Host age and expression of genes involved in red blood cell invasion in *Plasmodium falciparum* field isolates

Aida Valmaseda1,2, Quique Bassat1,2, Pedro Aide1, Alfons Jiménez1,4, Aina Casellas2, Sonia Machovo2, Ruth Aguilar2, Betuel Sigauqü2, Virander S. Chauhan6, Christine Langer7, James Beeson7, Chetan Chitnis6, Pedro L. Alonso1,2, Deepak Gaur5 & Alfredo Mayor1,2

*Plasmodium falciparum* proteins involved in erythrocyte invasion are main targets of acquired immunity and important vaccine candidates. We hypothesized that anti-parasite immunity acquired upon exposure would limit invasion-related gene (IRG) expression and affect the clinical impact of the infection. 11 IRG transcript levels were measured in *P. falciparum* isolates by RT-PCR, and IgG/IgM against invasion ligands by Luminex®, in 50 Mozambican adults, 25 children with severe malaria (SM) and 25 with uncomplicated malaria (UM). IRG expression differences among groups and associations between IRG expression and clinical/immunologic parameters were assessed. IRG expression diversity was higher in parasites infecting children than adults (p = 0.022). *eba140* and *ptramp* expression decreased with age (p = 0.003 and 0.007, respectively) whereas *p41* expression increased (p = 0.022). *pfhr5* reduction in expression was abrupt early in life. Parasite density decreased with increasing *pfhr5* expression (p < 0.001) and, only in children, parasite density increased with *p41* expression (p = 0.007), and decreased with *eba175* (p = 0.013). Antibody responses and IRG expression were not associated. In conclusion, IRG expression is associated with age and parasite density, but not with specific antibody responses in the acute phase of infection. Our results confirm the importance of multi-antigen vaccines development to avoid parasite immune escape when tested in malaria-exposed individuals.

By invading host cells, apicomplexan parasites can develop in a rich nutrient environment while being protected from host defenses. In *Plasmodium falciparum*, invasion of red blood cells (RBC) by merozoites is mainly driven by the cooperative function of ligands from the *P. falciparum reticulocyte-binding* homolog (*PfRh*) and the erythrocyte-binding antigen (*EBA*) families1–3. In addition, many ligands that do not belong to these families have been identified as having key roles during RBC invasion4–6. The expression of invasion-related genes (IRG) during schizont stages7,8 is tightly regulated as part of the parasite life cycle8,9. Depending on the nature of the RBC receptor used, the invasion phenotype can be classified as sialic acid (SA)-dependent or SA-independent10. Some invasion ligands have been linked to each of these two phenotypes in culture-adapted parasite lines namely *PfRh1, EBA175*, and *EBA140* to SA-dependent pathway11 and *PfRh2a, PfRh2b* and *PfRh4* to SA-independent pathway12–17. Inverse correlations in expression between genes involved in opposite pathways (*pfhr1* vs. *pfhr2b*) have been observed in natural isolates from Kenya18 and Tanzania19. However, it was observed that the same invasion phenotype does not always involve the same gene expression pattern18.

The essential process of RBC invasion is extremely rapid20, probably for minimizing the exposure of merozoites to host-derived antibodies targeting invasion ligands. This immune recognition can lead to inhibition of the...
parasite replication or killing and therefore constitute a selection pressure on the parasite population\(^a\). Antibodies to a range of merozoite invasion ligands are acquired through exposure to malaria and have been associated with protection from clinical illness\(^a\). In return, the parasite has developed immune evasion strategies through the high polymorphism\(^a\), functional redundancy or gene variant expression of critical invasion ligands\(^a,13,23\). Indeed, the parasites that survive during an infection are thought to express variants corresponding to gaps in the repertoire of host antibodies\(^24\), and variation in invasion ligand use by the parasite during RBC invasion has been shown to mediate evasion of inhibitory antibodies\(^25,26\). Consequently, this immune pressure may modulate IRG expression, as it is hypothesized for eba175\(^27,28\), and changes in EBA175 expression and/or use by the parasite alter the susceptibility to acquired inhibitory antibodies\(^25\). Interestingly, few studies have reported that particular transcripts of IRG correlate with parasitemia, age or endemicity levels\(^28,29\). Moreover, differences have been reported in multiplication rates and invasiveness in natural isolates from Thai patients with severe (SM) and uncomplicated (UM) malaria\(^30,31\). However, no differences have been found between SM and UM patients in other parasite populations regarding multiplication rates, RBC selectivity, invasion phenotypes\(^32,33\) or IRG expression levels\(^18,28\).

The immune escape of parasites caused by exposure to vaccine-induced anti-malarial immunity can be minimized if the IRGs whose expression is modulated by immune pressure are identified\(^40\). Therefore, this kind of studies can help to adapt vaccination strategies based on the predicted change in malaria transmission intensity (and therefore in accumulated exposure) caused by malaria control and elimination campaigns. Here we aimed to study the gene expression levels of IRGs in \(P. falciparum\) isolates collected from Mozambican individuals with different levels of previous exposure to malaria parasites, defined by their age or their antibody responses against \(P. falciparum\) antigens. We hypothesized that anti-parasite immunity acquired after years of exposure would limit the expression of IRG by the infecting parasites and determine the clinical impact of the infection. To address this, we compared IRGs expression in parasites infecting adults and children from the same area and studied the association with their immune status and clinical outcomes.

