Molecular analysis of two local falciparum malaria outbreaks on the French Guiana coast confirms the msp1 B-K1/varD genotype association with severe malaria.

Eric Legrand, Beatrice Volney, Anne Lavergne, Caroline Tournegros, Loïc Florent, Doris Accrombessi, Micheline Guillotte, Odile Mercereau-Puijalon, Philippe Esterre

To cite this version:

Eric Legrand, Beatrice Volney, Anne Lavergne, Caroline Tournegros, Loïc Florent, et al.. Molecular analysis of two local falciparum malaria outbreaks on the French Guiana coast confirms the msp1 B-K1/varD genotype association with severe malaria.. Malaria Journal, BioMed Central, 2005, 4, pp.26. 10.1186/1475-2875-4-26. pasteur-00590991

HAL Id: pasteur-00590991
https://hal-riip.archives-ouvertes.fr/pasteur-00590991

Submitted on 30 Jun 2011

HAL is a multi-disciplinary open access archive for the deposit and dissemination of scientific research documents, whether they are published or not. The documents may come from teaching and research institutions in France or abroad, or from public or private research centers.
Malaria Journal

Research

Molecular analysis of two local falciparum malaria outbreaks on the French Guiana coast confirms the msp1 B-K1/varD genotype association with severe malaria

Eric Legrand*1, Beatrice Volney1, Anne Lavergne1, Caroline Tournegros1, Loïc Florent1, Doris Accrombessi1, Micheline Guillotte2, Odile Mercereau-Puijalon2 and Philippe Esterre1

Address: 1Centre National de Référence sur la Chimiorésistance du Paludisme dans la région Antilles – Guyane, Institut Pasteur de la Guyane, BP 6010, F-97306 Cayenne-Cedex, France and 2Unité d’Immunologie Moléculaire des Parasites, Unité de Recherche Associée 2581 du Centre National de la Recherche Scientifique, Institut Pasteur, Paris, France

Email: Eric Legrand* - elegend@pasteur-cayenne.fr; Beatrice Volney - bvolney@pasteur-cayenne.fr; Anne Lavergne - alavergne@pasteur-cayenne.fr; Caroline Tournegros - cnrcp@pasteur-cayenne.fr; Loïc Florent - cnrcp@pasteur-cayenne.fr; Doris Accrombessi - cnrcp@pasteur-cayenne.fr; Micheline Guillotte - mguillot@pasteur.fr; Odile Mercereau-Puijalon - omp@pasteur.fr; Philippe Esterre - pesterre@pasteur-cayenne.fr

* Corresponding author

Abstract

Background: Plasmodium falciparum outbreaks can occur in the coastal area of French Guiana, where the population is essentially non-immune. Two sporadic outbreaks were observed, including one with severe malaria cases. To characterize these outbreaks and verify previous observations of specific genotype characteristics in severe malaria in this area, all cases from each outbreak were studied.

Methods: P. falciparum genotypes for six genetic loci were determined by PCR amplification from peripheral blood parasites. The msp1/block2 and msp2 genotypes were determined by DNA sequencing. Microsatellite and varD genotyping was based on size polymorphism and locus-specific amplification.

Results: The outbreak including severe malaria cases was associated with a single genotype. The other mild malaria outbreak was due to at least five distinct genotypes.

Conclusion: Two distinct types of outbreak occurred despite systematic and sustained deployment of malaria control measures, indicating a need for reinforced vigilance. The varD/B-K1 msp1 linkage and its association with severe malaria in this area was confirmed.

Background

The annual malaria incidence in French Guiana has increased ten-fold during the last 30 years, reaching now-adays approximately 3%, with 60% due to Plasmodium falciparum and 40 % due to Plasmodium vivax. French Guiana is an area with multidrug resistant P. falciparum malaria. Transmission occurs in isolated foci located in settlement pockets within the Amazonian forest and along the rivers. The main malaria-endemic areas are located along the Maroni and Oyapock rivers, which serve as natural
frontiers with Surinam and Brazil, respectively [1]. As a result of three-monthly insecticide spraying campaigns, there are virtually no malaria cases along the coast, where 80% of the population resides. However, outbreaks, which usually are of short duration and affect a small number of patients, are occasionally reported in the coastal area.

