Schistosomiasis Coinfection in Children Influences Acquired Immune Response against *Plasmodium falciparum* Malaria Antigens

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**Abstract**

**Background:** Malaria and schistosomiasis coinfection occurs frequently in tropical countries. This study evaluates the influence of *Schistosoma haematobium* infection on specific antibody responses and cytokine production to recombinant merozoite surface protein-1-19 (MSP1-19) and schizont extract of *Plasmodium falciparum* in malaria-infected children.

**Methodology:** Specific IgG1 to MSP1-19, as well as IgG1 and IgG3 to schizont extract were significantly increased in coinfection compared to *P. falciparum* mono-infected children. Stimulation with MSP1-19 lead to a specific production of both interleukin-10 (IL-10) and interferon-γ (IFN-γ), whereas the stimulation with schizont extract produced an IL-10 response only in thecoinfected group.

**Conclusions:** Our study suggests that schistosomiasis coinfection favours anti-malarial protective antibody responses, which could be associated with the regulation of IL-10 and IFN-γ production and seems to be antigen-dependent. This study demonstrates the importance of infectious status of the population in the evaluation of acquired immunity against malaria and highlights the consequences of a multiple infection environment during clinical trials of anti-malaria vaccine candidates.

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**Introduction**

*Plasmodium falciparum* (*Pf*) malaria remains one of the major public health issues in tropical countries and the vast majority of childhood deaths due to malaria occur in sub-Saharan Africa [1].

Protective immunity to *Pf* malaria is slowly acquired after several infections and is dependent on the intensity and duration of the individual exposure to the parasite [2]. The *P. falciparum* merozoite surface protein 1 (MSP1) and especially its highly conserved C-terminal EGF-like module pair, known as MSP1-19 is one of leading candidate antigens for a vaccine against the malaria parasite blood stage [3,4]. In infected humans, humoral immune responses to blood stage parasites play a primary role in providing protection against malaria [5] and they are largely dependent on cytophilic type immunoglobulin (Ig) antibodies (Abs) such as IgG1 and IgG3 isotypes [4]. In addition, a specific cellular immune response and its associated cytokines, play a key protective or pathological role during malaria. Some cytokines, such as interferon-γ (IFN-γ) and Interleukin-10 (IL-10), are known to be directly involved in the production of specific isotypes of anti-*P. falciparum* Ab responses [6]. Hartgers et al. recently demonstrated that in Ghanaian school children, there was an increase in specific IL-10 production in helminth-infected individuals, compared to non-helminth infected [7]. Therefore, the regulation of the specific production of cytokines induced by such parasite infection could have a substantial impact on the development of malaria protection, as well as on its pathological consequences [7–8].

Intrinsic features of the immune system that develop with age but also with the presence of other chronic infections may influence malaria immunity [9–11]. Indeed, a chronic coinfection, such as schistosomiasis, may have a key impact on the acquired immune response to *Plasmodium* infection [9]. Previous studies, relating the complexity of interactions between host response to helminths and malaria infection, suggested possible consequences...
on age-dependent malaria morbidity [12–14]. In addition, the presence of *Schistosoma* coinfection during uncomplicated *P. falciparum* malaria unbalances the regulation of the associated inflammatory response [15]. Coinfection with helminthic parasites could then constitute a confusing factor in the assessment of efficacy of malaria-control intervention, including vaccine clinical trials [7,12].

The present study evaluates the impact of coinfection by *S. haematobium* (SH) on the specific isotype Ab response and its associated cytokine production to PIMSP1-19 and to schizont extracts of *P. falciparum* asexual blood stage antigens (AgS) in children living in a particular malaria endemic area, where schistosomiasis appeared 15 years ago [15].

**Materials and Methods**

**Subjects**

The studied population was a cohort of 79 malaria infected children living in the same area of the Senegal River basin (villages of Lampsar, Taba Tache and Taba Dar Salam), as previously described [16]. Two groups were identified in this cohort: children infected by *P. falciparum* without a confirmed schistosomiasis (PF) (n = 40; age mean = 11 years; range: 7–13; villages of Taba Tache and Taba Dar Salam) and malaria infected children presenting a coinfection with *S. haematobium* (PF-SH) (n = 39; age mean = 10 years; range: 7–13; villages of Lampsar). Comparable mean age and sex ratio were obtained. Absence of clinical morbidity for malaria and schistosomiasis was a criterion of selection. The detection of schistosomiasis infection was performed using the referent parasite criteria (presence of eggs in urine or/and in feces). Schistosoma infected patients presenting pathological cases of schistosomiasis were treated but not included in the study. The Taba Tache and Taba Dar Salam villages are located down-river from Diama dam on the Senegal river where schistosomiasis is absent as previously described [13–14] and as confirmed by the present study.

