Little is known about severe imported *Plasmodium falciparum* malaria in industrialized countries where the disease is not endemic because most studies have been case reports or have included <200 patients. To identify factors independently associated with the severity of *P. falciparum*, we conducted a retrospective study using surveillance data obtained from 21,888 *P. falciparum* severe imported malaria cases, France, 1996–2003.
RESEARCH

Each year, a growing number of persons from industrialized countries travel to developing countries. Among these millions of travelers, 20% to 70% report some illness associated with their travel and 3% report fever (7). Malaria appears to be the most common cause of fever in returned travelers. Because of its potential severity, Plasmodium falciparum infection must be considered in all febrile persons who return from an area where malaria is endemic. Depending on the country of importation and the year, 2%–16% of P. falciparum imported malaria infections are severe cases according to the World Health Organization (WHO) definition, and 10% of severe cases end with the death of the patient despite appropriate antimalarial treatment (2).

Globally, epidemiologic and pathophysiologic research studies on malaria are mainly based on disease in children <5 years of age in areas where malaria is endemic. Little is known about severe imported malaria, which primarily affects nonimmune adults. Most previous studies that have focused on severe imported malaria have been case reports or have included <200 patients (3–12). Recently, we published an analysis of risk factors associated with death in a series of 96 malaria-related deaths of 21,888 patients with imported P. falciparum malaria (2). Characteristics independently associated with death were older age, being a native of an area where malaria is nonendemic, infection occurring in eastern Africa, and absence of appropriate chemoprophylaxis. This database, including the characteristics of 832 case-patients with severe disease, provides a unique opportunity to analyze the risk factors for severe imported malaria in France. We conducted a retrospective analysis of the main features of severe imported malaria cases compared with nonsevere cases that occurred during 1996–2003 in France. Increased knowledge in this area may lead to improvement in terms of prevention and patient management.

Materials and Methods

Data Sources

 Imported malaria is not a mandatory notifiable disease in metropolitan (mainland) France. Data for this study were collected by a reporting network of 120 selected hospital laboratories and were analyzed by the French National Reference Center for Imported and Autochthonous Malaria Epidemiology (CNREPIA). Participants of the network were asked to report imported malaria cases whenever asexual forms of P. falciparum were seen by laboratory observations of a patient’s blood film. Data from the national medical informatics systems and from 2 exhaustive studies (National Quality Control Survey) suggested that these cases represented 50%–55% of the total number of imported P. falciparum malaria cases in France during the study period (13,14). A standard 57-item questionnaire, completed by clinicians and biologists for each reported case, collected basic demographic, epidemiologic, clinical, and parasitologic information (including prophylaxis and treatment).

Data Collection

The study population consisted of all P. falciparum–infected patients reported to CNREPIA during 1996–2003. We used WHO criteria for the definition of severe P. falciparum malaria as the primary outcome (10,11). The 1990 WHO definition (15) was used for cases occurring before 2000 (Table 1); the revised 2000 definition (16) was used for those occurring after 1999 (Table 2).

Table 1. Criteria used before 2000 to define severe malaria, 1990 WHO definition

<table>
<thead>
<tr>
<th>Major criteria</th>
<th>Unroutable coma</th>
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</thead>
<tbody>
<tr>
<td></td>
<td>Glasgow Coma Scale score of ≤9</td>
</tr>
<tr>
<td></td>
<td>Repeated generalized seizures</td>
</tr>
<tr>
<td></td>
<td>Circulatory collapse, systolic blood pressure &lt;80 mm Hg</td>
</tr>
<tr>
<td></td>
<td>despite adequate volume repletion</td>
</tr>
<tr>
<td></td>
<td>Pulmonary edema with presence of criteria for acute respiratory distress syndrome or acute lung injury</td>
</tr>
<tr>
<td></td>
<td>Spontaneous bleeding and/or disseminated intravascular coagulation</td>
</tr>
<tr>
<td></td>
<td>Acidemia, pH &lt;7.35, or acidosis, serum bicarbonate &lt;15 mmol/L</td>
</tr>
<tr>
<td></td>
<td>Severe anemia, hemoglobin &lt;5 g/dL</td>
</tr>
<tr>
<td></td>
<td>Renal impairment, serum creatinine &gt;265 μmol/L</td>
</tr>
<tr>
<td></td>
<td>Hypoglycemia, blood glucose, &lt;2.2 mmol/L</td>
</tr>
<tr>
<td></td>
<td>Macroscopic hemoglobinuria (if unequivocally related to malaria)</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Minor criteria</th>
<th>Impaired consciousness but rousable</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Extreme weakness</td>
</tr>
<tr>
<td></td>
<td>Temperature &gt;40°C</td>
</tr>
<tr>
<td></td>
<td>Parasitemia &gt;5%</td>
</tr>
<tr>
<td></td>
<td>Jaundice or total bilirubin &gt;50 μmol/L</td>
</tr>
</tbody>
</table>
Table 2. Criteria for severe malaria, World Health Organization definition revised in 2000