The *P. falciparum* parasite population of French Guiana presents a remarkably low degree of polymorphism, with a clonal type structure [2]. The parasite diversity is so restricted that the circulation of specific genotypes could be followed within the area, making it possible to investigate the clinical impact of specific parasite genotypes. A comparative analysis of parasite genetic characteristics in isolates collected from mild and severe malaria patients has highlighted a significant linkage disequilibrium between a particular *msp1/block2* allele that was called B-K1 and a particular *var* gene called *varD* in isolates from patients with severe *P. falciparum* malaria [3].

To verify this association and better understand the possible cause of local outbreaks in the coastal area located away from the main endemic sites, two *P. falciparum* outbreaks were studied, one with five cases, including two cases of severe malaria and one death, the other with nine mild malaria cases. The isolates from each case were characterized for both outbreaks using a six-locus genotyping approach, including *varD*, *msp1/block2*, *msp2* [2-7], as well as three microsatellite loci [8].

**Methods**

**Parasite isolates and patient information**

The outbreaks occurred in Macouria and Matoury, both located on the French Guiana coast, an area with unstable, low transmission, away from the main endemic areas along the Maroni and the Oyapock (Fig. 1). The international airport close to Cayenne, the main city, is in Matoury. The isolates were collected from patients with *P. falciparum* malaria in health centres and sent to the laboratory for routine drug susceptibility phenotyping and genotyping analysis. The policy of the malaria control service is to get an *in vitro* susceptibility profile for each diagnosed *P. falciparum* malaria case. To this end, a 5 mL blood sample was systemically collected by venepuncture in EDTA tubes and maintained at 4°C during shipment to the laboratory. The samples were immediately processed for *in vitro* susceptibility assays and a 0.5 mL aliquot was frozen at -20°C for subsequent DNA preparation. The five Macouria isolates were collected in June and July 2001 from three mild (E57, E64 and E72) and two severe cases (E61, a fatal case, and E62). Patient E57 had visited the endemic Macapa region in Brazil one month before the malaria attack. The other four Macouria patients did not report having left the area. The nine Matoury isolates, collected from February to June 2002 originated from mild malaria cases, none of whom reported a recent visit to an endemic area.

**DNA extraction and genotyping**

DNA was extracted as described [4]. Genotyping was performed by PCR using the primers listed in Figure 2. For *msp1/block2*, *msp2* and *varD*, PCR was performed in a 50 µL final volume containing the DNA template, 1.5 mM MgCl$_2$, 2 µM each primer, 200 µM dNTP, and 1.5 units of AmpliTaq Gold polymerase (Applied Biosystems, Foster City, CA, USA). The PCR conditions for *msp1/block2* and *msp2* were: one cycle at 95°C for 5 min, followed by 40 cycles at 94°C for 1 min, 56°C for 2 min, 72°C for 2 min, and final extension at 72°C for 10 min. The *varD* PCR conditions were: one cycle at 94°C for 5 min, followed by 35 cycles at 94°C for 10 sec, 61°C for 1 min 30 sec, 72°C for 2 min, and one cycle at 72°C for 10 min. The PCR products were analysed by agarose gel electrophoresis and directly sequenced (Qbiogen, Every, France). Sequences were aligned using the Clustal W program, followed by manual analysis using the ED editor of the MUST package [9].

Microsatellite typing was done using the C1M4, C3M27 and C4M69 loci, located on chromosomes 1, 3 and 4, respectively. The locus-specific primers described by Su et al. [8] were used (see Figure 2). Amplification was done in 15 µL reaction volume containing the DNA template, 1.5 mM MgCl$_2$, 2 µM each primer, 200 µM each dNTP, and 0.5 Units Taq polymerase (Promega, Madison, WI, USA). The samples were subjected to 1 cycle at 94°C for 7 min, followed by 35 cycles at 94°C for 30 sec, 52°C for 30 sec, 47°C for 30 sec, 72°C for 30 sec, and a final extension step at 72°C for 15 min. The PCR products were analysed by 3% Metaphor agarose gel (FMC Bioproduct, Rockland, Maine, USA) electrophoresis and stained with ethidium bromide. Fragment size was calculated using the Taxotron software (P.A.D. Grimont, Institut Pasteur, Paris).

