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**Citation for published version:**
Kyriacou, HM, Stone, GN, Challis, RJ, Raza, A, Lyke, KE, Thera, MA, Koné, AK, Doumbo, OK, Plowe, CV & Rowe, JA 2006, 'Differential var gene transcription in Plasmodium falciparum isolates from patients with cerebral malaria compared to hyperparasitaemia' Molecular and Biochemical Parasitology, vol. 150, no. 2, pp. 211-218. DOI: 10.1016/j.molbiopara.2006.08.005

**Digital Object Identifier (DOI):**
10.1016/j.molbiopara.2006.08.005

**Link:**
Link to publication record in Edinburgh Research Explorer

**Document Version:**
Publisher's PDF, also known as Version of record

**Published In:**
Molecular and Biochemical Parasitology

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Download date: 24. Apr. 2019
Differential var gene transcription in Plasmodium falciparum isolates from patients with cerebral malaria compared to hyperparasitaemia

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Received 10 July 2006; received in revised form 4 August 2006; accepted 8 August 2006
Available online 5 September 2006

Abstract

The Plasmodium falciparum variant erythrocyte surface antigens known as PfEMP1, encoded by the var gene family, are thought to play a crucial role in malaria pathogenesis because they mediate adhesion to host cells and immuno-modulation. Var genes have been divided into three major groups (A, B and C) and two intermediate groups (B/A and B/C) on the basis of their genomic location and upstream sequence. We analysed expressed sequence tags of the var gene DBL/H9251 domain to investigate var gene transcription in relation to disease severity in Malian children. We found that P. falciparum isolates from children with cerebral malaria (unrousable coma) predominantly transcribe var genes with DBL/H92511-like domains that are characteristic of Group A or B/A var genes. In contrast, isolates from children with equally high parasite burdens but no symptoms or signs of severe malaria (hyperparasitaemia patients) predominantly transcribe var genes with DBL/H92510-like domains that are characteristic of the B and C-related var gene groups. These results suggest that var genes with DBL/H92511-like domains (Group A or B/A) may be implicated in the pathogenesis of cerebral malaria, while var genes with DBL/H92510-like domains promote less virulent malaria infections.

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Keywords: Plasmodium falciparum; Virulence; Pathogenesis; Cerebral malaria; Hyperparasitaemia; PfEMP1

1. Introduction

Plasmodium falciparum malaria continues to exact a huge toll on health and life throughout the tropics, predominantly among young children in sub-Saharan Africa [1]. The clinical manifestations of falciparum malaria differ markedly from infection to infection; disease symptoms often remain mild, but in some cases severe complications such as cerebral malaria, severe malarial anaemia or respiratory distress develop [2]. The reasons why certain children develop life-threatening complications, while others are able to tolerate very high parasite burdens without severe clinical symptoms remain unclear [3].

Severe malaria has previously been associated with expression of a restricted and antigenically conserved subset of variant erythrocyte surface antigens [4,5], suggesting that expression of certain surface molecules may be associated with specific disease manifestations. The major candidate molecules are the PfEMP1 family (Plasmodium falciparum erythrocyte membrane protein one) encoded by the var genes [6,7]. PfEMP1 is thought to play a crucial role in host–parasite interaction because it mediates adhesion to host cells to bring about sequestration and immuno-modulation [3]. Each parasite has a repertoire of 50–60 var genes [8] and variant surface antigen expression is regularly switched to avoid host recognition [9]. The extracellular portion of PfEMP1 variants contains between two and seven cysteine-rich domains known as Duffy-binding-like (DBL) domains [8].
DBL domains have been classified into five types, \( \alpha-\varepsilon \), on the basis of consensus motifs, plus a sixth heterogeneous group, DBL-X [8,10]. The PfEMP1 DBL\( \alpha \) domain occurs towards the N-terminus of the protein and is the most well-conserved DBL domain. Two distinct types of DBL\( \alpha \) domain have been described (\( \alpha \) and \( \alpha_1 \)), which differ in the number of conserved cysteines and other hydrophobic residues [11]. For clarification purposes we have here named the two distinct types of DBL\( \alpha \) domain as DBL\( \alpha_0 \) and DBL\( \alpha_1 \).