**Diagnosis**

**PF infection.** *PF* infection was detected by Quantitative Buffy Coat (QBC) (Becton Dickinson) and parasites were then counted and identified on blood smears. These tests are commonly used in malaria endemic areas and blood smears represent the referent criterion to detect malaria infection [17]. Only red blood cells infected by *PF* were observed on positive slides. No other *Plasmodium* species infection was detected. The studied population was considered positive for malaria when *PF* was detected with the QBC test and confirmed by blood smear observation, over a one-month period. In coinfection children, the mean parasitaemia of *P. falciparum* was not significantly different from the respective *PF* mono-infected groups (Mann-Whitney U-test). Mean of parasitaemia was 1617 per mm³/blood for *PF* infected children versus 340 per mm³/blood for coinfectected children. During pre-selection, subjects showing high positivity in QBC (+ + +) were excluded and immediately treated for malaria. Children presenting clinical symptoms of mild or severe malaria morbidity were not selected. The studied population did not therefore present clinical symptoms of malaria morbidity. None of the selected children had received anti-malaria treatment in the month preceding the study.

**Schistosoma haematobium infection.** The presence of SH eggs was evaluated in three samples of urine (urine filtration) using microscopy. In the studied area and population, no *Schistosoma mansoni* (Sm) infection was detected using Kato-Katz method.

In the villages of Taba Tache and Taba Dar Salam, schistosomiasis infection has never been previously detected [16; D. De Clercq, unpublished data]. In addition to the complete absence of schistosome eggs in faeces and urine, we have confirmed the absence of *Schistosoma* infection in these villages by evaluating the schistosome worm circulating anodic antigen (CAA) in sera, as previously described [16].

In Lampsar village, the presence of Sm and Sh eggs was evaluated in three samples of faeces (Kato–Katz method) or urine (urine filtration), respectively. The intensity mean of urinary Sh was 57 eggs/10 ml of urine.

None of the selected children had received anti-schistosome treatment in the previous 6 months.

The study followed ethical principles and was approved by the ethics committee of the Senegalese Ministry of Health. Written informed consent was obtained from the study population by their parents or guardians.

**P. falciparum and schistosome antigens**

The *P. falciparum* antigens were recombinant PIMSP1-19 and schizont total antigens. PIMSP1-19 was produced in *Pichia pastoris* as a hexa-His fusion [18]. PIMSP1-19 was provided by J. McCormick and W. Morgan from National Institute for Medical Research (London, UK). The schizont antigenic preparation of *P. falciparum* is a soluble extract of schizont lysate obtained from infected erythrocyte cultures [19].

Soluble Egg Antigen (SEA) is a total soluble extract of schistosome eggs obtained from experimental infection in guinea pigs by the Antigen Production Laboratory at the Pasteur Institute of Lille, France.

**Human Ab levels**

Specific antibody levels to PIMSP1-19, schizont extract or SEA in sera were determined by enzyme-linked immunosorbent assay (ELISA).

PIMSP1-19 protein (0.5 μg/ml), schizont extract (0.5 μg/ml) or SEA (5 μg/ml) were coated on 96-well plates (Nunc, Denmark) for 2 h 30 at 37°C. Plates were then blocked in PBS containing 0.5% gelatin (Merck, Germany). The sera diluted in PBS/Tween were incubated at 4°C overnight at a 1/100 dilution for IgG2, IgG3 and IgG4, 1/50 for IgA and IgE, and at 1/1,000 for IgG1 detection to schizont extract. For MSP1-19 antigen, a 1/100 dilution for IgG2, IgG3, IgG4 and IgA, 1/50 for IgE and 1/1,500 for IgG1 were used and for SEA, a 1/1,000 dilution for IgG1 and 1/100 for IgG3. Corresponding biotinylated mAbs to human Ig isotypes (SBA, AL or BD Pharmingen, CA, USA), were then incubated at a 1/1,000 dilution for IgG2, IgG4; 1/2,000 for IgG1, 1/750 for IgG3; 1/1,500 for IgA and 1/250 for IgE (1 h 30 min at 37°C). Streptavidin-horseradish peroxidase-conjugated for IgG1, IgG2, IgG3 and IgG4 and IgA detection (1/1,500, 30 min at 37°C; Amersham, Les Ulis, France) and ExtrAvidin® peroxidase conjugate for IgE detection (1/5,000, 30 min at 37°C; Sigma, St-Louis, MO) were then added. These ELISA conditions have been determined according to the results of previous experiences. Colorimetric development was allowed by means of ABTS (2,2′-azino-bis(3-ethylbenzothiazoline 6-sulphonic acid) diammonium; Sigma) in 50 mM citrate buffer (pH = 4 containing 0.003% H₂O₂), and absorbance (OD) was measured at 405 nm for IgG1, IgG2, IgG3, IgG4 and IgA. IgE detection was revealed by OPD (1,2 - Phenylene Diamine Dihydrochloride, Dako; Glostrup, Denmark). OD was added for 30 minutes, followed by the addition of 2N H₂SO₄ to stop the reaction and OD was measured at 492 nm. The same ELISA procedures were performed in parallel for 30 sera from uninfected European subjects used as negative controls. Individual results were
expressed as ΔOD value calculated for each isotype test according to the formula: ΔOD = ODx-ODn, where ODx is the individual OD value of infected patient and ODn is the arithmetic mean of individual OD value for the 30 uninfected control individuals (ODn value for IgE = 0.107; IgG1 = 0.225; IgG2 = 0.123; IgG3 = 0.124; IgG4 = 0.111, IgA = 0.159 using schizont extract; IgE = 0.09, IgG1 = 0.156, IgG2 = 0.121, IgG3 = 0.128, IgG4 = 0.120, IgA = 0.181 using MSP1-19; IgG1 = 0.151, IgG3 = 0.147 using SEA).