**Materials and Methods**

### Study area and population.

The study was conducted in the Manhiça District, southern Mozambique\(^35\), where a demographic surveillance system (DSS) set up in 1998 by the Centro de Investigação em Saúde de Manhiça (CISM), currently provides accurate demographic information on its circa 178,000 inhabitants. The region has a warm and rainy season from November to April, and a cooler and drier season the rest of the year. Malaria transmission is perennial with some seasonality. *Plasmodium falciparum* is the predominant species, and *Anopheles funestus* the main vector. Entomological inoculation rate in 2002 was 38 infective bites/person/year\(^35\).

Patients attending the Manhiça District Hospital with a clinical diagnosis of *P. falciparum* malaria were recruited into the study after obtaining written informed consent. Samples were collected between April-November 2006 (children; \(n = 25\) SM patients and \(n = 25\) UM patients) and August 2006 to May 2008 (adults, \(n = 50\), UM). Clinical malaria was defined as the presence of fever (axillary temperature \(\geq 37.5^\circ C\)) with an axillary parasitemia of \(P. falciparum\) \(\geq 500/\mu L\) on thin blood film examination. SM was defined when children had, at least, one of the following symptoms: cerebral malaria (Blantyre Coma Score \(\leq 2\)), severe anemia (packed cell volume \(< 15\%) or hemoglobin \(< 5\) g/dL), acute respiratory distress (lactate \(> 5\) mM and/or chest indrawing or deep breathing), prostration, hypoglycemia (blood glucose \(< 2,2\) mM) and multiple seizures (\(\geq 2\) convulsions in the preceding 24 h)\(^46\). Individuals with UM were those with clinical malaria not showing any of the mentioned signs of severity. Parasite density was determined by optic microscopy. Peripheral blood was collected by venipuncture before treatment. Following centrifugation, the RBC pellet was cryopreserved in liquid nitrogen\(^46\). The studies were approved by the Mozambican National Bioethics Committee and the Hospital Clinic of Barcelona Ethics Review Committee and all experiments were carried out in accordance with relevant guidelines and regulations.

### Parasite culture and quantification of transcript levels.

Cryopreserved *P. falciparum* isolates were thawed and matured as previously described\(^46\). Cultures were monitored by Giemsa-stained thin smears for 18–56 hours until most schizont stages were observed, as determined by the agreement of two trained researchers. Infected RBCs were collected, washed with PBS, mixed with 20 V of TRIzol\(^\text{TM}\) and RNA was obtained following manual extraction with phenol/chloroform phase separation. Total RNA was treated with Turbo DNA-free\(^\text{TM}\) kit (Ambion) at 37 °C for 45 minutes and reverse transcribed with oligo(dT) using Maxima H reagents (Thermo scientific).

Quantitative PCR (qPCR) was performed on an ABI PRISM-7500 real-time system (Applied Biosystems) using 2\(\mu\)L of cDNA in a final volume of 20\(\mu\)L, including 10\(\mu\)L of Power SYBR green master mix (Applied Biosystems) and primers for specific genes at the convenient concentration (see Supplementary Table 1). Gene-specific primers were designed to match conserved regions only in *P. falciparum* based on the coding sequence (GeneBank) and rechecked with sequences available in [Plasmodb.org](http://www.plasmodb.org) on October 2015. Human DNA was not amplified with any primer. To obtain transcript concentration from each sample, the mean Ct (duplicates) of the sample was interpolated to a 7-point 10-log standard curve with 3D7 genomic DNA of known concentrations, for each of the studied genes. Samples with a Ct \(> 30\) for the housekeeping gene (Seryl tRNA synthetase) were discarded, and samples with a Ct higher than the higher Ct from each gene’s standard curve, were considered as not expressed. Each IRG transcript level was normalized as a proportion of the sum of all transcripts concentrations in each isolate and reported as a percentage of expression (relative transcript levels). Expression was considered only if relative transcript level was higher than 0. Multiplicity of infection in children was measured as described elsewhere\(^37\).

### Antibody measurement.

Apical asparagine-rich protein (AARP), *Plasmodium* Thrombospondin-related apical merozoite protein (PTRAMP), EBA175PfF2, PfRh1, PfRh2a/b, PfRh4\(_{30}\), PfRh4\(_{C\text{-terminal}}\), PfRh5, P41 and
Cysteine-rich protective antigen (CyRPA) were produced at the International Centre for Genetic Engineering and Biotechnology (ICGEB)38–42. PfRh22030–2528 and regions III–V of EBA175 (EBA175 III–V) and EBA140 (EBA140 III–V) were produced at the Burnet Institute26, 43, 44. Recombinant proteins were expressed in Escherichia coli. IgG and IgM responses targeting P. falciparum antigens were measured by quantitative suspension array technology (qSAT) in multiplex using xMAP44 beads (Luminex Corporation) coupled to each antigen. Positive, negative and background controls were added to each plate. Multiplexed beads were incubated with plasma samples, and antibody levels were measured as described elsewhere45, 46. Median Fluorescence Intensity (MFI) was obtained from the InVitrogen Luminex® platform (xPONENT® Software, at least 100 counts/analyte) and normalized for inter-plate variability by multiplying individual values by the median value of a positive control assayed in all plates and dividing by each plate’s value. Negative values were substituted by the minimum positive value divided by two in order to log-transform these variables.