On the basis of the full genomic sequence of the 3D7 P. falciparum clone, var genes have been classified into three major groups (A, B and C) and two intermediate groups (B/A and B/C) defined by the presence of one of three conserved 5' upstream sequences (UpsA, UpsB or UpsC), and by the gene’s position and orientation within a chromosome (summarised in Table 1) [8,12,13]. The Group A and B/A genes are larger and have a more complex domain structure than the other groups [12]. All of the Group A genes have a DBL\( \alpha_1 \) type N-terminal domain, and two out of four B/A genes have a hybrid DBL\( \alpha \) domain with some DBL\( \alpha_1 \)-like features including the lower number of conserved cysteine residues [11]. All of the 3D7 Group B, B/C and C var genes have DBL\( \alpha_0 \) N-terminal domains [11,12]. Amplification of expressed DBL\( \alpha \) sequence tags using degenerate primers [14] gives approximately 300–400 base pairs of sequence that can be used to assign expressed var genes to either DBL\( \alpha_1 \)-like sequences (Group A or B/A) or DBL\( \alpha_0 \)-like sequences (Group B, C, B/C or B/A) on the basis of the number of conserved cysteine residues in the amplified tag sequence (Table 1).

It has been suggested that the var gene groups may recombine separately [12,13], and there is some evidence for different functional properties between groups. Binding to the endothelial receptor CD36 is a property of Group B, B/C and C genes but not Group A genes [11], whereas binding to uninfected erythrocytes to mediate rosetting (a virulence-associated phenotype) [15,16] is a property of the DBL\( \alpha_1 \) domain of some Group A genes [17,18]. A recent experiment showed up-regulation of some Group A var genes in a P. falciparum laboratory clone selected using semi-immune serum to express variant surface genes [17,18]. We aimed to investigate the hypothesis that parasite virulence and clinical disease severity in African children are associated with the transcription of distinct subsets of P. falciparum var genes. We examined P. falciparum isolates from patients with one of the commonest and most virulent forms of severe malaria, cerebral malaria (unrousable coma), compared to isolates from controls with equally high parasite burdens but non-virulent disease (hyperparasitaemia patients). Our data show that P. falciparum isolates from cerebral malaria patients are significantly more likely to transcribe var genes with DBL\( \alpha_1 \)-like domains characteristic of Group A and B/A var genes than isolates from patients with non-severe hyperparasitaemia, which predominantly transcribe var genes with DBL\( \alpha_0 \)-like domains.

2. Materials and methods

2.1. Sample collection and RNA extraction

The samples were collected as part of the Bandiagara Malaria Project case-control study of severe malaria that has been described in detail previously [26,27]. Blood samples were collected from children with malaria after informed consent from parents or guardians, and all protocols received institutional review board approval. Blood samples were suspended in glycerol after lymphocyte separation via density centrifugation and frozen to \(-70^\circ\text{C}\). A blood spot from each isolate was made on 3 MM Whatman paper for genomic DNA extraction. Frozen samples were shipped to Edinburgh where they were thawed by standard methods and cultured for 8–24 h until the parasites were a mixture of late ring stage and early pigmented-trophozoite stage (morphologically equivalent to approximately 16–20 h post-invasion in a laboratory strain). This was done in order to study parasite stages in which the full length var mRNA is present [28], while avoiding the early ring stage, when some authors claim multiple non-specific var transcription occurs [29]. The cultured cells were solubilised in Trizol and stored at \(-70^\circ\text{C}\). RNA was extracted as described [28].

