TRANSFUSION-TRANSMITTED MALARIA: CASE REPORT OF ASYMPTOMATIC DONOR HARBORING *Plasmodium malariae*

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**SUMMARY**

Malaria in Brazil is endemic in the Amazon region, but autochthonous cases with low parasitaemia occur in the Atlantic Forest area of the country. According to Brazilian legislation no test is mandatory for blood donors from non-endemic areas. However if they have traveled to malaria transmission regions they are deferred for six months before they can donate. This report describes a transfusion-transmitted malaria case in Sao Paulo, Brazil, where one recipient received infected blood and developed the disease. He lived in Sao Paulo and had no previous transfusion or trips to endemic areas, including those of low endemicity, such as Atlantic Forest. Thick blood smears confirmed *Plasmodium malariae*. All donors lived in Sao Paulo and one of them (Donor 045-0) showed positive hemoscopy and PCR. This asymptomatic donor had traveled to Juquia, in the Atlantic Forest area of Sao Paulo State, where sporadic cases of autochthonous malaria are described. DNA assay revealed *P. malariae* in the donor’s (Donor 045-0) blood. Serum archives of the recipient and of all blood donors were analyzed by ELISA using both *P. vivax* and *P. falciparum* antigens, and IFAT with *P. malariae*. Donor 045-0’s serum was *P. malariae* IFAT positive and the *P. vivax* ELISA was reactive. In addition, two out of 44 donors’ archive sera were also *P. vivax* ELISA reactive. All sera were *P. falciparum* ELISA negative. This case suggests the need of reviewing donor selection criteria and deferral strategies to prevent possible cases of transfusion-transmitted malaria.

**KEYWORDS:** Transfusion-transmitted malaria; PCR; Malaria serology; Malarial DNA; Atlantic Forest.

**INTRODUCTION**

Malaria occurs in more than 100 countries in the tropical and subtropical regions including parts of Asia, Africa, Oceania, Central and South America. Half of the world’s population is at risk of malaria and an estimated 243 million cases led to nearly 863,000 deaths in 2008, mostly in children living in sub-Saharan Africa. In Brazil, 306,908 cases were notified in 2009, being 306,342 in the Amazon region, where malaria is endemic. *P. vivax* is the most common species (84.4%), followed by *P. falciparum* (14.6%), mixed infections (0.9%) and *P. malariae* (0.02%). However, it is important to note that *P. malariae* may be misdiagnosed due to the difficulty of differentiating it from *P. vivax*. Outside of the Amazon region the National Malaria Control Program notified 154 autochthonous cases in 2008.

Data about the frequency of transfusion transmitted malaria show values that vary from less than 0.2 in non-endemic countries to 50 or more cases per million in endemic countries and the species most frequently associated with transfusional cases are *P. falciparum, P. malariae* and *P. vivax*. The donors implicated in this kind of transmission are invariably semi-immune, with parasite levels below the detection threshold of currently available assays. As the result of the asymptomatic persistence of parasites, transmission has been documented after the last exposure as long as 44 years for *P. malariae*, five years for *P. vivax* and eight years for *P. falciparum*. Besides that, parasites can survive in blood at temperatures between 2 °C and 6 °C for until three weeks, with the estimated inoculum in transfusions from one to 10 parasites per donation. Whole blood and red blood cell units are the primary carriers of malaria transmitted by transfusion, although platelets and leukocytes may contain variable numbers of red cells and have, rarely, transmitted malaria.

In endemic regions transfusion transmitted malaria has frequently been described. In a recent prospective study in Pakistan, healthy volunteer blood donors were screened by thick blood smear and showed 0.57% positivity, with one case of *P. falciparum* and eight due to *P. vivax*. In a study conducted in India 11,736 units of blood were screened from 2008 to 2009, with 0.03% positivity using rapid diagnostic tests that was confirmed by microscopy. In Nigeria, where blood donors are not routinely tested for malarial infection, potential blood donors were screened, resulting in a positivity of 20.2% by microscopy, 3.8% by OptiMAL and 57.8% by Clinotech.

