Malaria Transmission and Naturally Acquired Immunity to PfEMP-1

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Why there are so few gametocytes (the transmission stage of malaria) in the blood of humans infected with Plasmodium spp. is intriguing. This may be due either to reproductive restraint by the parasite or to unidentified gametocyte-specific immune-mediated clearance mechanisms. We propose another mechanism, a cross-stage immunity to Plasmodium falciparum erythrocyte membrane protein 1 (PIEMP-1). This molecule is expressed on the surface of the erythrocyte infected with either trophozoite or early gametocyte parasites. Immunoglobulin G antibodies to PIEMP-1, expressed on both life cycle stages, were measured in residents from an area where malaria is endemic, Papua New Guinea. Anti-PIEMP-1 prevalence increased with age, mirroring the decline in both the prevalence and the density of asexual and transmission stages in erythrocytes. These data led us to propose that immunity to PIEMP-1 may influence malaria transmission by regulation of the production of gametocytes. This regulation may be achieved in two ways: (i) by controlling asexual proliferation and density and (ii) by affecting gametocyte maturation.

Transmission of malaria parasites from the human host to the anopheline vector involves the production of gametocytes. These stages arise after the commitment of asexually dividing erythrocytes (RBCs) to a pathway of sexual development. Plasmodium falciparum gametocytes develop through several morphologically distinct stages, designated I to V, within the host RBC over a period of 10 to 14 days (25). Stage I to IV gametocytes sequester from the peripheral circulation during maturation (22, 42, 47). Stage V gametocytes circulate in the bloodstream and after a further 2 to 3 days become infectious to the mosquito vector.

Observations of the natural history of malaria infection in humans point to two important features of the transmission biology of malaria. First, blood slide surveys have shown that both the prevalence and density of P. falciparum gametocytes decline in an age-specific manner in hosts living in areas of intense malaria transmission (14, 37). This decline may result from the development of naturally acquired immunity to gametocytes, although no age-dependent mechanisms of immune-mediated clearance of gametocytes have yet been identified (for a review, see reference 45). Second, studies of within-host parasite dynamics as well as population surveys have shown that there are far fewer gametocytes in the peripheral blood than the circulating asexual forms known as trophozoites (14, 28, 30, 37, 46). This paucity of transmission stages reflects, in part, the life history of zoites (14, 28, 30, 37, 46). This paucity of transmission stages may result in an age-specific manner in hosts living in areas of intense malaria transmission (14, 37). Nonetheless, these aspects of the parasite’s biology cannot fully explain why there are so few transmission stages.

Taylor and Read (45) have put forward two mechanisms to explain the low prevalence and density of gametocytes relative to those of asexual parasitemias: (i) natural selection favors reproductive restraint such that only low numbers of gametocytes are ever produced, and (ii) a gametocyte-specific immune mechanism(s) acts in the clearance of gametocytes at some stage in their development. We favor a third mechanism, one involving naturally acquired immunity to the variant surface antigen designated P. falciparum erythrocyte membrane protein 1 (PIEMP-1) (27).

PIEMP-1 is expressed on the surface of trophozoite-infected RBCs (32) and mediates adhesion to CD36 and other host adhesion ligands (for a review, see reference 20). This molecule is highly immunogenic (29) and undergoes clonal antigenic variation (5, 41) with variant forms differing in both antigenic and adhesive characteristics (43). By analogy with animal model experiments, the sequential expression of different antigenic variants is believed to mediate the persistence of the parasite within the human host (8). In humans, variant-specific agglutinating antibodies reactive to the surface of the trophozoite-infected RBC have been observed; seroconversion occurs after an acute P. falciparum infection (21, 33). This variant-specific immunity is acquired in an age-dependent manner (12, 21, 24) and is associated with protection from clinical disease (12, 34). It is believed that these agglutinating antibodies are directed against PIEMP-1 (12, 50). It has recently been demonstrated with a PIEMP-1 deletion mutant that this is the case in sera from Papua New Guinea (40). Cytophilic immunoglobulin G (IgG) antibodies have been shown to mediate the recognition of PIEMP-1 (40).