<table>
<thead>
<tr>
<th>var gene group</th>
<th>Upstream sequence</th>
<th>Position</th>
<th>Orientation (direction of transcription)</th>
<th>No. of genes in 3D7</th>
<th>No. of DBL domains</th>
<th>DBL( \alpha_0/\alpha_1 )</th>
<th>No. of cysteines in amplified tag</th>
</tr>
</thead>
<tbody>
<tr>
<td>A</td>
<td>UpsA</td>
<td>Subtelomeric</td>
<td>Telomeric</td>
<td>10</td>
<td>2–5</td>
<td>DBL( \alpha_1 )</td>
<td>2</td>
</tr>
<tr>
<td>B/A</td>
<td>UpsB</td>
<td>Subtelomeric</td>
<td>Centromeric</td>
<td>4</td>
<td>4–7</td>
<td>DBL( \alpha_1 ) or DBL( \alpha_0^{*} )</td>
<td>2 or 4</td>
</tr>
<tr>
<td>B</td>
<td>UpsB</td>
<td>Subtelomeric</td>
<td>Centromeric</td>
<td>21</td>
<td>2–3</td>
<td>DBL( \alpha_0 )</td>
<td>4</td>
</tr>
<tr>
<td>B/C</td>
<td>UpsB</td>
<td>Centromeric</td>
<td>Telomeric</td>
<td>10</td>
<td>2–3</td>
<td>DBL( \alpha_0 )</td>
<td>4</td>
</tr>
<tr>
<td>C</td>
<td>UpsC</td>
<td>Central</td>
<td>Telomeric</td>
<td>13</td>
<td>2–3</td>
<td>DBL( \alpha_0 )</td>
<td>4</td>
</tr>
</tbody>
</table>

* Two of the 3D7 Group B/A var genes (PFL0020w and MAL6P1.4) have a DBL\( \alpha_0 \) domain and two (PF08_0140 and MAL6P1.316) have a hybrid DBL\( \alpha_0/\alpha_1 \) domain that is DBL\( \alpha_0 \)-like at the N-terminus but DBL\( \alpha_1 \)-like in the expressed sequence tag region (i.e. it has two rather than four conserved cysteine residues).
2.2. DNA extraction and genotyping

DNA was extracted from blood spots on filter paper using chelex-100 extraction [30]. The minimum number of genotypes per isolate was estimated by genotyping PCR with primers to MSP1 and MSP2 [31].

2.3. cDNA preparation

RNA was treated for 30 min at room temperature with 1.5 units DNAase (Gibco) to remove any contaminating genomic (g) DNA. cDNA was prepared using Superscript First Strand Synthesis System (Invitrogen) with random hexamers according to manufacturers instructions.

2.4. RT-PCR

Reverse-transcriptase (RT)-PCR was used to amplify a region of 300–400 bp of the var DBLα domain using unbiased degenerate primers αAF and αBR [14,21]. These primers are capable of amplifying the entire var gene repertoire of 3D7 apart from the strain-transcending var2CSA gene implicated in malaria in pregnancy [32] and the type 3 var genes that are of unknown function. Amplification conditions were as described [14] with a hot start (95°C, 5 min) followed by 35 cycles of 95°C, 20 s; 42°C, 20 s; 60°C, 1 min. Samples without RT were used in all reactions to exclude gDNA contamination.

2.5. PCR for upstream regions

Upstream PCRs were carried out on gDNA using upstream primers UpsA (5′-TAT TYH ATK TAT TAY ATT TGT TGT A) UpsB (5′-GTT AGA ACA TTT AAA ATT ATA) and UpsC (5′-AVA GAW ATA TGR TAG ATA YAG), based on sequences from 3D7, and a gene-specific DBLα primer for each isolate. Amplification conditions were 35 cycles of 94°C, 5 s; 46°C, 15 s; 60°C, 2 min. The upstream sequence of the predominant var gene from nine isolates was studied. The remaining upstream sequences were not studied due to lack of parasite material.

2.6. Cloning and sequencing

PCR and RT-PCR products were run on agarose gels and extracted (Gel Extraction Kit, QIAGEN) then cloned (TA cloning kit, Invitrogen), and used to transform One Shot TOP10F competent cells (Invitrogen). Transformed cells were grown overnight and individual white colonies were selected for culture. Plasmids were extracted from overnight cultures using a miniprep kit (QIAGEN), and were sequenced using BigDye terminator reaction mix (Applied Biosciences).

