selection on eba genes in South America is stronger than in Africa. This is even most likely the opposite. Indeed, as shown by Ford et al. (2007), the level of immunity against eba genes in South America is far lower than in the Cameroonian population, a pattern probably due to the fact that in low endemic areas (such as the South American region), the immune system of the host is less frequently boosted by the parasites and so less efficient in comparison with the one in high endemic areas (such as Africa).

Another possibility to explain the variations in the relative levels of selection acting on eba genes in South America compared with Africa could be linked to changes in the genetic variability of the erythrocyte receptors themselves. This could at least explain part of the selection acting on eba-140 and eba-181, for which it was demonstrated that the polymorphisms observed in region II (most of which being nonsynonymous polymorphisms) could change the binding affinities of the protein to their specific receptors (Maier et al. 2009). We could thus imagine that some of these variants were selected and maintained within populations because they provided a better affinity to some of the host receptor variants. Modifications in the relative diversity of the host receptors in the South American human populations compared with the African ones may thus have changed the levels of diversifying selection acting on the different genes. A comparison of the genetic diversity of erythrocyte surface receptors in South America and Africa could provide an answer.

Overall, it is not easy from the different elements provided in the literature (that are often contradictory) to highlight one factor more than another to explain the variations of selection pressures acting on *eba* genes between the New and the Old world. It is very likely the interplay between the host immune response, the receptor polymorphisms and the ability of the parasite to silence or express its invasion proteins that determines the nature and level of selection acting on *eba* genes and hence their level of polymorphism within populations. This is all the more true that the new human environment met in South America by the parasite is not homogeneous.

Amerindian vs. others P. falciparum populations

In our study, two of the *P. falciparum* populations came from villages inhabited mainly by Amerindians

(Camopi and Trois Sauts), another one came from a village inhabited mainly by Maroons and the others originated from human populations of mixed origins. Did this environmental heterogeneity generate local adaptation? In particular, did populations infecting Amerindians experience particular patterns of selection at eba genes? This possibility was considered for several reasons. Amerindian populations are genetically strongly differentiated from populations of African and European origins (Rosenberg et al. 2002; Manica et al. 2005). This differentiation concerns the whole genome and glycophorin genes, the receptors of *eba* genes at the surface of the red cells, are no exception (Tarazona-Santos et al. 2011). In addition, genetic diversity in Amerindian populations is lower than in populations of African origin (Prugnolle et al. 2005a,b). This concerns in particular the genes of the immune system like those of the HLA complex that play a role in the recognition of nonself (Prugnolle et al. 2005a,b), which could thus limit the level of immune-mediated diversifying selection acting on eba genes.

Patterns of genetic variability in the two Plasmodium Amerindian populations are, on average, similar to what is observed for the other South American populations except for eba-175 (see Table 1 and Fig. S1, Supporting information). For this gene indeed, no genetic variability was observed for the population of Trois Sauts and a smaller diversity than in the other populations for Camopi. In addition to that, negative Tajima's *D* and Fu and Li's *D** values were obtained for Camopi, in contrast to what was observed for the other populations for the same gene or for the other genes in Camopi (mostly positive values). The patterns of genetic variability observed in these two Amerindian populations could reflect a phenomenon of recent positive selection acting on eba-175 in these populations. This possibility needs however to be confirmed by more thorough analyses, for instance, by genotyping more genetic markers around the region of *eba-175*.

Conclusion

Our study shows that the level and or type of selection acting on *eba* genes in South America differs from the one acting on the same genes in Africa. A change in the relative expression of *eba* genes and hence of the invasion pathways used by the parasite to infect red blood cells in South America could, at least partly, explain these changes in the patterns of selection. This could also be due to changes in the genetic composition of the receptors expressed at the surface of the erythrocyte or in the host immune system. From what we currently know, we still cannot determine with insurance which factor is responsible for selection, most likely a combination of factors, perhaps not the same for the different genes and certainly variable from one region of the world to another. Further studies should thus be performed to clearly identify these factors.

Acknowledgements

This study was funded by ANR Grant MGANE 07 SEST 012 (Programme Sante-Environnement et Santé-Travail- Malaria Genetic Adaptation to a New Environment), the ANR grant ORIGIN 2012 ICIC, CNRS, IRD, CROUS (Centre Régional des Oeuvres Universitaires et Scolaires), Ministère des Affaires Etrangères, Institut Kurde de Paris. We thank Institut Pasteur and Institut de Veille Sanitaire for the samples sequencing. We also thank the Program of 'Employment of Newly Graduated Doctors of Science for Scientific Excellence' (Grant Number CZ.1.07/2.3.00/30.009) cofinanced from European Social Fund and the state of budget of the Czech Republic who partially supported Erhan Yalcindag. We would like to also thank Masaryk University, Parasitology Research Group, for their support. Samples from Peru were collected within a project funded by the Directorate General for Development Cooperation (DGCD) of the Belgian Government (framework agreement 02, 2003-2007), project 95501 in collaboration with Dr. Alejandro Llanos-Cuentas. DG is supported by DGCD (Framework Agreement 03, 2008-2010, project 910000) and U.S. Public Health Service Grants NIH/NIAID RO1 AI067727 and ICEMR Grant (1U19AI089681). We thank the programme CNRS-Amazonie Ecologie de la Santé. Thanks to Muriel Dietrich for her valuable comments during the preparation of this article.

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E.Y., E.E., F.J.A., F.R., and F.P. designed research; E.Y., C.A., P.D. and F.P. performed research; S.B., L.D., A.A., P.B., U.D., D.F., D.G., A.M., D.M., L.M., O.N., V.V., A.W., B.C., E.L. and C.C. contributed new reagents/ analytic tools; E.Y., V.V., E.E., P.D. and F.P. analyzed data; and E.Y., V.V., E.E., F.J.A., F.R. and F.P. wrote the paper.

Data accessibility

DNA sequences : GenBank Accession numbers; *Eba-140* : KJ419351 - KJ419496, *Eba-175* : KJ419497 -KJ419632, *Eba-181* : KJ419633 - KJ419770, *Adsl* : KJ419771 -KJ419857 and *Serca* : KJ419858 - KJ419915.

SNP dataset, sequences alignments of each gene and sampling details for each population: DRYAD doi:10. 5061/dryad.59645.

Supporting information

Additional supporting information may be found in the online version of this article.

Table S1 Characteristics of the *Plasmodium falciparum* samples collected in Africa and South America, with their geographical coordinates, total sample size and mono-infected sample size.

Table S2 Characteristics of the *Plasmodium falciparum* samples retrieved from literature, with their geographical coordinates, total sample size and mono-infected sample size.

Table S3 Primers used for first and second PCR.

Table S4 Estimates of genetic differentiation (F_{ST}) obtained between pairs of populations and *P*-value of the test of differentiation for each gene.

Fig. S1 Average nucleotide diversity, Tajima's D, Fu and Li's D^* and Fay and Wu's H measured in the *P. falciparum* populations collected in South America.