**Definitions and statistical analysis.** Age was categorized as < 2.5 years; 2.5 years to ≤ 5; 14 to < 22 years and 22 to ≤ 61 years according to quartile distribution. Seropositive threshold was defined as the mean plus 3 standard deviations of the value of the negative population given by finite mixture models estimated with log-transformed antibody levels. Breadth of antibody responses was defined as the sum, for each individual, of the number of specific antibodies above the seropositive threshold. The ratio for genes from the SA-dependent/SA-independent pathways was obtained from eba175, eba140 and pfrh1 for SA-dependent and from pfrh2a, pfrh2b and pfrh4 for SA-independent pathways. The diversity of gene expression in each P. falciparum isolate was defined as the number of genes with transcripts detected by qPCR.

To compare mean levels or prevalence between groups, we used Student’s t-test (or Wilcoxon rank-sum test if the variable was not normally distributed) and chi-square test, respectively. Spearman's correlations were calculated to assess strength of the relationship between IRG levels and parasite density. Comparisons between more than two groups were done with Kruskal-Wallis test for non-normally distributed data. To assess the association between variables, we performed crude and adjusted linear regression models, including interaction terms with age. Wald’s test was used to assess the overall effect of age on relative transcript levels. Reported p-values were not adjusted for multiple comparisons21, 47 and therefore, the interpretation of results is based on the standard significance level (p < 0.05) and on the magnitude of the effect. Statistical analyses were performed using Stata (StataCorp. 2015. Stata Statistical Software: Release 14. College Station, TX: StataCorp LP) and graphs with Prism7 (GraphPad).

**Results**
The 100 clinical *Plasmodium falciparum* isolates thawed for *ex vivo* culture developed and reached late trophozoite/schizont stages. Three of them (3%) were discarded due to a Ct value > 30 for the housekeeping gene (Seryl tRNA synthetase). All plasma samples from adults and children were tested for specific IgG and IgM levels against the recombinant proteins produced from the studied genes (except for 1 adult and IgM determination in 8 children with UM and 4 children with SM due to unavailability of enough plasma volume). Out of the 96 samples analyzed for transcript and IgG antibody levels, 46 were adults (48%) and 50 children (52%). No statistically significant differences in terms of parasitemia, gender or parasites’ maturation time between groups were found (Table 1).

**Expression of invasion-related genes and specific antibody levels.** The most abundant transcripts in the studied isolates were from *p41*, with median relative transcript levels of 15.2% (interquartile range [IQR] 6.4%, 29.5%), followed by *eba140* (12.6% [5.6%, 22.1%]) and *ptrymp* (7.6% [5.7%, 10.7%]), followed by *cyrpa* (6.1% [4.4%, 10.7%]), *eba175* (5.9% [0.2%, 14.2%]) and *aarp* (4.5% [9%, 12%]). Genes from the *pfrh* family were the least expressed (see Supplementary Table 2). No consistent correlation was found for relative transcript levels of genes involved in the SA-dependent pathway (*eba175*, *eba140*, *pfrh1*) or the SA-independent pathway (*pfrh2a*, *pfrh2b*, *pfrh4*), nor among them (see Supplementary Table 3). No differences were found in the ratio for relative transcript levels reflecting SA-dependent or SA-independent pathways among adults, children with SM and children with UM (Kruskal-Wallis test, p-value = 0.735). Relative transcript levels did not differ in infections with different levels of multiclonality (Kruskal-Wallis test, all p-values > 0.05).

**Table 1.** Demographic, clinical and parasitological factors of the infected individuals included in this study.

|                      | Adults | Children |
|----------------------|--------|----------|
| **N**                | 46     | 50       |
| **Age (years), median (IQR)** | 22 (18, 34) | 2.6 (1.3, 3.5) |
| **Sex (male % (n))**  | 47.8% (22) | 60% (30) |
| **Parasitemia (OM, parasites/μL)** | 47.8% (22) | 60% (30) |
| **Maturation time (hours at 37°C), median (IQR)** | 31.3 (28, 39) | 32 (30, 39) |

- *Wilcoxon rank-sum test; bChi-square test. Abbreviations: UM, Uncomplicated malaria; SM, Severe malaria; IQR, Interquartile range; OM, Optic Microscopy.*
Relative transcript levels of invasion-related genes, by age and specific antibody levels. The median of different transcripts detected within the parasite isolates studied was 10 (9, 12). This median was higher in children (11 [11, 12]) compared to adults (10 [9, 12]) (p = 0.022, Fig. 1a). Diversity in IRG expression was independent of parasite density (Spearman’s rho = 0.07, p = 0.497). Breadth of IgG antibody responses was higher in adults (p < 0.001) and, mirroring the higher diversity in IRG expression in children than in adults, children also showed higher breadth in IgM antibody responses than adults (Wilcoxon rank-sum test, p = 0.003). Median values with the 95% confidence intervals are shown. (c,d) Specific IgG (c) and IgM (d) levels in adults and children. *p < 0.05; **p < 0.01; ***p < 0.001. Grey dots represent values for children and empty dots values for adults. Median values are represented by horizontal black lines.
Relative transcript levels of *eba140* and *ptramp* were lower in older than in younger individuals (*p* = 0.003 and *p* = 0.007, respectively), while *p41* relative transcript levels increased with age (*p* = 0.022) (Fig. 2a). The expression of *eba140*, *ptramp* and *p41* by infecting parasites showed a gradual change as the age of the host increased. In contrast, *pfrh5* relative transcript levels were higher in children than in adults (Wilcoxon rank-sum test, *p* = 0.013), and decreased abruptly during firsts years of life and reached the lowest levels in parasite infecting young adults (Wilcoxon rank-sum test, *p* = 0.023, Fig. 2b). Among children, *p41* was more expressed in parasite isolates from children with SM (*p* = 0.048) than with UM (Fig. 3). This difference was slightly higher when only considering those samples with *p41* transcripts detected (*p* = 0.043). Interestingly, higher levels of relative transcript levels for *eba175* in parasites causing UM compared to SM was only found when considering parasites with *eba175* transcripts detected (*p* = 0.03).