2.7. Sequence analysis

Sequences were analysed using Lasergene software (DNASTAR Inc). Contigs were created with a minimum percentage match of 95% to classify sequences for each isolate. Amino acid sequences were aligned using MUSCLE [33] and the alignment converted into a nucleotide alignment with Tranalign (http://bioweb.pasteur.fr/docs/EMBOSS/tranalign.html)

2.8. Phylogenetic network

A phylogenetic network was generated rather than a traditional phylogenetic tree because var genes are subject to recombination. The network allows visualisation of ambiguous and conflicting signals in the dataset due to recombination or other factors [34]. Splits computed from the data are represented as parallel edges rather than single branches and the network provides an explicit representation of evolutionary history [34]. The phylogenetic network was generated in SplitsTree 4.4 [34] using the Neighbor-Net distances transformation [35] and equal angle splits transformation [36].

3. Results

Twenty-six P. falciparum field isolates were obtained from a vaccine trial site in Mali with intense seasonal transmission of P. falciparum (up to 20–60 infected bites per person per month at the peak of the July–December transmission season [37]). Blood was obtained from children with cerebral malaria (unrousable coma with a Blantyre coma score ≤2 and no other detectable cause of coma), non-severe hyperparasitaemia (>500,000 parasites per microlitre of blood but with no symptoms or signs of severe disease) or uncomplicated malaria (fever and P. falciparum parasitaemia with no evidence of severe malaria or hyperparasitaemia). Hyperparasitaemia is generally considered to be an indication of severe malaria using World Health Organisation criteria [38]. However, African children with hyperparasitaemia and no other symptoms or signs of severe disease have a very low mortality rate, and hyperparasitaemia in sub-Saharan Africa can thus be considered as a form of non-severe malaria [2,26].

There were no significant differences in patient age across the three clinical categories (Table 2). The cerebral malaria and hyperparasitaemia patients did not differ significantly in parasitaemia, but the uncomplicated malaria patients had significantly lower parasitaemias than the other two groups (Table 2). We therefore considered the hyperparasitaemia isolates to provide a better control group for the cerebral malaria isolates than the uncomplicated malaria isolates, because the major difference between the cerebral and hyperparasitaemia isolates is in parasite virulence and disease severity rather than parasite burden. Parasite rosette frequency was significantly associated with disease severity, as expected in isolates from Africa (Table 2) [15,16]. Multiple genotypes were present in most isolates (range 1–5), and there was no significant difference in the number of genotypes between the three clinical categories (Table 2).

The transcribed DBLα var gene sequences from each isolate were amplified by reverse transcriptase-PCR using unbiased degenerate primers [14]. The RT-PCR products were cloned and 14–19 recombinant plasmids with var gene inserts were sequenced per isolate. Identical sequences from the same isolate were defined as isolate sequence a, b, c etc. in decreasing order of abundance (GenBank accession numbers DQ367086–DQ367226). The number of distinct DBLα var gene
sequences detected per isolate varied between 1 and 14, and almost every isolate showed a predominant gene (Fig. 1). Identical or almost identical sequences from different isolates were rare, indicating minimal overlap in var gene repertoires between isolates (see Supplementary material for further details). There was no significant difference in the number of distinct DBLα sequences per isolate detected in each clinical category (cerebral malaria: median 5, range 1–9; hyperparasitaemia: median 6.5, range 2–14; uncomplicated malaria: median 4, range 2–10, $\text{P} = 0.72$, Kruskal–Wallis test). Further experiments to validate the reproducibility of these data are shown in the Supplementary material (Fig. S1).