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The risk of transfusion-transmitted malaria differs widely among low endemic countries, where the imported infection occurs in individuals that have traveled to or migrated from endemic regions\(^1\). In a survey conducted in 14 national transfusion services centers in different countries, the use of serological tests for malaria in donor screening was reported as a routine only in four countries (United Kingdom, Denmark, Finland and New Zealand). Those with positive serology are confirmed by rapid tests for detection of antigens in New Zealand and nucleic acid testing (NAT) in the United Kingdom where no case of transfusion-transmitted malaria was diagnosed during the last 5 years\(^2\).

According to Brazilian guidelines there are different recommendations for endemic and non-endemic regions. The endemic area is classified into low, medium or high risk, based on the Annual Parasite Index. In endemic areas those donors who have had malaria in the 12 months preceding the donation and those coming from areas of high malaria risk are rejected, as well as those donors that presented fever or are suspected of having malaria in the last 30 days. Donors from areas of medium and low risk can be accepted after a negative thick film or rapid malaria test. In non-endemic areas testing is not required nor performed, but donors who were in an endemic area with active transmission in the last six months and those that have had malaria or resided in endemic areas in the last three years are excluded. In both endemic and non-endemic areas donors who have had infection with *P. malariae* are excluded\(^3\). Some transfusion services use antibody assays such as Indirect Fluorescent Antibody Test (IFAT) and enzyme-linked immunosorbent assay (ELISA) in addition to antigenic tests. Molecular tests are used only for diagnosis or research in reference centers. Even in non-endemic areas and with all the precautionary measures being taken, exclusion of all ‘malaria risk’ individuals is not possible.

A further limitation to travel-based restrictions is that the long periods of asymptomatic carriage, particularly associated with *P. malariae*, also mean that donors can harbor parasites even after their exclusion period. Besides that, current strategies to reduce the likelihood of transfusion transmitted infections in non-endemic countries invariably involve discarding blood from potentially exposed donors, possibly leading to wastage of donations.

In the last 10 years cases of transfusion-transmitted malaria have been reported in a number of countries: two in France\(^4\); two in Italy, three in Brazil\(^5\)\(^6\) and one in USA\(^7\), one in Republic of Korea\(^8\); one in United Kingdom\(^9\). We describe a case of transfusion-transmitted malaria due to *P. malariae* from an asymptomatic donor attended in a private blood bank in the city of Sao Paulo. On August 12\(^{th}\) 2008, a patient was admitted for cardiac surgery and received five units of red blood cells, 10 of cryoprecipitate, 20 of platelets and 15 of fresh frozen plasma from 50 different donors. This patient had always lived in Sao Paulo city and denied any previous transfusion before this surgery or of having traveled to malaria endemic areas, including low-endemicity ones, such as Atlantic Forest. The patient had negative serological tests for malaria before transfusion (Table 1). On October 27\(^{th}\) 2008 he developed fever, chills, arthralgia and sweating. Laboratory tests were carried out and the thick films from peripheral blood showed the presence of *P. malariae*. As this recipient received five units of red blood cells, the respective donors were recalled for epidemiological investigation and to perform thick films, PCR and serology. Four out of five donors attended the recall, all of them living in Sao Paulo. Only one donor (045-0) presented positive tests (Table 1, Fig. 1, Fig. 2). Since 1983 this donor traveled on a number of occasions to Juquia City, located in the Atlantic Forest area of Sao Paulo State, and where sporadic cases of autochthonous malaria are described. His last visit was in December 2007. Previously, he had last donated blood in 1990. On examination his thick film showed scarce parasites without clear species determination (Fig. 1A). Nested PCR based on the detection of species-specific sequences of small-subunit rRNA genes of *Plasmodium*\(^{10}\) was done, and revealed the presence of *P. malariae* in the donor’s blood (Fig. 1B). Sera archives of the recipient and of 48 blood donors were recovered and analyzed for the presence of malarial antibodies using an IgG IFAT with *P. malariae* antigen, an ELISA with *P. vivax* recombinant antigen (rPvMSP1\(_{19}\)) and an ELISA using total extract of *P. falciparum* asexual forms. The *P. malariae* IFAT\(^{11}\) was positive only in donor 045-0 (Fig. 1C). The *P. vivax* ELISA was reactive with the sera from the infected donor (045-0) and two of the other 48 donors’ archive sera (Fig. 2). All archive sera were negative in the *P. falciparum* ELISA. Both the blood donor and the recipient were treated successfully.