No consistent relationships between relative transcript levels and specific antibodies (IgG and IgM) against the recombinant merozoite antigens were found (Supplementary Table 4). Despite IgG breadth being higher in adults than in children and IgM breadth higher in children than in adults, no general association was found between breadth of IgG or IgM antibody responses and IRG relative transcript levels (Supplementary Figure 1a,b, respectively). Interaction with age was only significant for *ptramp* relative transcript levels and IgG breadth. The association between IgG breadth and *ptramp* relative transcript levels was only significant in children (Coefficient −0.93 [95% confidence interval −0.29, −1.70]; *p* = 0.007). Therefore, data presented in Supplementary Figure 1 was not stratified.

**Transcript levels of invasion-related genes are associated with parasite density.** Parasite density had a proportional increase of 1.3 (95% CI 1.1, 1.5; *p* = 0.007) with a 10% increase in *p41* relative transcript levels. In contrast, parasite density decreased with increasing levels of *eba175* (Coef. 0.7 [95% CI 0.5, 0.9]; *p* = 0.013) and *pfrh5* (Coef. 0.2 [95% CI 0.1, 0.5]; *p* < 0.001), (Fig. 4a). The relationship of *eba175* and *p41* relative transcript levels with parasite density was affected by the fact of being an adult (age groups from 14 to < 22 and 22 to ≤ 61 years old) or children (age groups from 0 to < 2.5 and 2.5 to < 5 years old) (p-values of interaction = 0.021 and 0.005, respectively). After stratifying the analysis, it was observed that in children, but not in adults, parasite density was positively associated with *p41* relative transcript levels (Coef. 1.8 [95% CI 1.3, 2.5]; *p* = 0.001) and negatively associated with *eba175* (Coef. 0.5 [95% CI 0.3, 0.8]; *p* = 0.005) (Fig. 4a). In contrast, the reduction of parasite density with increasing relative transcript levels of *pfrh5* was similar in children (Coef. 0.2 [95% CI 0.03,
bias our results, can be discarded. Thus, it is shown that older individuals have a higher ability to limit the density of IRG transcripts during an infection compared to children. The observed lower parasite densities, contrary to what was previously described: the more exposure to the parasite densities, the more exposure to the parasite. This expression might then be shaped in the first rounds of invasion depending on the host's antimalarial immunity.

The diversity of IRG expression was higher in children than in adults. Such diversity was not associated with parasite density and therefore a sensitivity issue due to cDNA available during the qPCR procedure, which could bias our results, can be discarded. Thus, it is shown that older individuals have a higher ability to limit the diversity of IRG transcripts during an infection compared to children and presumably select the parasites expressing a reduced IRG repertoire. Among the specific IRG analyzed, eba140, ptramp and pfrh5 were the highest in parasites infecting young individuals, while p41 relative transcript levels were the highest in older individuals. This points out how different histories of previous exposure can modulate the expression of genes involved in RBC invasion, probably due to the development of antimalarial immunity upon exposure to the parasite.

The diversity of IRG expression was higher in children than in adults. Such diversity was not associated with parasite density and therefore a sensitivity issue due to cDNA available during the qPCR procedure, which could bias our results, can be discarded. Thus, it is shown that older individuals have a higher ability to limit the diversity of IRG transcripts during an infection compared to children and presumably select the parasites expressing a reduced IRG repertoire. Among the specific IRG analyzed, eba140, ptramp and pfrh5 were the highest in parasites infecting young individuals, while p41 relative transcript levels were the highest in older individuals. This points out how different histories of previous exposure can modulate the expression of genes involved in RBC invasion, probably due to the development of antimalarial immunity upon exposure to the parasite.

The diversity of IRG expression was higher in children than in adults. Such diversity was not associated with parasite density and therefore a sensitivity issue due to cDNA available during the qPCR procedure, which could bias our results, can be discarded. Thus, it is shown that older individuals have a higher ability to limit the diversity of IRG transcripts during an infection compared to children and presumably select the parasites expressing a reduced IRG repertoire. Among the specific IRG analyzed, eba140, ptramp and pfrh5 were the highest in parasites infecting young individuals, while p41 relative transcript levels were the highest in older individuals. This points out how different histories of previous exposure can modulate the expression of genes involved in RBC invasion, probably due to the development of antimalarial immunity upon exposure to the parasite.

The diversity of IRG expression was higher in children than in adults. Such diversity was not associated with parasite density and therefore a sensitivity issue due to cDNA available during the qPCR procedure, which could bias our results, can be discarded. Thus, it is shown that older individuals have a higher ability to limit the diversity of IRG transcripts during an infection compared to children and presumably select the parasites expressing a reduced IRG repertoire. Among the specific IRG analyzed, eba140, ptramp and pfrh5 were the highest in parasites infecting young individuals, while p41 relative transcript levels were the highest in older individuals. This points out how different histories of previous exposure can modulate the expression of genes involved in RBC invasion, probably due to the development of antimalarial immunity upon exposure to the parasite.