There was a highly significant difference between clinical categories in the percentage of var genes amplified that were DBLα1-like. 80.1% of the var gene sequences amplified from the cerebral malaria isolates were DBLα1-like, compared to only 25.7% of the sequences from the hyperparasitaemia isolates and 40.5% of the sequences from the uncomplicated malaria patients ($P < 0.001$, Chi-square test, Fig. 1). If only the predominant gene from each isolate was examined, a marked difference remained between the cerebral malaria and hyperparasitaemia isolates ($P = 0.013$, Fishers Exact test, Fig. 2), but not between any other group (uncomplicated versus hyperparasitaemia, $P = 0.294$; uncomplicated versus cerebral, $P = 0.131$, Fishers Exact test). There was a significant positive correlation between the proportion of DBLα1-like sequences and the rosette frequency ($\text{Rho} = 0.520$, Spearman Rank correlation).

The contrast between the var genes transcribed by the isolates from cerebral malaria patients compared to hyperparasitaemia patients was also demonstrated by mapping clinical categories across a phylogenetic network of DBLα sequence tags (Fig. 3). A phylogenetic network was used rather than a traditional phylogenetic tree because var genes are subject to recombination, a process that can generate conflicting signals in tree-building and can be a significant source of error [34]. For reference, a selection of DBLα sequence tags from the three major Groups A, B and C from the fully sequenced $P. falciparum$ laboratory clone 3D7 [8] were included, plus four 3D7 genes with intermediate characteristics (B/A or B/C). In the phylogenetic network the sequences fell into two major clades (Fig. 3). All of the sequences to the left of the dotted line are DBLα0-like (i.e. they contain four cysteine residues in the expressed sequence tag), whereas all of the sequences to the right of the dotted line are

### Table 2

<table>
<thead>
<tr>
<th>Malaria disease category</th>
<th>Age (months) (mean ± S.D.)</th>
<th>Parasitaemia (%) (mean ± S.D.)</th>
<th>Rosette frequencya (%) (median:25th, 75th percentiles)</th>
<th>Genotypes (mean:range)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Uncomplicated</td>
<td>45.8 ± 28.0</td>
<td>3.3 ± 1.5</td>
<td>8.0:0.0, 15.8</td>
<td>2.4 (1–4)</td>
</tr>
<tr>
<td>Hyperparasitaemia</td>
<td>42.5 ± 16.9</td>
<td>11.9 ± 3.9</td>
<td>9.5:0.0, 18.0</td>
<td>2.6 (1–4)</td>
</tr>
<tr>
<td>Cerebral malaria</td>
<td>35.2 ± 22.0</td>
<td>12.3 ± 6.3</td>
<td>22.0:19.8, 46.8</td>
<td>2.2 (1–5)</td>
</tr>
</tbody>
</table>

$a$ Percent of mature-stage infected erythrocytes that bind two or more uninfected erythrocytes.
$b$ Uncomplicated: cerebral $P = 0.46$, uncomplicated: hyperparasitaemia $P = 0.72$, cerebral:hyperparasitaemia $P = 0.68$, Student’s $t$-test.
$c$ H: hyperparasitaemia; C: cerebral malaria; U: uncomplicated malaria. Student’s $t$-test.
$d$ Kruskal–Wallis test.
$e$ Uncomplicated: cerebral $P = 0.72$, uncomplicated: hyperparasitaemia $P = 0.72$, cerebral:hyperparasitaemia $P = 0.49$, Student’s $t$-test.
Fig. 2. Alignment of DBL0 domains. A MUSCLE alignment of the predominant gene transcribed by *P. falciparum* isolates from cerebral malaria and hyperparasitaemia patients. The second half of the expressed sequence tag is shown and the position of the residues that distinguish DBL0-like domains (cysteines present) from DBL1-like domains (cysteines missing) are arrowed, and the cysteine residues are boxed. The predominant gene in 8/9 isolates from cerebral malaria patients is of the DBL0-like type, whereas the predominant gene in 6/8 hyperparasitaemia isolates is of the DBL1-like type. This difference is significant by Fishers Exact test ($P = 0.013$). Symbols: (*) indicates conserved residues; (: ) indicates conservative substitution; (·) indicates semi-conservative substitution.