**DISCUSSION**

In Sao Paulo State, from 1980 to 2007, 821 autochthonous cases of malaria were reported; 91.6% occurred in the eastern region and these were predominantly due to *P. vivax*, as determined by microscopy; 9.6% of cases were asymptomatic\(^8\). It is important to note that the diagnosis of *P. malariae* is not possible.

### Table 1

<table>
<thead>
<tr>
<th>Tests</th>
<th>Receptor</th>
<th>Infected donor</th>
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<tbody>
<tr>
<td></td>
<td>Before transfusion</td>
<td>After transfusion</td>
</tr>
<tr>
<td>PCR</td>
<td>ND</td>
<td>ND</td>
</tr>
<tr>
<td>TBS</td>
<td>ND</td>
<td><em>P. malariae</em></td>
</tr>
<tr>
<td>IFAT-<em>Pm</em></td>
<td>Negative</td>
<td>ND</td>
</tr>
<tr>
<td>ELISA <em>Pf</em></td>
<td>Negative</td>
<td>ND</td>
</tr>
<tr>
<td>ELISA <em>Pv</em>-MSP1(_{19})</td>
<td>Negative</td>
<td>ND</td>
</tr>
</tbody>
</table>

PCR - polymerase chain reaction; TBS - thick blood smear; IFAT - indirect fluorescent assay test; ELISA - Enzyme-linked immunosorbent assay; ND - not done.
Symptomless infections with low parasitemia, confirmed by molecular tools as *P. malariae*, can occur in the Atlantic Forest area. Transfusion-transmitted malaria is a possibility associated with donors who either live in or have traveled to this area, reinforcing the need to monitor these donations. Three recent cases related to *P. malariae*, two of them in splenectomized receptors, including one lethal, originated from infectious donations from asymptomatic donors. Tests carried out to detect the donors involved in the transmission showed high titers against *P. malariae* in IFAT and positive PCR. They had no report of traveling to Amazon region, although they had visited the Atlantic Forest area in the State of Sao Paulo, where sporadic autochthonous cases have persistently been reported. Similar cases may have occurred without being identified.

According to Brazilian legislation no malaria tests are mandatory in blood donors living in non-endemic areas, but they are deferred for donation for six months if they have traveled to endemic or autochthonous malaria transmission regions.

This report is relevant because the transmission occurred despite a careful interview with the blood donors when they were asked about previous residence in and travel to endemic areas. This measure was not able to prevent transfusion transmission of malaria, since the donor had traveled to an area with very low transmission rates and where symptomless malaria cases occur, mainly due to *P. malariae*. The limited knowledge about the epidemiology of the disease in the low transmission regions by health professionals also contributes to failure in donor screening. In this case, the donor remained asymptomatic and had traveled to a low transmission area eight months before blood donation, indicating that the six months period of deferral on its own is not sufficient in such situations. Furthermore, increasing the deferral period to exclude candidates whose displacement occurred more than six months before the donation could affect blood supply drastically. The use of nested PCR in this case allowed the diagnosis of very low parasitemia in the blood donor, since this method can detect one parasite per microlitre. This result was corroborated with the homologous antigen *P. malariae* in IFAT. This serological technique as well as ELISA is commonly used to detect malarial antibodies. DODERER et al. compared two serological assays and obtained clinical sensitivity of 84.2% and specificity of 99.6% by DiaMed ELISA and 70.5% and 99.6% respectively, using the IFAT method. SEED et al. used an enzyme-linked immunoassay (Newmarket) and the results pointed out to the usefulness of a screening strategy combining antibody testing with a 6-month cellular component restriction period for donors with a declared malarial risk. Recently the restriction period was shortened to 4-month combined with a sensitive antibody screening test for donors with malarial risk and the risk of transfusional malaria in Australia remains low. In France, a multicentre study was performed in nine blood banks to compare IFAT and DiaMed ELISA, obtaining a rate of concordance of 92.6% in retrospective samples and 97% in the prospective group, corroborating with the potential use of ELISA test in blood bank screening, as an alternative to the IFAT in non-endemic areas. Since all human *Plasmodia* have been implicated in transfusional malaria, a new assay based on detection of antibodies anti MSP1α of *P. falciparum*, *P. malariae*, *P. ovale* and *P. vivax* is now available.