**Adsorption procedure of sera with schistosome antigens**

Individual sera of *Pf/Sh* coinfected subjects were diluted with PBS/Tween solution and incubated with schistosomes Soluble Egg Antigens (SEA) at a final concentration of 200 μg/ml. The mixture (sera-SEA) was incubated with gentle shaking for 5 h at room temperature to allow the binding of specific Abs to SEA. Specific IgG1 and IgG3 responses to SEA, to schizont extract of *Pf* or to PMSP1-19 were evaluated by ELISA (as described above) before and after this adsorption procedure.

**Whole blood culture and cytokine production measurement**

Sterile conditions were maintained during the whole blood culture procedure, as originally described [20]. After white blood cell count, blood samples were diluted in RPMI-1640 medium (Gibco, France) supplemented with 2 mM L-glutamate, 0 μg/ml Gentamicin and 1 mM Sodium Pyruvate in order to obtain 2 × 10^6 mononuclear cells/ml. Blood cells were then distributed at the rate of 1 ml/well in 24-well plates (Nunc). Antigens were added to each culture well to a final concentration of 1 μg/ml for MSP1-19 and 10 μg/ml for schizont extract. After 3 days of incubation at 37°C in 5% CO2, the content of each culture well was centrifuged, and supernatants were harvested and conserved at −80°C. Supernatant concentrations of IL-10, IL-12, IL-13, IFN-γ and TGF-β were measured at day 3 by means of ELISA kits according to the manufacturer’s instructions (Immunotech, France). Specific cytokine production after Ag stimulation were expressed as picograms, nanograms or International Units (IU) per milliliter after the subtraction of the amount detected in medium control cultures (non-stimulated conditions) for each individual. Identical culture conditions were assessed with mononuclear cells of 30 uninfected control individuals in which no cytokine production was detected after Ag stimulation (data not shown).

**Statistical analysis**

All data were analysed with Graph Pad Prism (Graph Pad Software, CA, USA). After verifying that values did not assume a Gaussian distribution, the nonparametric Mann-Whitney U-test was used to compare values between both independent groups. Correlation between specific Ab levels and the intensity of *P. falciparum* infection was analysed using Spearman’s rank correlation coefficient. All differences were considered significant at *p* < 0.05.

**Results**

**Specific Ab responses to PMSP1-19 and schizont extract**

The comparison of specific isotype Ab response to MSP1-19 between children having been solely infected by *P. falciparum* (*Pf*) and those coinfected with *S. haematobium* (*Pf/Sh*) was assessed (Figure 1). IgG1 Ab levels were significantly higher in *Pf/Sh*-infected children, compared to *Pf*-infected children (*P* < 0.05) (Figure 1A). The specific IgG3 response was detected in a few *Pf/Sh*-infected children but no statistical difference was observed between both groups (Figure 1A). No relevant anti-MSP1-19 IgG2, IgG4 and IgA responses were observed in either group and the levels of specific IgE Ab were similar between *Pf* and *Pf/Sh*-infected children (Figure 1A and 1B).

To evaluate the antigenic restriction of the observed isotype variations according to infectious status, Ab responses against whole schizont extract were also assessed. As observed with the MSP1-19 Ag, specific IgG2, IgG4, IgA, and IgE responses to total schizont extract were very low or similar between *Pf* and *Pf/Sh* groups (Figure 2A and 2B). In contrast, specific IgG1 and IgG3 Ab levels were significantly higher in *Pf/Sh*-infected children compared to *Pf*-infected group (Figure 2A). These results suggest that the observed difference in IgG1 responses is not restricted to MSP1-19 antigen but is also found with *P. falciparum* total extract.

Figure 1. Specific Ab responses to PfMSP1-19 in children infected by *P. falciparum* (*Pf* - *n* = 40) and coinfected with *S. haematobium* (*Pf-Sh* - *n* = 39) as determined by ELISA. Results represent individual values of OD and bars are the arithmetic mean for each group. Statistical significance between each group is indicated. (A) isotypes IgG1, IgG2, IgG3 and IgG4, (B) isotypes IgA and IgE.

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IgG3 observed in haematobium (comparison shows that the mean OD values were similar between doi:10.1371/journal.pone.0012764.g002 isotypes IgG1, IgG2, IgG3 and IgG4, (B) isotypes IgA and IgE. Each group. Statistical significance between each group is indicated. (A) represent individual values of Absence of cross-reactivity of anti-Plasmodium falciparum (Pf) and Schistosoma haematobium (Pf/Sh) coinfection.