The diversity of IRG expression was higher in children than in adults. Such diversity was not associated with parasite density and therefore a sensitivity issue due to cDNA available during the qPCR procedure, which could bias our results, can be discarded. Thus, it is shown that older individuals have a higher ability to limit the diversity of IRG transcripts during an infection compared to children and presumably select the parasites expressing a reduced IRG repertoire. Among the specific IRG analyzed, eba140, ptramp and pfrh5 were the highest in parasites infecting young individuals, while p41 relative transcript levels were the highest in older individuals. This points out how different histories of previous exposure can modulate the expression of genes involved in RBC invasion, probably due to the development of antimalarial immunity upon exposure to the parasite.

The diversity of IRG expression was higher in children than in adults. Such diversity was not associated with parasite density and therefore a sensitivity issue due to cDNA available during the qPCR procedure, which could bias our results, can be discarded. Thus, it is shown that older individuals have a higher ability to limit the diversity of IRG transcripts during an infection compared to children and presumably select the parasites expressing a reduced IRG repertoire. Among the specific IRG analyzed, eba140, ptramp and pfrh5 were the highest in parasites infecting young individuals, while p41 relative transcript levels were the highest in older individuals. This points out how different histories of previous exposure can modulate the expression of genes involved in RBC invasion, probably due to the development of antimalarial immunity upon exposure to the parasite.

The diversity of IRG expression was higher in children than in adults. Such diversity was not associated with parasite density and therefore a sensitivity issue due to cDNA available during the qPCR procedure, which could bias our results, can be discarded. Thus, it is shown that older individuals have a higher ability to limit the diversity of IRG transcripts during an infection compared to children and presumably select the parasites expressing a reduced IRG repertoire. Among the specific IRG analyzed, eba140, ptramp and pfrh5 were the highest in parasites infecting young individuals, while p41 relative transcript levels were the highest in older individuals. This points out how different histories of previous exposure can modulate the expression of genes involved in RBC invasion, probably due to the development of antimalarial immunity upon exposure to the parasite.
affected by anti-parasite host immunity. However, the mechanisms remain still unknown, since p41 is the only gene studied here that has not been directly linked to invasion. More studies will be useful to both determine the role of p41 in the parasite’s biology as well as the effect of specific antibodies in protection against malaria severity.

In the case of eba175, its expression levels were slightly higher in children with UM compared to SM, although it did not reach statistical significance. This is in accordance with previous results showing that invasion by parasites causing UM had the higher dependence on trypsin (and a trend to neuraminidase-sensitive phenotype). Interestingly, isolates from The Gambia showed that SA-dependent invasion is more common among infections in children. Also, although it is not always linked to SA-dependent pathway, it has been hypothesized that eba175 can be more expressed in non-immune individuals. In addition, the acquisition of inhibitory antibodies against antigens from the SA-dependent pathway may be acquired early in life. Here, we show that IgG levels against antigens from the SA-independent pathway (PfRh2 and PfRh4, Fig. 1c) are much higher in adults than in children; early antibody responses (IgM) against PfRh4 are lower in adults; and that expression of the SA-dependent pathway antigen eba140 is lower in adults. Taken together, these evidences suggest the possibility that what is being selected in parasites infecting immune individuals is the invasion phenotype rather than invasion ligands themselves, although further studies are needed.

This study has some limitations that need to be considered. First, we cannot discard that differences in the relative transcript levels between parasites collected from children and adults are due to different duration of the infections, which could affect IRG expression levels. Second, the maturation process of multi-clonal infections could lead to the selection of those clones with higher maturation rates. Given that there is no reference gene to normalize for maturation stage, transcript levels were expressed as relative to the sum of all genes studied to control for temporal transcript patterns during cell cycle development, as has been done in previous studies. Third, there may be a lack of concordance between transcript and protein levels for the merozoite proteins expressed by field isolates. Also, it has not been possible to study the invasion phenotype, which would have helped to clarify its relationship with previous malaria exposure. Finally, the possibility of type I errors derived from analyses reporting non-adjusted p-values should be taken into account, and, although the overall results of our study suggest that expression of genes is modulated by factors modified by age, other studies would be required to confirm this statement.

In conclusion, this study shows that the expression of invasion ligands used by P. falciparum to invade RBCs can be modulated by the age of the infected host, possibly as a surrogate of immunity. These results suggest that the efficacy of malaria vaccines aiming to block the invasion process may differ between individuals with different levels of malaria exposure. Also, the observation that children (or non-immune individuals) are infected by parasites expressing a more diverse IRG repertoire, supports the potential need of combining several RBC binding ligands in a multi-antigenic vaccine to avoid that the parasite escapes the immune responses induced by single-target vaccines. Finally, results reported here suggest that the loss of immunity associated with reductions in malaria transmission after elimination or control campaigns, could lead to a change in the repertoire of IRG expressed by the remaining parasite population, and therefore to changes in the efficacy of invasion-blocking vaccines.