Fig. 3. Phylogenetic network showing relationships between *var* gene DBL0 sequence tags transcribed by *P. falciparum* isolates, generated using Neighbour-Net [35]. Sequences transcribed by isolates from African children with cerebral malaria (CM, red), hyperparasitaemia (HYP, blue), and uncomplicated malaria (U, green) are compared to a selection of *var* genes from the laboratory clone 3D7. For the 3D7 genes, the gene name is preceded by A, B, C, BA or BC to indicate the group to which the gene belongs. The sequences fall into two major clades, with the DBL0-like sequences to the left of the dotted line and the DBL1-like sequences to the right. For the predominant gene from nine of the Malian isolates, the *var* gene upstream region was determined by PCR and is indicated as a boxed letter.
DBLα1-like (containing two cysteine residues in the expressed sequence tag) (Fig. 3). A phylogenetic network containing the full repertoire of 3D7 var gene sequence tags plus the Malian sequences was also generated and gave the same separation into two distinct clades of DBLα0-like and DBLα1-like sequences (data not shown due to the complexity of the figure. This is available from the authors on request). Sequences from the cerebral malaria patients (red in Fig. 3) are significantly concentrated into the “DBLα1-like clade” ($\chi^2 = 12.72$ on 1 d.f., $P < 0.001$), while sequences from the hyperparasitaemia patients (blue) are significantly concentrated in a “DBLα0-like clade” ($\chi^2 = 5.6$ on 1 d.f., $P < 0.025$). Sequences from the uncomplicated malaria patients (green) showed no bias in distribution between the two clades (Fig. 3).

Further separation of the DBLα expressed sequence tags into the six sequence groups recently proposed by Bull et al. [21] did not reveal any further differences between the three disease categories (see Supplementary material, Fig. S2). Investigation of the var gene upstream regions for the predominant var gene from nine isolates showed that four field isolate var genes with DBLα0-like domains had UpsB or UpsC upstream regions, whereas four out of five var genes with DBLα1-like domains had UpsA-type upstream regions and the fifth (Hyp4A) had an UpsB sequence linked to a hybrid DBLα0/α1 domain and was therefore a Group B/A var gene (Fig. 3).

4. Discussion

The major finding of this study is that P. falciparum isolates from Malian children with cerebral malaria (a virulent form of disease) predominantly transcribe var genes with DBLα1-like domains that are characteristic of Group A or B/A var genes, whereas isolates from patients with hyperparasitaemia (a non-virulent form of disease) predominantly transcribe var genes with DBLα0-like domains. These results provide the first firm evidence from P. falciparum clinical isolates to support the hypothesis put forward previously that Group A var genes (which all have DBLα1 domains) could be implicated in the pathogenesis of severe malaria in African children [19].

Previous studies on var gene transcription in clinical isolates have given conflicting results. A role for var genes with DBLα1-like domains in the multi-organ failure type of severe malaria that occurs in adults in low malaria transmission settings was suggested by a study of 10 Brazilian isolates [20]. A study in Papua New Guinea, however, showed upregulation of Group B var genes in clinical disease (severe and uncomplicated) compared to asymptomatic infections [22]. The “severe disease” patients in the Papua New Guinea study had a variety of different clinical syndromes, and strictly defined cerebral malaria (Blantyre coma score $\leq 2$) was not included. A recent study from Tanzania also investigated a mixed group of severe malaria patients and found increased abundance of both Groups A and B var genes in severe compared to uncomplicated and asymptomatic malaria patients [23]. Analysis of a sub-group of seven patients with neurological alterations (Blantyre coma score $\leq 3$, therefore this group does not fulfill a strict definition of cerebral malaria) did suggest a trend towards increased abundance of Group A var genes, but this was not statistically significant [23]. A study of sequence diversity in Kenyan field isolates did not find any association between var gene group and disease manifestation, however only six isolates from assorted severe malaria patients were studied, only one of whom had cerebral malaria [21]. As pointed out previously, severe malaria is a heterogeneous group of syndromes that may have different underlying pathogenic mechanisms, therefore strict definitions of disease are essential in studies addressing malaria pathogenesis [24]. Appropriate controls groups with equivalent parasite burdens are also required [24, 25], and the study reported here is the first to investigate var gene transcription using isolates from non-severe hyperparasitaemia patients as a control group rather than uncomplicated malaria patients with low parasite burdens. Although it remains possible that there are genuine geographical differences in the relationship between var gene transcription and disease severity, we suggest that the reason why a clear pattern emerged from the current study, even though the sample numbers are not large, is because a strictly defined severe disease syndrome (cerebral malaria) was investigated and a non-severe malaria control group with equivalent parasite burdens (hyperparasitaemia) was used.