Figure 2. Specific Ab responses to schizont extract in children infected by Plasmodium falciparum (Pf - n = 40) and coinfected with S. haematobium (Pf-Sh - n = 39) as determined by ELISA. Results represent individual values of ΔOD and bars are the arithmetic mean for each group. Statistical significance between each group is indicated. (A) isotypes IgG1, IgG2, IgG3 and IgG4, (B) isotypes IgA and IgE. doi:10.1371/journal.pone.0012764.g002

Absence of cross-reactivity of anti-Plasmodium falciparum IgG1 and IgG3 from coinfected patients with schistosome antigens

We cannot exclude that antigenic cross-reactions can at least partly account for increased levels of anti-Plasmodium falciparum IgG1 and IgG3 observed in Pf/Sh coinfected group. Adsorption experiments with total schistosome antigens (SEA) of sera from coinfected group were therefore performed in order to verify the possible presence of IgG1 and IgG3 Abs cross-reacting with schistosome and Plasmodium antigens.

First, results show a complete adsorption of anti-SEA IgG1 and IgG3 Abs after incubation of the Pf/Sh patient sera with SEA, indicating the efficiency of the experimental procedure (Table 1). Using the same individual samples, we have evaluated the specific IgG1 and IgG3 Ab levels against both Plasmodium falciparum Ags (schizont extract and MSP1-19) before and after Abs adsorption. The comparison shows that the mean OD values were similar between SEA adsorbed and non adsorbed sera for both schizont extract and MSP1-19 Ags (Table 1). These data show that increased levels of IgG1 and IgG3 isotypes recognizing malaria antigens in the coinfected group could not be explained by a cross reactivity to Ags from both parasites.

Influence of intensity of Plasmodium falciparum infection on the variation of anti-Plasmodium falciparum IgG1 and IgG3 levels observed between groups of children

The correlation between specific Ab levels and the parasitaemia due to Plasmodium falciparum infection was first analysed using Spearman’s rank correlation coefficient, without taking into account the coinfection status. This analysis did not reveal any significant correlation (data not shown). This suggests that the observed quantitative difference in the anti-Plasmodium falciparum Abs responses between both groups were not due to differences in the intensity of blood malaria parasites but could be related to the influence of Schistosoma coinfection.

In addition, we have compared individual IgG1 and IgG3 responses against schizont Ags from both groups (coinfected ss. mono-infected) matched on the level of Pf parasitaemia. We have obtained 25 matched-pairs with parasitaemia values varying from 3 to 3780 infected erythrocytes per mm$^3$ of blood. The Wilcoxon t-test was used to compare the means of the matched-pairs. Regarding Ab responses against schizont Ags, the obtained value was t=3.75 for IgG1 (p<0.001) and t=3.23 for IgG3 (p<0.001), indicating that both groups were significantly different regarding their IgG1 and IgG3 levels when patients were matched on their parasitaemia.

Taken together these results indicate that the level of Plasmodium falciparum parasitaemia should not be considered as a confounding factor implicated in the observed variation of specific IgG1 and IgG3 responses according to infectious status.

Production of cytokines in response to PfMSP1-19 and schizont extract

The production of specific IFN-γ, IL-10, IL-12, IL-13 and TGF-β cytokines after in vitro stimulation of mononuclear cells from studied individuals by schizont or MSP1-19, was evaluated in children (Table 2). These cytokines have been selected according to previous studies showing their role in the immune response regulation during malaria infection and for some of them, their role in the regulation of Ab isotype production [7]. The specific production of IL-12, IL-13 and TGF-β to both Plasmodium falciparum antigens was not significantly different between Pf and Pf/Sh groups. In contrast, specific IL-10 and IFN-γ production after MSP1-19 or schizont extract stimulation, showed differences between both groups. IL-10 production in presence of both Ags was significantly higher in Pf/Sh-infected children than in Pf-infected children (p = 0.025, p = 0.002, respectively). As for specific IFN-γ secretion, difference between both groups was only observed for the schizont extract and was significantly higher in Pf/Sh-group compared to Pf-group (p = 0.02). Nevertheless, high IFN-γ production to MSP1-19 was observed in few coinfected children (12/39) but the mean value of IFN-γ secretion in this group was not statistically higher than the one observed in Pf-group (p = 0.282). Background production of IFN-γ and IL-10 (medium alone) was similar between both groups (IFN-γ: 1.25 [mean] ±2.38 [SEM] UI/ml in Pf group, and 4.95±10.71 UI/ml in Pf/Sh group; NS) (IL-10: 248.1±219 ng/ml in Pf group, and 185.2±140.3 ng/ml in Pf/Sh group; NS). These last results indicate that the differences of cytokine production according to infection status were observed only after the stimulation by Plasmodium antigens.
responses to schizont extract were higher in Pf/Sh potential antigenic cross-reactivity. Indeed, the IgG1 and IgG3 increasing IgG1 and IgG3 Abs levels. This increase was not demonstrated that S. haematobium [21,22]. It is believed that these cytophilic IgG subclasses can IgG1 and IgG3 are thought to play a key role in protection increased in coinfected children suggesting that the influence of children compared to the Pf infected children have never been infected by Schistosoma. We demonstrated that S. haematobium coinfection influenced the humoral immune response against malaria antigens by specifically increasing IgG1 and IgG3 Abs levels. This increase was not dependent on the age, on the intensity of malaria infection or on a potential antigenic cross-reactivity. Indeed, the IgG1 and IgG3 responses to schizont extract were higher in Pf/Sh-infected children compared to the Pf-infected group. However, only specific IgG1 Ab response to the MSP1-19 antigen was significantly increased in coinfected children suggesting that the influence of coinfection could be antigen-dependent. During human malaria, IgG1 and IgG3 are thought to play a key role in protection [21,22]. It is believed that these cytophilic IgG subclasses can neutralize parasites directly, by inhibiting parasite invasion or growth in erythrocytes, or indirectly by a mechanism involving cooperation between parasite-opsonising antibodies and monocytes, through binding to the Tcc receptor IIA [23–25]. The subclass response to MSP1-19 was biased towards IgG1 with a minor component from the IgG3 subclass [26]. In addition, the presence of specific IgG1 instead of IgG3 was associated with clinical protection against malaria infection [27,28].