Figure 4. Association of invasion-related genes (IRG) relative transcript levels and parasite density. (a) Effects of 10% increase in relative transcript levels on parasite density by optic microscopy (linear regression models adjusted by age and clinical presentation) are shown with the 95% confidence interval for each specific IRG. Black symbols represent significant associations between IRG relative transcript levels and parasite density. From left to right: all population, only children and only adults. (b) Parasite density by optic microscopy according to tertile categorization of pfRh5 relative transcript levels in the parasite population.
References

1. Loposticki, S. et al. Reticulocyte and erythrocyte binding–like proteins function cooperatively in invasion of human erythrocytes by malaria parasites. Infect. Immun. 79, 1107–1117 (2011).

2. Gaur, D., Mayer, D. C. G. & Miller, L. H. Parasite ligand–host receptor interactions during invasion of erythrocytes by Plasmodium merozoites. International Journal for Parasitology 34, 1413–1429 (2004).

3. Gaur, D. & Chitnis, C. E. Molecular interactions and signaling mechanisms during erythrocyte invasion by malaria parasites. Current Opinion in Microbiology 14, 422–428 (2011).

4. Weiss, G. E., Crabb, B. S. & Gilson, P. R. Overlaying Molecular and Temporal Aspects of Malaria Parasite Invasion. Trends in Parasitology. doi:10.1016/j.pt.2015.12.007 (2016).

5. Wright, G. J. & Rayner, J. C. Plasmodium falciparum Erythrocyte Invasion: Combining Function with Immune Evasion. PLoS Pathog 10, e1003943 (2014).

6. Gaur, D., Chitnis, C. E. & Chauhan, V. S. In Advances in Malaria Research 17, 33–86 (John Wiley & Sons, Inc., 2016).

7. Blair, P. L. et al. Transcripts of developmentally regulated Plasmodium falciparum genes quantified by real-time RT-PCR. Nucleic acids research 30, 2224–2231 (2002).

8. Cortés, A. et al. Epigenetic silencing of Plasmodium falciparum genes linked to erythrocyte invasion. PLoS Pathog 3, e107 (2007).

9. Fang, L. et al. Epigenetic control of the variable expression of a Plasmodium falciparum receptor protein for erythrocyte invasion. Proc. Natl. Acad. Sci. USA. 107, 2224–2229 (2010).

10. Hadley, T. J. et al. Falciparum malaria parasites evade erythrocytes that lack glyco phosphorin A and B (MkMk). Strain differences indicate receptor heterogeneity and two pathways for invasion. The Journal of clinical investigation 80, 1190–1193 (1987).

11. Triglia, T., Duraisingham, M. T., Good, R. T. & Cowman, A. F. Reticulocyte-binding protein homologue 1 is required for sialic acid-dependent invasion into human erythrocytes by Plasmodium falciparum. Molecular Microbiology 55, 162–174 (2005).

12. Stubbs, J. et al. Molecular mechanism for switching of P. falciparum invasion pathways into human erythrocytes. Science (New York, NY) 309, 1384–1387 (2005).

13. Duraisingham, M. T. et al. Phenotypic variation of Plasmodium falciparum merozoite proteins directs receptor targeting for invasion of human erythrocytes. The EMBO Journal 22, 1047–1057 (2003).

14. Taylor, H. M., Grainger, M. & Holder, A. A. Variation in the expression of a Plasmodium falciparum protein family implicated in erythrocyte invasion. Infect. Immun. 70, 5779–5789 (2002).

15. Gaur, D. et al. Upregulation of expression of the reticulocyte homology gene 4 in the Plasmodium falciparum clone D42 is associated with a switch in the erythrocyte invasion pathway. Molecular and Biochemical Parasitology 145, 205–215 (2006).

16. Gaur, D. et al. Recombinant Plasmodium falciparum reticulocyte homology protein 4 binds to erythrocytes and blocks invasion. Proceedings of the National Academy of Sciences 104, 17789–17794 (2007).

17. Sahar, T. et al. Plasmodium falciparum reticulocyte binding–like homologue protein 2 (PRH2) is a key adhesive molecule involved in erythrocyte invasion. PLoS ONE 6, e17102 (2011).

18. Nery, S. et al. Expression of Plasmodium falciparum genes involved in erythrocyte invasion varies among isolates cultured directly from patients. Molecular and Biochemical Parasitology 149, 208–215 (2006).

19. Be, A. K. et al. Variant merozoite protein expression is associated with erythrocyte invasion phenotypes in Plasmodium falciparum isolates from Tanzania. Molecular and Biochemical Parasitology 153, 66–71 (2007).

20. Cowman, A. F. & Crabb, B. S. Involvement of red blood cells by malaria parasites. Cell 124, 755–766 (2006).

21. Richards, J. S. et al. Identification and prioritization of merozoite antigens as targets of protective human immunity to Plasmodium falciparum malaria for vaccine and biomarker development. J. Immunol. 191, 795–809 (2013).

22. Barry, A. E. & Arnott, A. Strategies for designing and monitoring malaria vaccines targeting diverse antigens. Frontiers in immunology 5, 359 (2014).

23. Mayer, D. C. G. et al. Glycophorin B is the erythrocyte receptor of Plasmodium falciparum erythrocyte–binding ligand, EBL-1. Proc. Natl. Acad. Sci. USA. 106, 5348–5352 (2009).

24. Abdi, A. I. et al. Global selection of Plasmodium falciparum virulence antigen expression by host antibodies. Sci Rep. 6, 1–9 (2016).

25. Persson, K. E. M. et al. Variation in use of erythrocyte invasion pathways by Plasmodium falciparum mediates evasion of human inhibitory antibodies. J. Clin. Invest. 118, 342–351 (2008).