We also investigated var gene transcription in parasite isolates from children with uncomplicated malaria and lower parasite burdens that are equivalent to the control groups used in previous studies [20–23]. Isolates from these patients showed a mixed pattern of var gene transcription with four out of nine isolates transcribing mostly DBLα1-like var genes, while five out of nine isolates transcribed mostly DBLα0-like var genes (Figs. 1 and 3 and Fig. S2). This is consistent with the idea that non-hyperparasitaemic uncomplicated malaria represents an earlier stage in the natural history or pathogenesis pathway of a malaria infection. At this stage, the host–parasite process has progressed beyond asymptomatic infection and produced symptoms, but it may yet diverge along several possible pathways ranging from spontaneous resolution to hyperparasitaemia, cerebral malaria, other severe syndromes, and death. Our data support the hypothesis that parasites expressing DBLα1-like var genes influence the direction of the disease process toward cerebral malaria, presumably increasing the predilection for parasites to sequester and cause microvascular obstruction in the brain. Other factors are likely to influence this process including host genotype [39], and it is possible that parasite isolates transcribing “cerebral malaria-type” DBLα1-like var genes may fail to cause severe disease in some children because of host genetic polymorphisms that can modify parasite virulence phenotypes [40].

This study and previous similar investigations [20–23] use peripheral blood samples to examine P. falciparum var gene transcription profiles in clinical isolates. One unusual feature of malaria infection is that only immature ring stage parasites are found in the peripheral blood, whereas the mature blood stage parasites (pigmented trophozoites and schizonts) are sequestered in the microvasculature [3]. Therefore it is a major concern in studies of this type whether the parasites in the peripheral blood adequately represent the sequestered population. Ideally studies aimed at examining the relationship between
parasites from patients in other disease categories [49], which could also contribute to the lower virulence of these infections.

In conclusion, we have shown that var genes with DBLα1-like domains (characteristic of Group A and some B/A var genes) are predominantly transcribed in *P. falciparum* isolates from Malian cerebral malaria patients, whereas var genes with DBLα0-like domains are predominantly transcribed in isolates from patients with a non-virulent form of disease (hyperparasitaemia). These findings suggest fundamental differences in the roles played by the different var genes groups in host–parasite interaction that could be related to the contrasting cytoadhesive and immuno-modulatory functions of the var gene groups. Further research is required to examine the disease-associations, functions and diversity of var gene subsets in different geographical areas, and we emphasise the importance of strict clinical definitions and appropriate control groups in future work.

**Acknowledgements**

We are grateful to Sue Kyes, Andrew Read and Brian Charles worth for comments on the manuscript and to the Bandiagara Malaria Project Team and the patients and their guardians for participation in the study. We are also grateful to Martin Jones for bioinformatics advice. This work was supported by the Wellcome Trust (Ph.D. studentship to H.M.K. and Senior Research Fellowship to J.A.R., grant number 067431), the NERC (research grants to G.N.S., and a Ph.D. studentship to R.J.C.), the National Institutes of Health (contract no. N01-AI-85346) and the Fogarty International Center (training grant no. D43TW001589).

**Appendix A. Supplementary data**

Supplementary data associated with this article can be found, in the online version, at doi:10.1016/j.molbiopara.2006.08.005.

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