Non-endemic countries have discussed measures in order to prevent transfusional malaria. In France, a successful strategy has been in use since 1986, based on self-exclusion, questionnaire, medical interview and serological screening prior to blood donation. Individuals who have had malaria in the past are permanently deferred. These measures lead to the reduction of five- to 10-fold in the number of cases. Three transfusional accidents notified from 1999 to 2006 were due to failure on data obtained by the questionnaires and lack of prescription of serology. According to French and British hemovigilance networks, no transfusional malaria in both countries was a consequence of missed serological positivity. Blood transfusion services, in a number of other non-endemic countries, are also...
currently considering the adoption of serology. Therefore, an adequate strategy for minimizing the risk of malaria transmitted by blood in non-endemic areas without losing donations is a combination of suitable donor selection and deferral together with screening for malarial antibodies.\(^\text{19}\)

In our case, it is important to review the criteria adopted in the clinical-epidemiological trials recommended by Brazilian guidelines, in order to ensure that the appropriate and effective donor selection guidelines are developed to implement suitable screening strategies to ensure blood safety. This review must consider including additional and relevant questions such as: place of birth, previous malaria history, residence during donor’s lifetime and travel history, even if they have traveled only to areas of low endemicity. Recent and accurate data on malaria prevalence and incidence must be available for Transfusion Centers to avoid failure in the identification of low transmission areas where symptomless cases can occur. In the Atlantic Forest of Sao Paulo State \(P. malariae\) was first reported based on molecular diagnosis after the investigation of a case of transfusion-transmitted malaria.\(^\text{21}\) This species of \(Plasmodium\) is genetically indistinguishable of \(P. brasilianum\) with respect to the CSP gene\(^\text{13}\) and also to two other nuclear genes, SSUrRNA and the protein-encoding MSP-1\(^\text{13}\), DUARTE \textit{et al.}\(^\text{13}\) tested by PCR blood samples from \textit{Alouatta guariba clamitans} from the Sao Paulo Atlantic Forest and detected 5.6% of \(P. malariae\) and 5.6% of \(P. vivax\). Since there are many summer holiday residences near the forest it is thought that malaria probably occurs as a zoonosis in this area.

The introduction of a serological test for malaria, together with good donor selection guidelines, could lead to the deferral of those donors most likely to have asymptomatic infections and thus reduce the risk of transfusion-transmitted malaria in Brazil

**RESUMO**

**Malária transfusional: relato de caso de doador assintomático infectado por Plasmodium malariae**

No Brasil a malária é endêmica na Amazônia, porém casos autóctones com baixas parasitemias ocorrem na área costeira de Mata Atlântica. De acordo com a legislação brasileira, não são obrigatórios testes para detecção de malária em doadores de sangue de áreas não-endêmicas; entretanto são excluídos por seis meses aqueles com relato de deslocamento para áreas de transmissão. Este trabalho descreve um caso de malária transfusional ocorrido em São Paulo, Brasil, em que um paciente recebeu sangue infectado, desenvolvendo a doença. Ele residia em São Paulo e não apresentava histórico de transmissão anterior ou deslocamentos para áreas endêmicas, incluindo as de baixa endemicidade, como a Mata Atlântica. A gota espessa revelou \textit{Plasmodium malariae}. Os doadores eram residentes em São Paulo e um deles (045-0) apresentou RIFI negativo com antígenos de \(P. falciparum\) e RIFI positivo com \(P. malariae\). O doador 045-0 apresentou RIFI positiva para \(P. malariae\). ELISA-P. vivax foi reagente no doador infectado (045-0) e em dois dos 44 doadores. Todos os soros foram negativos com antígeno de \(P. falciparum\). Este caso aponta a necessidade de revisão dos critérios de triagem clínico-epidemiológica para evitar casos transfusionais e também adequar as estratégias de exclusão de doadores de sangue.

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