**Results and discussion**

PCR amplification of the *msp1/block2* and *msp2* *P. falciparum* loci generated a single band for all isolates (Fig. 3). Each PCR band was sequenced. This showed three distinct alleles for each locus. All Macouria samples carried the same 464-bp *msp1* block2 allele, which belonged to the K1-type allelic family, corresponding to the allele called B-K1 in a previous study [3]. A smaller (431 bp) K1-type allele, previously called A-K1 [3], and a 398 pb RO33-type allele were observed in six and three Matoury isolates, respectively. All three RO33-type sequences were identical to the RO33 allele (accession number Y00087). Alignment of the B-K1 and A-K1 deduced protein sequence with sequences available in the databases identified several alleles observed in other regions of the world showing...
up to 99% and 97% identity, respectively. However, no isolate in the database had precisely similar tri-peptide repeats as B-K1 or A-K1. This is in line with recent evidence indicating that \( msp1/ \) block 2 (and \( msp2 \)) repeats evolve at a faster rate than SNPs in the non-repeated domains [10]. Numerous isolates showed in addition single nucleotide polymorphisms in the non-repeated sequences (Fig. 4).

All three \( msp2 \) alleles belonged to the 3D7-type family and were arbitrarily called 3D7a (591 bp), b (561 bp) and c (603 bp). The 3D7a allele was observed in all five
Macouria isolates and in five of nine Matoury isolates. The 3D7b and 3D7c allele were observed in one and three Matoury isolates, respectively (Fig. 3). Alignment of the deduced protein sequence showed that allele 3D7c clustered with a specific group of isolates showing typical particularities in the repeat region in particular the NPPA tetrapeptide interspersed within G-, A-, S- rich repeats, mainly GAGASG (Fig. 5). However, the 3D7c allele was
peculiar within this group in having the largest number of both types of repeats NNPA and presenting GGSAs repeats as well. The 3D7a and 3D7b also presented each a specific arrangement of repeats as well as SNPs in the non-repeated regions.

To further explore the genetic make-up of the *P. falciparum* isolates, three microsatellite loci were studied. A single 140-bp C1M4 product was observed for all isolates, except *isolates* three microsatellite loci were studied. A single

---

**Figure 4**

Deduced amino acid sequence of the French Guiana A-K1 and B-K1 *msp1/*block2 alleles and alignment with alleles described in other areas. Single letter codes are used for amino acids. The name of the reference laboratory strains are indicated. The accession number for K1, FCB1, Palo Alto, NF7 and RO71 were X03371, AF286876, M37213, M19144, X61930. For 3D7, the sequence of the genome sequence (locus PFI475w) is shown. For field isolates the Genbank accession number was used. The reference sequence used was K1. Deletions are indicated by a dash and mutant residues are in red, except for the 3D7-types which are in blue.
infections. This confirms previous findings in this area, where a very high percentage of single infections was observed [2,3], and is in line with reports from other areas of South America [5,6].

French Guiana is a hypoendemic area, where the population is limited, if any, immunity. There are between 6,000–7,000 mild malaria cases and only between four and 20 severe malaria cases each year [1,3]. Previous studies have highlighted specific genotype characteristics in P. falciparum isolates from severe cases in this setting, with the B-K1 msp1 allele in strong linkage disequilibrium with a particular var gene, called varD [3]. The var genes code for the variant adhesins that mediate the cytoadhesion of parasitized erythrocytes to specific host receptors that undoubtedly contribute to severe malaria. As highlighted in Figure 3, all five Macouria isolates, including the two severe cases, had the same 5-loci genotype, suggesting that a single virulent isolate was present. These isolates were undoubtedly a single virulent isolate was present. These isolates were