In the present study, we could also rule out the possible Abs cross reactivity between total Sh and Pf antigens, by assessing SEA adsorption experiments. In addition, statistical analysis showed the absence of an effect of the intensity of P. falciparum parasitaemia on these specific Ab responses to malaria antigens. Altogether, the data suggest that chronic schistosome infection in malaria-infected children could affect the specific immune response to P. falciparum. In particular, S. haematobium coinfection appeared to influence the specific isotype Ab responses involved in the development of protective immunity to P. falciparum infection. Moreover, we and others have described that P. falciparum coinfection also influenced the anti-schistosome immune response in S. haematobium-infected children [29,30]. In this case, anti-schistosome IgG3 level was increased in coinfected individuals in contrast to the other IgG subclasses. In regard to our present results, the co-occurrence of schistosomiasis and malaria in individuals could have a particular effect on the IgG3 isotype regulation. The specific IgG3 response to MSP1-19 has been observed in 1/3 of coinfected children (12/39) whereas this isotype response was not detected in children infected only by Pf. Several immuno-epidemiological studies indicate that the presence of anti-MSP1-19 IgG3 response could be dependent to the studied site. For example, a very low IgG3

Table 1. Absence of cross reactivity of anti-P. falciparum IgG1 and IgG3 from coinfected patients with Schistosoma antigens.

| Schistosome Ag | P. falciparum Ags |     |     |     |     |
|---------------|------------------|-----|-----|-----|-----|
|               | SEA              | SE  | PMSP1-19 |     |     |
|               | NA   | A    | N A   | A    | A  |
| IgG1*         | 1.024±0.222     | 0.019±0.09* | 0.336±0.063 | 0.323±0.061 | 1.885±0.319 | 1.762±0.311 |
| IgG3          | 0.413±0.115     | 0.013±0.004* | 0.259±0.065 | 0.342±0.079 | 0.683±0.257 | 0.704±0.250 |

*: Results are mean ± SEM of individual specific production (after subtracting the amount detected in unstimulated medium control cultures for each individual).

Discussion

Following the study of the influence of schistosomiasis coinfection on the regulation of inflammatory factors in uncomplicated P. falciparum malaria [16], we have here evaluated the influence of coinfection in children on the acquired specific immune response to total Plasmodium Ags extract (schizont stage) and to the specific PMSP1-19 vaccine candidate. The greatest interest of this study is that the selected children lived in the same low malaria area (confirmed by the similar parasitemia between groups), without clinical manifestations, but living in villages from the same studied area, where schistosomiasis was monitored since several years (13–14). In particular, it indicated that Pf mono-infected children have never been infected by Schistosoma. We demonstrated that S. haematobium coinfection influenced the humoral immune response against malaria antigens by specifically increasing IgG1 and IgG3 Abs levels. This increase was not dependent on the age, on the intensity of malaria infection or on a potential antigenic cross-reactivity. Indeed, the IgG1 and IgG3 responses to schizont extract were higher in Pf/Sh-infected children compared to the Pf-infected group. However, only specific IgG1 Ab response to the MSP1-19 antigen was significantly increased in coinfected children suggesting that the influence of coinfection could be antigen-dependent. During human malaria, IgG1 and IgG3 are thought to play a key role in protection [21,22]. It is believed that these cytophilic IgG subclasses can neutralize parasites directly, by inhibiting parasite invasion or growth in erythrocytes, or indirectly by a mechanism involving cooperation between parasite-opsonising antibodies and monocytes, through binding to the Tcc receptor IIA [23–25]. The subclass response to MSP1-19 was biased towards IgG1 with a minor component from the IgG3 subclass [26]. In addition, the presence of specific IgG1 instead of IgG3 was associated with clinical protection against malaria infection [27,28].