The genes involved in the expression of PIEMP-1 molecules have been identified as a multigene family designated var (4, 43, 44). The differential expression of var genes is associated with the expression of antigenically distinct PIEMP-1 molecules with different adhesive properties (43). Trophozoites and gametocytes share the same repertoire of var genes and express the same PIEMP-1 variants at the surface of the infected...
The surface expression of PfEMP-1 is restricted to young gametocytes (stages I and IIa). Based on these molecular data, it has been proposed that immunity against PfEMP-1 variants may limit the numbers of asexual parasites with the potential to become gametocytes and prevent the maturation of sexual stages (27).

To determine whether such an immune mechanism exists and whether it could account for age-specific patterns of gametocytemia, an immunoenpidemiological study was designed. IgG antibody responses were measured to the surface of trophozoite- and stage I and IIa early-gametocyte-infected RBCs of three P. falciparum isolates. Plasma samples from residents aged 2 to 60 years from Madang, Papua New Guinea, where intense year-round transmission of malaria occurs (14), were screened for these responses. Age-specific patterns of immunity are discussed in the context of the potential role of PfEMP-1-specific IgG antibodies in the regulation of the numbers of transmission stages.

**Materials and Methods**

Study population and plasma collection. The study was conducted in five rural villages situated along the Gogol River basin in Madang Province on the north coast of Papua New Guinea (for details, see reference 17). This area experiences intense year-round transmission of malaria (14). In 1993, a cross-sectional microlimetric survey of 555 individuals aged 2 to 60 years was completed. Thick and thin blood smears were made at the time of plasma collection. Thin films were fixed in methanol and then both thick and thin films were stained with 4% Giemsa stain for 20 min, washed, dried, and stored. Both asexual and gametocyte parasitemias were scored by counting the number of parasites per 200 leukocytes (WBCs). A parasite-negative slide was one on which 2,000 WBCs had been viewed and no parasites had been seen. Parasite densities were converted to parasites per microliter of blood by assuming 8,000 WBCs per ml. Plasma was isolated from EDTA-treated blood samples by Histopaque (Sigma, Poole, United Kingdom) separation. Fifty-nine individuals were selected for measurement of antibody recognition to PfEMP-1 from the cross-sectional study population of 555 individuals and divided by age into the following groups (each designation indicates the age range, in years, for that group): 2-to-4, 5-to-9, 10-to-14, 15-to-19, 20-to-30, and 31-to-60. Hyperimmune plasma (HIP) pools were made from samples from 10 immune adults from the cross-sectional study population, and normal human serum (NHS) was made from samples from three non-malaria-exposed Europeans (Oxford BTS, Oxford, United Kingdom). All serum and plasma aliquots were stored at −70°C.