26. Persson, K. E. M. et al. Erythrocyte-binding antigens of Plasmodium falciparum are targets of human inhibitory antibodies and function to evade naturally acquired immunity. J. Immunol. 191, 785–794 (2013).

27. Bowyer, P. W. et al. Variation in Plasmodium falciparum erythrocyte invasion phenotypes and merozoite ligand gene expression across different populations in areas of malaria endemicity. Infect. Immun. 83, 2575–2582 (2015).

28. Gomez-Escobar, N. et al. Erythrocyte invasion and merozoite ligand gene expression in severe and mild Plasmodium falciparum malaria. 201, 444–452 (2010).

29. Mensah-Brown, H. E. et al. Analysis of Erythrocyte Invasion Mechanisms of Plasmodium falciparum Clinical Isolates Across 3 Malaria-Endemic Areas in Ghana. J. Infect. Dis. 212, 1288–1297 (2015).

30. Chotivanich, K. et al. Parasite multiplication potential and the severity of Falciparum malaria. Journal of Infectious Diseases 181, 1206–1209 (2000).

31. Simpson, J. A., Silamut, K., Chotivanich, K., Pukrittayakamee, S. & White, N. J. Red cell selectivity in malaria: a study of multiple-infected erythrocytes. Transactions of the Royal Society of Tropical Medicine and Hygiene 93, 165–168.

32. Deans, A.-M. et al. Invasion pathways and malaria severity in Kenyan Plasmodium falciparum clinical isolates. Infect. Immun. 75, 3014–3020 (2007).

33. Deans, A.-M. et al. Low multiplication rates of African Plasmodium falciparum isolates and lack of association of multiplication rate and red blood cell selectivity with malaria virulence. Am. J. Trop. Med. Hyg. 74, 524–563 (2006).

34. Genton, B. et al. A recombinant blood-stage malaria vaccine reduces Plasmodium falciparum density and exerts selective pressure on parasite populations in a phase 1–2b trial in Papua New Guinea. Journal of Infectious Diseases 185, 820–827 (2002).

35. Alonso, P. L. et al. Efficacy of the RTS,S/AS02A vaccine against Plasmodium falciparum infection and disease in young African children: randomised controlled trial. The Lancet 364, 1411–1420 (2004).

36. Mayor, A. et al. Association of severe malaria outcomes with platelet-mediated clumping and adhesion to a novel host receptor. PLoS ONE 6, e19422 (2011).

37. Rovira-Vallbona, E. et al. Low antibodies against Plasmodium falciparum and imbalanced pro-inflammatory cytokines are associated with severe malaria in Mozambican children: a case-control study. Malar J 11, 181 (2012).

38. Pandey, A. K. et al. Identification of a potent combination of key Plasmodium falciparum merozoite antigens that elicit strain-transcending parasite-neutralizing antibodies. Infect. Immun. 81, 441–451 (2013).

39. Pandey, K. C. et al. Bacterially expressed and refolded receptor binding domain of Plasmodium falciparum EBA-175 elicits invasion inhibitory antibodies. Molecular and Biochemical Parasitology 123, 23–33 (2002).

40. Mayor, A. et al. Functional and immunological characterization of a Duffy binding–like alpha domain from Plasmodium falciparum erythrocyte membrane protein 1 that mediates rosetting. Infect. Immun. 77, 3857–3863 (2009).

41. Reddy, K. S. et al. Bacterially expressed full-length recombinant Plasmodium falciparum RH5 protein binds erythrocytes and elicits potent strain-transcending parasite-neutralizing antibodies. Infect. Immun. 82, 152–164 (2014).

42. Reddy, K. S. et al. Multiprotein complex between the GPI-anchored CYPRA with PRH5 and PRhir is crucial for Plasmodium falciparum erythrocyte invasion. Proceedings of the National Academy of Sciences 112, 201415466 (2015).
43. Reiling, L. et al. Evidence that the erythrocyte invasion ligand PlRh2 is a target of protective immunity against Plasmodium falciparum malaria. *J. Immunol.* **185**, 6157–6167 (2010).
44. Ahmed Ismail, H. et al. Acquired antibodies to merozoite antigens in children from Uganda with uncomplicated or severe Plasmodium falciparum malaria. *Clin. Vaccine Immunol.* **20**, 1170–1180 (2013).
45. Moncunill, G. et al. Cytokine and antibody responses to Plasmodium falciparum in naive individuals during a first malaria episode: effect of age and malaria exposure. *PLoS ONE* **8**, e55756 (2013).
46. Campo, J. I. et al. Impact of the RTS.S malaria vaccine candidate on naturally acquired antibody responses to multiple asexual blood stage antigens. *PLoS ONE* **6**, e25779 (2011).
47. Perneger, T. V. What’s wrong with Bonferroni adjustments. *BMJ* **316**, 1236–1238 (1998).
48. Akpogheneta, O. I., Dunyo, S., Pinder, M. & Conway, D. J. Boosting antibody responses to Plasmodium falciparum merozoite antigens in children with highly seasonal exposure to infection. *Parasite Immunol.* **32**, 296–304 (2010).
49. Wilson, D. W. et al. Quantifying the importance of MSP1-19 as a target of growth-inhibitory and protective antibodies against Plasmodium falciparum in humans. *PLoS ONE* **6**, e27705 (2011).
50. Drew, D. R. et al. A novel approach to identifying patterns of human invasion-inhibitory antibodies guides the design of malaria vaccines incorporating polymorphic antigens. *BMC Med.* **14**, 144 (2016).
51. Bachmann, A. et al. Mosquito Passage Dramatically Changes var Gene Expression in Controlled Human Plasmodium falciparum Infections. *PLoS Pathog* **12**, e1005538 (2016).
52. Vola, J. C. et al. Essential Role of the PIRH5/PIRipr/CyRPA Complex during Plasmodium falciparum Invasion of Erythrocytes. *Cell Host & Microbe* **20**, 60–71 (2016).
53. Weaver, R. et al. The association between naturally acquired IgG subclass specific antibodies to the PIRH5 invasion complex and protection from Plasmodium falciparum malaria. *Sci Rep* **6**, 33094 (2016).
54. Doolan, D. L., Dobaño, C. & Baird, J. K. Acquired immunity to malaria. *Clinical Microbiology Reviews* **22**, 13–36, Table of Contents (2009).
55. Leenstra, T. et al. Dehydroepiandrosterone sulfate levels associated with decreased malaria parasite density and increased hemoglobin concentration in pubertal girls from western Kenya. *Journal of Infectious Diseases* **188**, 297–304 (2003).
56. Taechalertpaisarn, T. et al. Biochemical and functional analysis of two Plasmodium falciparum blood-stage 6-cys proteins: P12 and P41. *PLoS ONE* **7**, e41937 (2012).
57. Baum, J., Pinder, M. & Conway, D. J. Erythrocyte invasion phenotypes of Plasmodium falciparum in The Gambia. *Infect. Immun.* **71**, 1856–1863 (2003).
58. Lobo, C.-A. et al. Invasion profiles of Brazilian field isolates of Plasmodium falciparum: phenotypic and genotypic analyses. *Infect. Immun.* **72**, 5886–5891 (2004).