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Table 2. Specific cytokine production to PMSP1-19 and Schizont extract in children infected by P. falciparum (Pf, n = 40) or coinfected by P. falciparum and S. haematobium (Pf-Sh, n = 39).

| PMSP1-19 | Schizont Extract |     |     |     |     |
|----------|------------------|-----|-----|-----|-----|
|          | Pf              | Pf/ Sh | P value* |   |    |
| IL-10* (ng/ml) | 14.73±5.21 | 22.42±4.89 | 0.025 | 36.64±10.99 | 72.99±10.79 | 0.002 |
| IL-12 (pg/ml)  | 17.24±5.63 | 16.82±6.06 | 0.207 | 20.47±7.47 | 26.23±8.72 | 0.386 |
| IL-13 (pg/ml)  | 2.34±1.33 | 3.07±1.09 | 0.219 | 4.22±1.69 | 4.10±2.08 | 0.495 |
| IFN-γ (pg/ml)  | 0.26±0.11 | 4.22±2.05 | 0.282 | 1.72±0.48 | 11.3±5.34 | 0.020 |
| TGF-β (pg/ml)  | 6.72±1.83 | 5.23±1.89 | 0.286 | 12.65±2.31 | 10.81±1.60 | 0.259 |

*: Results are mean ± SEM of individual specific production (after subtracting the amount detected in unstimulated medium control cultures for each individual).

P value between Pf and Pf/Sh groups (nonparametric Mann-Whitney U test). Differences are considered significant for P<0.05.

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antibody response to MSP1-19 antigen was observed in some malaria endemic areas [26,28,31]. This difference might be due to different genetic background of these populations and/or to a different intensity of transmission in the studied areas or to environmental factors. Indeed, it was speculated that the differential evolution of plasma IgG1 and IgG3 antibodies resulted from a different catabolism and/or use of these antibodies [32]. Thus, history of malaria exposure, seasonal transmission and the short life of specific IgG3 to MSP1-19 antigen cannot be excluded to explain the presence or the absence of this specific IgG3 response. Nevertheless, our results indicate that the presence of specific IgG3 to MSP1-19 among some children living in an endemic malaria area was observed when schistosomiasis coinfection has been established. It could be of great interest to confirm the observations of our study in other coinfection contexts. Our study indicates that it will be crucial to systematically consider the coinfection status of patients in studies concerning the anti-malaria immunity. It strengthens the necessity to screen patients for helminth infections and their immune influence before and during malaria vaccine trials, as previously suggested [9].

We have demonstrated that the influence of *S. haematobium* infection on specific immune response to *Plasmodium* Ags involves specific cytokine production. Helminth infections seemed to influence IL-12, IL-13 and TGF-β production under stimulation with both malaria Ags. In contrast, specific IFN-γ and IL-10 production to the schizont extract were significantly higher in the coinfected group, which showed also a predominant anti-schizont IgG1 and IgG3 Ab levels. In addition, specific IL-10 production to MSP1-19 was significantly higher in the coinfected group compared to children only infected by *Pf* as observed for specific IgG1 response. In *vitro* human studies prove that stimulation by MSP1-19 preferentially elicits IL-10 production [33]. Moreover, IL-10 production specific to schistosome antigens was very relevant in children infected by schistosomes [34]. This cytokine is known to favour the production of IgG1 and IgG3 isotypes, especially during malaria infection [33,35]. In addition, specific IgG3 response has been closely associated with the production of IFN-γ during human infectious diseases, such as *Borreia burgdorferi* [36] or *S. haematobium* chronic infections [F. Remoue, unpublished data]. In the acute uncomplicated malaria, it has been demonstrated that peripheral blood CD4+ T cells producing both IFN-γ and IL-10 were significantly increased during drug-induced clearance of parasitaemia [37]. Moreover, it has been shown that T cells from vaccinated healthy volunteers and adults naturally exposed to malaria with MSP1-19 could produce both IFN-γ and IL-10 [38].

In our study, no statistical correlation of specific IFN-γ or IL-10 productions with IgG1 and/or IgG3 production has been observed. Nevertheless, we have shown an association between the production of these cytokines and the regulation of these specific isotypes according to the group of infection. With regard to previous studies [33–37] and the present results, we can suggest that IFN-γ, acting in synergy with IL-10, could be associated to events leading to the predominant IgG3 production as previously observed during others parasitic infections [F. Remoue, unpublished data]. In addition, it is likely that the production of anti-*P. falciparum* IgG1 and IgG3 Abs could be differently regulated by IL-10 and IL-10/IFN-γ, respectively. This hypothesis is now under investigation by analysing the *in vitro* isotype production after cytokine stimulation of T cells from infected individuals.

The influence of environment, genetic background or nutritional status cannot be ruled out to explain the variation of specific immunity observed in our study. Nevertheless, the concomitant infections such as schistosomiasis in children may influence the acquired anti-malaria Ab response and associated cytokine production. The presence of chronic schistosomiasis infection could specifically increase IL-10 and IFN-γ production in coinfected children, a cytokine production which would positively influence the production of cytophilic anti-malaria IgG1 and IgG3 isotypes, known to be protective during malaria infection. Altogether, this study represents an approach of the regulation of anti-malaria blood stage immunity by coinfection and indicates that the helminthic infection status has to be taken into account in the malaria epidemiological and vaccine studies.