Parasites. P. falciparum isolates Muz 37 and Muz 106 were collected from children presenting with acute symptomatic malaria at aid posts in Madang during a longitudinal cohort study from 1990 to 1991 (17). Isolate 1776 was collected in 1987 (21). RBCs were washed three times after buffy-coat depletion during a longitudinal cohort study from 1990 to 1991 (17). Isolate 1776 was collected in 1987 (21). RBCs were washed three times after buffy-coat depletion in RPMI-HEPES supplemented with 25 mM HEPES and supplemented with 25 mM sodium bicarbonate, 2.6 mM L-glutamine, 300 mM bovine serum albumin and 10 μg of gentamicin per ml (Gibco, Paisley, United Kingdom) (RPMMI-HEPES); the cells were then cryopreserved in liquid nitrogen for adaptation at WTCEID, Oxford. Adaptation of primary isolates to in vitro culture was performed at WTCEID, Oxford. Parasites were cultured according to the method of Trager and Jenson (48). Cells were grown in RPMI-HEPES supplemented with 10% blood type AB sera (from donors residing in a nonmalarious area) in an atmosphere of 5% CO2, 5% O2, and 90% N2 and subcultured into O-positive RBCs (Oxford BTS), and stablizes were frozen down. In some cases, feeder cells (peritoneal-wash mouse cells) were required for initial parasite growth (10^6 cells/ml, added at weekly intervals for up to 3 weeks) (adapted from the protocol outlined in reference 49). Isolates Muz 106, Muz 37, and 1776 were collected 2, 3, and 6 years, respectively, prior to the collection of plasma samples. It has previously been shown that on adaptation of new isolates to in vitro culture, there is a high risk of the loss of expression of PfEMP-1 from the infected RBC surface due to a subtelomeric deletion in chromosome 9 (19). Thus, parasites were selected over a C32 ameloblast in chromosome 9 (19), to minimize the number of null isolates. For trophozoite-infected RBCs, antibody detection was made with a rabbit anti-human IgG (Dako, Cambridge, United Kingdom), followed by a fluorescein isothiocyanate (FITC)-coupled swine anti-rabbit IgG (Dako) containing 50 μg of ethidium bromide per ml (Sigma). Cells were fixed for 1 h with 0.5% paraformaldehyde diluted in human tonicity phosphate-buffered saline–1% bovine serum albumin. A different staining regimen was used for staining human plasma bound to early-gametocyte-infected RBCs; a biotinylated sheep anti-human IgG (Sigma) was followed by fluorescein isothiocyanate-streptavidin (Sigma). This was to avoid the nonspecific binding of goat anti-mouse antibody, used for the detection of internally labelled gametocytes within the RBC, to externally labelled antibodies on the RBC surface. After surface staining, early-gametocyte-infected RBCs were fixed with 2% paraformaldehyde overnight at 4°C and permeabilized with 0.01% Triton X-100 to allow labelling of the early gametocyte within the infected RBC. Staining was done with a gametocyte-specific mouse monoclonal antibody, 2G7 (10), followed by phycoerythrin-conjugated goat anti-mouse IgG (Sigma). Both trophozoite- and gametocyte-infected RBCs were read on an EPICSXL counter (Couler Electronics), where 1,000 infected RBCs were counted. Mean fluorescence intensity (MFI) and the percentage of fluorescence-positive infected RBCs (IRBCs) were calculated from the following formula: (IRBCs of test plasma sample – IRBCs of test plasma sample) / (IRBCs in NHS – IRBCs in NHS).

**Data analysis.** Logistic regression analysis was used to look for age trends in parasitemias and antibody responses. The Spearman rank correlation coefficient test was used to compare the density of parasites to host age and to compare individual antibody titers to trophozoite- and early-gametocyte-infected RBCs from the same isolate and to trophozoite- or early-gametocyte-infected RBCs from different isolates. The Kruskal–Wallis H test was used to compare data between age groups. SPSS 7.0 for Windows was used for all statistical analyses.