**Acknowledgements**
Authors would like to thank to the individuals participating in the study and their parents/guardians; the staff of the hospital, clinical officers, field supervisors and data manager; Dr. Ariel Magallón-Tejada, Dr. Himanshu Gupta, Dr. Silvie Huijben, Dr. Mercedes Rubio, Dr. Carlota Dobaño and Dr. Alfred Cortés for their useful comments on this project. This work was supported by the Ministerio de Economía y Competitividad in Spain (PRI-PIBIN-2011-0893), the Instituto de Salud Carlos III [grant PI13/01478 2013 call for the Strategic Action on Health 2013–2016 cofunded by the Fondo Europeo de Desarrollo Regional (FEDER), and CES10/021-I3SNS to A.M.]. ISGlobal is a member of the CERCA Programme, Generalitat de Catalunya. The Manhiça Health Research Centre receives core support from the Spanish Agency for International Cooperation and Development. A.V. received a pre-doctoral fellowship from La Caixa foundation. A.M. is supported by the Department d’Universitats I Recerca de la Generalitat de Catalunya (AGAUR; 2014SGR263).

**Author Contributions**
A.V., A.J. and P.C. designed and performed the experiments; A.V. analyzed the data with input from A.C. and A.M.; A.V. wrote the main manuscript text and prepared the figures with the input of A.M. Q.B., P.A., S.M., R.A., B.S., P.A. and A.M.; A.M. provided samples and data from malaria-exposed individuals; V.C., C.L., J.B., C.C. and D.G. provided recombinant proteins. All authors reviewed the manuscript.

**Additional Information**
Supplementary information accompanies this paper at doi:10.1038/s41598-017-05025-5

**Competing Interests:** The authors declare that they have no competing interests.

**Publisher’s note:** Springer Nature remains neutral with regard to jurisdictional claims in published maps and institutional affiliations.

**Open Access** This article is licensed under a Creative Commons Attribution 4.0 International License, which permits use, sharing, adaptation, distribution and reproduction in any medium or format, as long as you give appropriate credit to the original author(s) and the source, provide a link to the Creative Commons license, and indicate if changes were made. The images or other third party material in this article are included in the article’s Creative Commons license, unless indicated otherwise in a credit line to the material. If material is not included in the article’s Creative Commons license and your intended use is not permitted by statutory regulation or exceeds the permitted use, you will need to obtain permission directly from the copyright holder. To view a copy of this license, visit http://creativecommons.org/licenses/by/4.0/.

© The Author(s) 2017
Minerva Access is the Institutional Repository of The University of Melbourne

Author/s:
Valmaseda, A; Bassat, Q; Aide, P; Cistero, P; Jimenez, A; Casellas, A; Machevo, S; Aguilar, R; Sigauque, B; Chauhan, VS; Langer, C; Beeson, J; Chitnis, C; Alonso, PL; Gaur, D; Mayor, A

Title:
Host age and expression of genes involved in red blood cell invasion in Plasmodium falciparum field isolates

Date:
2017-07-05

Citation:
Valmaseda, A., Bassat, Q., Aide, P., Cistero, P., Jimenez, A., Casellas, A., Machevo, S., Aguilar, R., Sigauque, B., Chauhan, V. S., Langer, C., Beeson, J., Chitnis, C., Alonso, P. L., Gaur, D. & Mayor, A. (2017). Host age and expression of genes involved in red blood cell invasion in Plasmodium falciparum field isolates. SCIENTIFIC REPORTS, 7 (1), https://doi.org/10.1038/s41598-017-05025-5.

Persistent Link:
http://hdl.handle.net/11343/256467

File Description:
published version

License:
CC BY