Figure 5
Deduced amino acid sequence of 3D7-a, b and c msp2 alleles and alignment with alleles described in other areas. Single letter codes are used for amino acids. The name of the reference laboratory strains are indicated. The 3D7 sequence (locus PF03003c) was used as reference sequence. For field isolates the Genbank accession number was used. The Genbank accession number for Palo Alto was M61122. Deletions are indicated by a dash and mutant residues as compared to 3D7 are in red. The various repeat sequence types are underlined or color-coded.
F40 and F96 had a different drug susceptibility profile (data not shown). The Matoury 4 and Matoury 5 genotypes differed by the C4M69 microsatellite locus.

These data confirm the varD/B-K1 msp1 linkage and its association with severe malaria in this area. It is important to note that this association was previously detected in isolates collected in 1994-6, more than six years before the Macouria epidemic. Such a stability over time in a species with a high recombination rate [8] is consistent with previous data pointing to a high selfing rate in this area [3]. The severe malaria cases studied here originated from a geographic area quite distinct from the previous cases reported where the varD/B-K1 association was observed, as illustrated in Figure 1. As observed previously [3], the varD/B-K1 msp1 association was not strictly specific for severe malaria, as the same genotype was also observed in the non-severe Macouria cases. Whether the patients with mild malaria received an earlier treatment compared to those with severe malaria or were less susceptible due to their genetic or immune make up is unclear. The occurrence of two severe cases at one day interval in patients infected with this strain suggests a particular inherent virulence. Whether this is due to the particular msp1 and/or varD allele present is unknown. It may reflect physical association with another locus implicated in virulence. It is worth noting that isolate F52, which harboured varD together with the A-K1 msp1 allele, was collected from a patient with very high peripheral parasitaemia (20%), also considered a sign of severity [11]. VarD is one of approximately 60 members of the P. falciparum var genomic repertoire [12]. The presence of a particular var gene is not synonymous with its expression. Expression of varD was demonstrated in a patient with fatal P. falciparum malaria in a previous study [3], but could not be studied here. The data indicate that future investigations on varD expression in severe and non-severe malaria are warranted. Work is in progress to characterize the full varD gene sequence.

The different genetic profiles of the isolates involved in the two outbreaks reveal distinct onset and dynamics scenarios. Genotyping strongly suggests that the Macouria outbreak was due to one single parasite strain, but the origin of this strain is uncertain. Transmission from E57, the first registered case, to the other cases is unlikely, due to the short delay between the clinical attack and the attack experienced by E61, E62 and E64 (12, 13 and 20 days later, respectively). E67 may possibly have subsequently transmitted the strain to E72, who experienced a malaria attack 37 days later. The diversity of the Matoury isolates indicates that the outbreak was certainly caused by several distinct strains. Matoury accommodates the largest airport in the country and consequently may serve as an occasional transmission focus for parasites originating in neighbouring endemic malaria areas. The possible cause and mode of transmission in this city at that time have not been identified.

**Conclusion**

The results point to two distinct types of outbreak in a region where malaria control measures are systematically deployed and sustained. Reinforced vigilance and rapid case notification are needed to ensure rapid deployment of vector control and personal protection measures to prevent such sporadic epidemics. The Macouria outbreak provoked two severe cases, including one death, a rare event in this area, where health facilities are well-equipped and treatment policy is regularly updated. Parasite genotyping confirmed the association of the B-K1 msp1/varD genotype with severe malaria, reinforcing the notion that some P. falciparum strains might cause more severe infections than others.

**Authors’ contributions**

A. Lavergne and E. Legrand did the sequences analysis. B. Volney, C. Tournegros, D. Acombressi and L. Florent did the laboratory experiments. M. Guilotte and O. Mercereau-Puijalon adapted the microsatellite typing to field isolates. O. Mercereau-Puijalon did the database search. All authors participated in the analysis and interpretation of data. E. Legrand and O. Mercereau-Puijalon wrote the manuscript.