**Results**

The age-specific density of asexual parasitemia and gametocytemia was measured in a cross-sectional survey of 555 individuals by blood slide positivity. This measure detects both young asexual stages (i.e., ring-infected RBCs) and mature-gametocyte-infected RBCs in the peripheral circulation. As expected from previous studies (37), the density of mature-gametocyte-infected RBCs was lower than that of asexual stages (Fig. 1); the density of gametocytes was an order of magnitude lower in most age classes (Fig. 1). An age-specific decrease in the densities of asexual parasitemia and gametocytemia was observed (Fig. 1). The decrease in asexual parasite density occurred in the 10- to 14-year-old age group such that individuals aged over 10 years had significantly lower parasite densities (<1,800/μl) than those under 10 (>3,000/μl) (χ^2 = 18.4; P < 0.001). For gametocyte densities, the drop occurred in the 5- to 9-year-olds such that densities were significantly lower in those over 5 years old (<200/μl) in comparison to 2- to 4-year-olds (>700/μl) (χ^2 = 4.7; P < 0.05).
IgG-specific antibodies to the surface of trophozoite- and early-gametocyte-infected RBCs from three Papua New Guinea isolates were measured in 59 individuals by flow cytometric methods. To account for the nonspecific binding of IgG antibodies, normal human serum reactivity was subtracted from test values according to the formula given in Materials and Methods. The age-specific prevalence of IgG antibodies to the surface of both trophozoite- and early-gametocyte-infected RBCs from three isolates is shown in Fig. 2. Age had a significant effect on IgG prevalence to trophozoite- and gametocyte-infected RBCs from Muz 37 and Muz 106 such that near-complete seroconversion was seen in those subjects who were over 10 years old (logistic regression; \( P < 0.05 \)). For isolate 1776, seroconversion to PfEMP-1 was acquired more slowly with age, reaching maximal prevalence in those subjects over 15 years old (\( P < 0.05 \) and \( P = 0.063 \) for trophozoite- and gametocyte-infected RBCs, respectively). In the 2- to 4-year-old age group, 2 of 10 individuals were seropositive for trophozoite- but not for gametocyte-infected RBCs of Muz 37. The inverse was true for isolate 1776, such that two sera were positive for gametocyte- but not for trophozoite-infected RBCs. On closer inspection of the data, both plasma samples responsive to Muz 37 and one of those responsive to 1776 were found to be borderline positive and were classed as seronegative. For all seropositive individuals, the MFI of the IgG response was similar across all age groups for all isolates and both life stages; for Muz 37, the range of reactivity for each age class was as follows: the MFI range was 40 to 229, 28 to 232, 60 to 328, 21 to 632, 19 to 257, and 21 to 345 for the 2-to-4, 5-to-9, 10-to-14, 15-to-19, 21-to-30, and 31-to-60 groups, respectively. Ten individual plasma and HIP samples never reacted to the surface of late-gametocyte-infected RBC (26).

Other studies have clearly demonstrated the variant specificity of the immune response to PfEMP-1 on trophozoite stage-infected RBCs (21, 33). Using the Spearman rank correlation coefficient test we compared the MFI values of samples from 29 patients to the surface of trophozoite- and early-gametocyte-infected RBCs. Children aged 2 to 14 years were chosen, since those older than 14 years were more likely be positive to all three isolates, as seen from age-prevalence profiles shown in Fig. 2. Antibodies to gametocyte stages from Muz 37, Muz 106, and 1776 did not correlate, as would be expected for variant-specific immunity. Curiously, though, plasma responses to the trophozoite stages of Muz 37 and Muz 106 did correlate \((r = 0.58; P < 0.01)\). This would suggest that only a subpopulation of variants from these two isolates converted to gametocytes, explaining the lack of correlation between antibody responses to gametocyte stages from Muz 37 and Muz 106.

Recent molecular data have demonstrated that asexual stages and gametocytes from one isolate share the same repertoire of var genes and that both stages express PfEMP-1 (27). If PfEMP-1 were the only major immunogenic variant surface antigen on both life-cycle stages, we would expect to see a strong correlation in individual immune responses to both life-cycle stages of a single isolate. Using all of the age cohort individuals, we compared plasma reactivity to the surface of trophozoite-infected RBCs and to the surface of early-gametocyte-infected RBCs from the same isolate. The IgG response to trophozoite-infected RBCs correlated significantly \((P < 0.01)\) with that to the homologous early-gametocyte-infected RBCs for each of the three isolates Muz 37, Muz 106, and 1776 \((r = 0.549, r = 0.468\) and \( r = 0.371\), respectively). Thus, we hypothesize that the major immunogen(s) on the surface of the gametocyte-infected RBC is the same as that on the trophozoite-infected RBC, i.e., PfEMP-1.