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**Author Contributions**

Conceived and designed the experiments: TOD FR NC DDC JPD OG GR. Contributed reagents/materials/analysis tools: TOD FR SP. Performed the experiments: TOD FR LG AMS OG GR. Conceived and designed the experiments: TOD FR NC DDC JPD OG. Performed the experiments: TOD FR SP. Analyzed the data: CONCEIVED AND DESIGNED THE EXPERIMENTS: TOD FR NC DDC JPD OG GR. Contributed reagents/materials/analysis tools: TOD GR SP. Performed the experiments: TOD FR LG AMS OG GR. Performed the experiments: TOD FR NC DDC JPD OG GR. Contributed reagents/materials/analysis tools: TOD FR LG AMS OG GR. Performed the experiments: TOD FR NC DDC JPD OG GR. Contributed reagents/materials/analysis tools: TOD GR SP. Performed the experiments: TOD FR LG AMS OG GR. Performed the experiments: TOD FR NC DDC JPD OG GR. Contributed reagents/materials/analysis tools: TOD GR SP. Performed the experiments: TOD FR LG AMS OG GR. Performed the experiments: TOD FR NC DDC JPD OG GR. Contributed reagents/materials/analysis tools: TOD GR SP. Performed the experiments: TOD FR LG AMS OG GR.

**References**

1. WHO (2009) Global malaria and elimination: report of a technical review. Geneva: World Health Organization.
2. Mbar K, Kanyanjui S (2006) Immune effector mechanisms in malaria. Parasite Immunol 28(1-2): 51–60.
3. Langhorne J, Ndungu FM, Sponaas AM, Marsh K (2008) Immunity to malaria: More questions than answers. Nat Immunol 9: 725–733.
4. Conway D J, Cavanagh DR, Tanabe K, Roper C, Mikes ZS, et al. (2000) A principal target of human immunity to malaria identified by molecular population genetic and immunological analyses. Nat Med 6(6): 689–92.
5. Yazdani SS, Mulkerrin P, Chauhan VS, Chitnis CE (2006) Immune Responses to Apical Blood-Stage of Malarial Parasites. Current Molecular Medicine 6: 187–203.
6. Garraud O, Mahanty S, Perraut R (2003) Malaria-specific antibody subclasses in immune individuals: a key source of information for vaccine design. Trends Immunol 24(1): 30–39.
7. Hargers FC, Obeng BB, Bokey D, Yazdanbakhsh M (2008) Immune responses during helminth-malaria co-infection: a pilot study in Ghanaian school children. Parasitology 135(7): 853–60.
8. Taylor-Robinson AW (1998) Immunoregulation of malarial infection: balancing the vices and virtues. Int J Parasitol 28(1): 11–20.
9. Nacher M (2001) Malaria vaccine trials in a wormy world. Trends Parasitol 17(2): 563–5.
10. Su Z, Segura M, Morgan K, Loresco-Oui JC, Stevenson MM (2005) Impairment of protective immunity to blood-stage malaria by concurrent nematode infection. Infect Immun 73(6): 3531–9.
11. Van Geertruyden JP, Van Ejik E, Yosaatmadja F, Kasongo W, Mulema G, et al. (2009) The relationship of Plasmodium falciparum hemeral immunity with HIV-1 immunosuppression and treatment efficacy in Zambia. Malar J 18; 8: 250.
12. Dowlo P, Tall A, Sokhna C (2003) Worms can worsen malaria: towards a new means to roll back malaria? Trends Parasitol 21(8): 359–62.
13. Sokhna C, Le Hesran JY, Mbaya PA, Akiana J, Camara P (2004) Increase of malaria attacks among children presenting concomitant infection by *Schistosoma mansoni* in Senegal. Malar J 3: 43.
14. Briand V, Watier L, Le Hesran JY, Garcia A, Cot M (2005) Coinfection with Plasmodium falciparum and Schistosoma haematobium: protective effect of schistosomiasis on malaria in senegalese children? Am J Trop Med Hyg 72(6): 702–7.
15. Picquet M, Ernould JC, Vercruyse J, Southgate VR, Mbaye A, et al. (1996) Royal Society of Tropical Medicine and Hygiene meeting at Manson House, London, 18 May 1995. The epidemiology of human schistosomiasis in the Senegal river basin. Trans R Soc Trop Med Hyg 90(4): 580–6.

16. Diallo TO, Remoué F, Schacht AM, Charrier N, Donnouer JP, et al. (2004) Schistosomiasis co-infection in humans influences inflammatory markers in uncomplicated Plasmodium falciparum malaria. Parasite Immunol 26(8-9): 365–9.

17. Paria SC, Bhadrapkar R, Elangoov S, Chaya DR (2009) A comparative study of blood smear, QBC and antigen detection for diagnosis of malaria Indian J Pathol Microbiol 52(2): 200–2.

18. Morgan WD, Birdshall B, Freerkeil TA, Gradwell MG, Burghaus PA (1999) Solution structure of an EGF module pair from the Plasmodium falciparum merozoite surface protein 1. J Mol Biol 291(1): 113–22.

19. Trager W, Jensen JB (1976) Human malaria parasites in continuous culture. Science 193(4248): 673–5.

20. Joseph S, Jones EM, Kimani G, Maitoka JK, Kamau T, et al. (2004) Cytokine production in whole blood cultures from a fishing community in an area of high endemicity for Schistosoma mansoni in Uganda: the differential effect of parasite worm and egg antigens. Infect Immun 72(2): 728–34.