**Acknowledgements**

This project was supported by the French Ministry of Health (InVS agency, Paris) and the French Academy of Sciences (Prix Louis D., 2000). The authors thank Prof. B. Carme and colleagues from the Parasitology Unit of the Central Hospital, Cayenne and Drs. P. Duval and B. Maubert, from the Biomedical Unit of the Institut Pasteur de Guyane, Cayenne for providing the blood samples.

**References**

1. Esterre P, Cordoliani G, Germanetto P, Robin Y: Epidémiologie du paludisme en Guyane française. Bull Soc Pathol Exot 1990, 83:193-205.
2. Ariey F, Chalvet W, Hommel D, Peneau C, Hulin A, Mercereau-Puijalon O, Duchemin J-B, Sarthou J-L, Reyes J-M, Fandeur T: Plasmodium falciparum parasites in French Guiana: limited genetic diversity and high selfing rate. Am J Trop Med Hyg 1999, 61:978-985.
3. Ariey F, Hommel D, LeScarf C, Duchemin JB, Peneau C, Hulin A, Sarthou J-L, Reyes JM, Fandeur T, Mercereau-Puijalon O: Association of severe malaria with a specific Plasmodium falciparum genotype in French Guiana. J Infect Dis 2001, 184:237-241.
4. Contamin H, Fandeur T, Bonnefoy S, Skouri F, Ntoumi F, Mercereau-Puijalon O: PCR typing of field isolates of Plasmodium falciparum. J Clin Microbiol 1993, 31:944-951.
5. Sallenaue-Sales S, Ferreira-da-Cruz HF, Faria CP, Cerruti C Jr, Daniel-Ribeiro CT, Zalis MG: Plasmodium falciparum : limited genetic diversity of MSP-2 in isolates circulating in Brazilian endemic areas. Exp Parasitol 2003, 103:127-135.
6. Tami A, Grundmann H, Sutherland C, McBride JS, Cavanagh DR, Campos E, Snounou G, Barnabe C, Tibayrenc M, Warhurst DC: Restricted genetic and antigenic diversity of Plasmodium falciparum under mesoendemic transmission in the Venezuelan Amazon. Parasitology 2002, 124:569-581.
7. Jelinek T, Kilian AH, Westermeier A, Proll S, Kabagambe G, Nothdurft HD, von Sonnenburg F, Loscher T: Population structure of recrudescent Plasmodium falciparum isolates from western Uganda. *Trop Med Int Health* 1999, 4:476-480.

8. Su X, Ferdig MT, Huang Y, Huynh CQ, Liu A, You J, Wootton JC, Wellems TE: A genetic map and recombination parameters of the human malaria parasite Plasmodium falciparum. *Science* 1999, 286:1351-1353.

9. Philippe H: MUST, a computer package of management utilities for sequence and tree. *Nucleic Acids Res* 1993, 21:5264-5272.

10. Tanabe K, Sakihama N, Kanejo A: Stable SNPs in malaria antigen genes in isolated populations. *Science* 2004, 303:493.

11. World Health Organization: Severe falciparum malaria. *Trans R Soc Trop Med Hyg* 2000, 94(Suppl 1):1-90.

12. Gardner MJ, Hall N, Fung E, White O, Berriman M, Hyman RW, Carlton JM, Pain A, Nelson KE, Bowman S, Paulsen IT, James K, Eisen JA, Rutherford K, Salzberg SL, Craig A, Kyes S, Chan M-S, Nene V, Shalom SJ, Suh B, Peterson J, Angiuoli S, Pertea M, Allen J, Selengut J, Haft D, Mather MW, Vaidya AB, Martin DMA, Fairlamb AH, Fraunholz MJ, Roos DS, Ralph SA, McFadden GI, Cummings LM, Subramanian GM, Mungall C, Venter JC, Carucci DJ, Hoffman SL, Newbold C, Davis RW, Fraser CM, Barel B: Genome sequence of the human malaria parasite Plasmodium falciparum. *Nature* 2002, 419:498-511.