**DISCUSSION**

The immunoepidemiological data presented in this paper provide the first description of an age-dependent naturally acquired immune response to the transmission stages of *P. falciparum*. Unlike transmission-blocking immunity, this re-
response is functional in the human host rather than in the mosquito vector. The observed IgG antibody response is generated by exposure to PfEMP-1 variants expressed on the surface of RBC infected either with gametocytes or mature asexual stages. This immunity has the potential to regulate the densities of both gametocytes and trophozoites by antibody-dependent cell-mediated cytotoxicity reactions. The magnitude of the asexual parasite population in a human host relative to the gametocyte population indicates that seroconversion to PfEMP-1 variants will occur predominantly as a result of cumulative exposure to trophozoite-infected RBCs. The size of the repertoire of PfEMP-1 variants in the parasite population of any area where disease is endemic is unknown but likely to be large, as there are 50 to 100 var genes per haploid genome (44). The high levels of incidence of malaria infection (9) and the 5 to 10 years of exposure required to seroconvert to any PfEMP-1 variant (21, 24) are consistent with a large repertoire of PfEMP-1 variants in this area of endemicity. The prevalence of these variant-specific responses increased in the age groups (5 to 14 years) where gametocyteemia and asexual parasitemia were declining. The decline in gametocyteemia compared to that in asexual parasitemia occurred earlier in this data set. Previous analyses of larger data sets from Madang (14, 17) demonstrated that both asexual and gametocyte densities peak in the 1- to 4-year-old subjects and decline with age thereafter. The small size of our data set and the greater variability in asexual parasitemia, compared to the low levels of gametocyte density near the limit of sensitivity, most likely account for the observed differences. This gives weight to the larger analyses performed in the same area of endemicity (14, 17), with the decline of asexual stage and gametocyte densities occurring in the age classes where seroprevalence to PfEMP-1 variants increases, as measured in this study.

Peak seroprevalence to isolate 1776 occurred later, with respect to age, in comparison to Muz 37 and Muz 106. This is consistent with the existence of rare and common variants proposed by Bull et al. (11). The apparent persistence of PfEMP-1 antibodies in the older age classes may be explained by long-lived immunity to PfEMP-1 variants or repeated boosting of PfEMP-1 immunity by low-density infections.

To date, immunity to the surface of the trophozoite-infected RBC is believed to be directed to PfEMP-1 (40, 50). Recently, however, data are emerging about the genes *stevor* and rif, which may also encode variant surface molecules in asexual blood stages (16). The products of rif (rifin proteins) are expressed on the surfaces of infected RBCs and are phenotypically variable (31). Immune sera, however, failed to immunoprecipitate proteins of molecular weight comparable to that of rifin proteins, and antisera raised to these proteins did not recognize the surfaces of infected RBCs (31). Whether these molecules are exposed at the exterior surface of the infected RBCs of both trophozoites and gametocytes and are immunogenic is thus unclear. The correlation between immune responses to trophozoite- and gametocyte-infected RBCs in individuals suggests that gametocytes will express the same variant antigenic repertoire. In a previous study, trophozoite-stage and gametocyte-stage parasites were coagglutinated by hyperimmune plasma (27), supporting the findings of our correlation analysis. Correlations, however, were never 100%, but in culture a clonal population of parasites can express multiple variants, as previously reported in our laboratory, at any one point in time (27), although only one variant is expressed by one individual parasite (15). Both asexual and gametocyte cultures were grown in parallel; the only difference was the addition of fresh RBCs to asexual cultures. Differences could not be due to mixed parasite populations, since microsatellite typing (1) of these isolates revealed that they were mixtures of genetically distinct parasite clones (39a). We cannot rule out the possibility that unknown antigens coordinately expressed with PfEMP-1 may also be recognized by plasma and may explain any differences observed. However, these plasmas do not recognize the surface of trophozoite-infected RBCs from the mutant cell line C10 (39a), which does not express PfEMP-1 at the infected RBC surface.