21. Aribot G, Roger C, Sarthou JL, Trape JF, Balde AT, et al. (1996) Pattern of immunoglobulin isotype response to Plasmodium falciparum blood-stage antigens in individuals living in a holoendemic area of Senegal (Diehmo, west Africa). Am J Trop Med Hyg 54: 449–457.

22. Tongren JE, Drakeley CJ, McDonald SL, Reyburn HG, Manjurano A, et al. (2004) Antigen detection for diagnosis of malaria. Indian J Med Microbiol 52(2): 200–21.

23. Bouharoun-Tayoun H, Oeuvray C, Lunel F, Druilhe P (1995) Mechanisms underlying the monocyte-mediated antibody-dependent killing of Plasmodium falciparum asexual blood stages. J Exp Med 182: 409–418.

24. Jafarshad A, Dziegiel MH, Lundquist R, Nielsen LK, Singh S, et al. (2007) A novel antibody-dependent cellular cytotoxicity mechanism involved in defense against malaria requires costimulation of monocytes FcgammaRII and FcgammaRIII. J Immunol 178: 3099–3106.

25. Tebo AE, Kremmer PG, Luty AJ (2001) Plasmodium falciparum: a major role for IgG3 in antibody-dependent monocyte-mediated cellular inhibition of parasite growth in vitro. Exp Parasitol 98: 20–28.

26. Cavanagh DR, Dobano C, Elhasaan IM, Marsh K, Elhasaan A, et al. (2001) Differential patterns of human immunoglobulin G subclass responses to distinct regions of a single protein, the merozoite surface protein 1 of Plasmodium falciparum. Infect Immun 69(2): 1207–1211.

27. Egan AF, Morris J, Barnish G, Allen S, Greenwood BM, et al. (1996) Clinical immunity to Plasmodium falciparum malaria is associated with serum antibodies to the 19kDa C-terminal fragment of the merozoite surface antigen, PMSP-1. J Infect Dis 173(3): 765–9.

28. Egan AF, Burghaus P, Drulhe P, Holder AA, Riley EM (1999) Human antibodies to the 19kDa C-terminal fragment of Plasmodium falciparum merozoite surface protein 1 inhibit parasite growth in vitro. Parasite Immunol 21(3): 133–9.

29. Mutapi F, Ndlovu PD, Hagan P, Woolhouse ME (2000) Antibody-dependent cellular cytotoxicity in children coinfected with malaria. Parasite Immunol 22(4): 207–9.

30. Remoué F, Diallo TO, Angél V, Hervé M, de Clercq D, et al. (2003) Malaria co-infection in children influences antibody response to schistosome antigens and inflammatory markers associated with morbidity. Trans R Soc Trop Med Hyg 97(5): 361–4.

31. Keitel WA, Kester KE, Asmam WK, White AG, Bond NH, et al. (1999) Phase I trial of two recombinant vaccines containing the 19kDa carboxy terminal fragment of Plasmodium falciparum merozoite surface protein 1 (msp-1(19)) and T helper epitopes of tetanus toxoid. Vaccine 18(3-6): 531–9.

32. Diallo TO, Spiegel A, Droué F, Lochooa L, Kaslow DC, et al. (2002) Short report: differential evolution of immunoglobulin G1/G3 antibody responses to Plasmodium falciparum MSP1(19) over time in malaria-immune adult Senegalese patients. Am J Trop Med Hyg 66(2): 137–9.

33. Garraud O, Droué F, Holm I, Nguer CM, Spiegel A, et al. (1999) Secretion of parasite-specific immunoglobulin G by purified blood B lymphocytes from immune individuals after in vitro stimulation with recombinant Plasmodium falciparum merozoite surface protein-119 antigen. Immunology 97(2): 204–10.

34. van den Biggelaar AH, van Ree R, Rodrigues LC, Lell B, Deelder AM, et al. (2000) Decreased atopy in children infected with Schistosoma haematobium: a role for parasite-induced interleukin-10. Lancet 356(9243): 1723–7.

35. Garraud O, Perraut R, Droué F, Nambei WS, Tall A, et al. (2002) Regulation of Antigen-Specific Immunoglobulin G Subclasses in Response to Conserved and Polymorphic Plasmodium falciparum Antigens in an In Vitro Model. Infect Immun 70(6): 2820–2827.

36. Widhe M, Ekerfelt C, Forsberg P, Bergström S, Ermerudh J (1998) IgG subclasses in Lyme borreliosis: a study of specific IgG subclass distribution in an interferon-gamma-predominated disease. Scand J Immunol 47(6): 575–81.

37. Winkler S, Willheim M, Baier K, Schmid D, Aichelburg A, et al. (1998) Reciprocal regulation of Th1- and Th2-cytokine-producing T cells during clearance of parasitemia in Plasmodium falciparum malaria. Infect Immun 66(12): 6040–4.

38. Lee EA, Palmer DR, Flaanagan KL, Resse WH, Othihambo K, et al. (2002) Induction of T helper type 1 and 2 responses to 19-kilodalton merozoite surface protein 1 in vaccinated healthy volunteers and adults naturally exposed to malaria. Infect Immun 70(3): 1417–21.