Observations of the intrahost dynamics of *P. falciparum* demonstrate that the asexual stages do not commit to sexual development until after a peak density of asexual parasitemia is reached (13). Recent data indicate that commitment is correlated with a decrease in parasite growth rate (20a). This life history strategy could be selected by immunity to PfEMP-1 variants. The asexual parasites expressing PfEMP-1 variants to which there is no pre-existing immunity will replicate successfully and also give rise to early gametocytes expressing the same PfEMP-1 variants (27). During subsequent development (stages Ib to V), PfEMP-1 expression is lost from the surface of the infected RBC, rendering the mature gametocyte safe from anti-PfEMP-1 immunity (27). Thus, the relatively few gametocytes observed within a host may be the consequence of immunity to predominantly PfEMP-1 expression, regulating the proliferation of the asexual population but with the potential to also regulate the maturation of transmission stages.

Variant-specific immunity to PfEMP-1 could also generate the independent transmission dynamics of genetically distinct parasite clones cohabiting the same human host. Such immunity would necessarily generate nonoverlapping infectious periods which would influence parasite mating patterns in the mosquito vector. The observed excess of homozygous infections in mosquitoes that arises from blood meals in human hosts harboring multiple infections from the same study area in Madang, Papua New Guinea (39), may be due to the above-mentioned variant-specific gametocyte dynamics.

Taylor and Read (45) have made a strong argument that the paucity of transmission stages in residents from areas of endemicity is due to reproductive restraint, possibly selected by density-dependent mechanisms directed against sexual stages (i.e., transmission-blocking immunity in the mosquito vector). They have argued that the observation of higher gametocyte densities induced in semi-immune hosts following intensive vector and drug control demonstrates that gametocyte-specific immunity, controlling sexual parasite densities, does not exist. Here, we describe a novel immune response to asexual and gametocyte stages with the potential to regulate populations of both life cycle stages. The ability to induce higher gametocyte densities in semi-immune individuals can occur as a result of the expansion-gametocyte conversion of the subpopulation of trophozoites to which there is no pre-existing variant-specific immunity.

Population studies of other *Plasmodium* spp. infecting humans have also shown a paucity of transmission stages (for a review, see reference 45). Serological studies demonstrate the existence of variant-specific antigens on the surface of *Plasmodium vivax*-infected RBCs (36), as well as *Plasmodium* spp. infecting nonhuman primates (3, 7, 8, 23). It may be that the same variant-specific immune mechanism is operational in *P. vivax* infection with consequent regulation of transmission.

The view has previously been held that the decline in gametocyteemia along with asexual parasitemia is regulated by the immune response to the asexual stages (6, 35). Here, we propose that immunity to PfEMP-1 could regulate both trophozoite and sexual stage densities. The similarity in the slope of the age-specific curve of trophozoite and gametocyte densities from a large data set such as the Garki project (37) supports
this opinion. There is some evidence, however, that gameto
cyte-specific immune responses can regulate gametocytemia independent of asexual-stage-specific immunity (2). A study in
Irian Jaya, comparing natives to transmigrants from Java, ob-
served that asexual stage prevalence did not differ significantly between natives and transmigrants but that gametocyte prev-
alence was lower among Irianese than Javanese subjects; one
year on, however, a similar decline in the prevalence of gameto-
cyttes was also observed in the Javanese population. In the
light of our data showing cross-stage immunity, one would ex-
pect that the prevalence of both parasitic stages would de-
cline with age in the same way. Recent observations by Bruce
(9) may explain this discrepancy. Bruce has shown, from re-
sampled parasite in children in an area of endem-
imcy in Papua New Guinea, that densities of asexual parasites remain stable regardless of the different numbers of infec-
tions between individuals. To explain this, Bruce proposes a mech-
anism of density-dependent regulation of asexual parasite
numbers. If gametocyte production is related to the number of
new infections acquired by the host, then density dependence could account for the observed differences in the behavior of asexual and gametocyte prevalence in transmigrants living in
Irian Jaya, given that the immunity generating density depen-
dence would be acquired rapidly after 1 year of exposure. Nonethe-
less, in areas of stable malaria transmission, immunity to
PIEMP-1 could contribute to the age-specific patterns of malaria parasitemia and indeed gametocytemia.

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Vol. 67, 1999 IMMUNE-MEDIATED REGULATION OF MALARIA TRANSMISSION 6373


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