<table>
<thead>
<tr>
<th>Condition</th>
<th>Definition</th>
</tr>
</thead>
<tbody>
<tr>
<td>Extreme weakness</td>
<td>Impaired consciousness, Glasgow Coma Scale score &lt;9</td>
</tr>
<tr>
<td>Pulmonary edema</td>
<td>Repeated generalized seizures, ≥2 within 24 h</td>
</tr>
<tr>
<td>Circulatory collapse</td>
<td>Systolic blood pressure &lt;80 mm Hg despite adequate volume repletion</td>
</tr>
<tr>
<td>Spontaneous bleeding and/or disseminated intravascular coagulation</td>
<td>Jaundice or total bilirubin &gt;50 µmol/L</td>
</tr>
<tr>
<td>Severe anemia</td>
<td>Macroscopic hemoglobinuria (if unequivocally related to malaria)</td>
</tr>
<tr>
<td>Hemoglobin &lt;5 g/dL</td>
<td>Hypoglycemia, blood glucose &lt;2.2 mmol/L</td>
</tr>
<tr>
<td>Acidemia, pH &lt;7.35, or acidosis, serum bicarbonate &lt;15 mmol/L</td>
<td>Hyperlactatemia, arterial lactate &gt;5 mmol/L</td>
</tr>
<tr>
<td>Acute renal failure, urine output of &lt;400 mL/24 h and serum creatinine &gt;265 µmol/L</td>
<td>Acute renal failure, urine output of &lt;400 mL/24 h and serum creatinine &gt;265 µmol/L and ≥2 generalized seizures within 24 h.</td>
</tr>
<tr>
<td>Parasitemia ≥4%</td>
<td></td>
</tr>
</tbody>
</table>

In the 1990 WHO definition, only major criteria were used to define severe malaria. In the revised WHO definition in 2000, major and minor criteria were grouped together (except temperature criteria) to expand the definition of severe malaria. In addition, 2 criteria were changed: acute renal failure with urine output of <400 mL/24 h and serum creatinine >265 µmol/L and ≥2 generalized seizures within 24 h.

The case severity rate per 100 patients was calculated for all relevant exposure variables. Various exposure categories created for the first study (2) were used in this analysis. Patients were divided into categories: European travelers (persons born and residing in a country in Europe not endemic for malaria); European expatriates (persons born in a country in Europe where malaria is not endemic and residing in a sub-Saharan African country where malaria is endemic); African travelers (persons born in a sub-Saharan African country where malaria is endemic); African residents (persons born and residing in a sub-Saharan African country where malaria is endemic); and others. Use of malaria chemoprophylaxis was categorized as reported by patients: no use; use of inappropriate drugs (chloroquine, proguanil alone, pyrimethamine, and sulfadoxine-pyrimethamine); and use of appropriate drugs (mefloquine, atovaquone-proguanil, doxycycline, and chloroquine-proguanil) according to recommendations from the Haut Conseil de la Santé Publique (www.hcsp.fr).

**Data Analysis**

Logistic regression was used to identify factors associated with severe malaria and to estimate odds ratios (ORs) and 95% confidence intervals (CIs) for the association between exposure variables and severe cases. Dummy variables were used for variables with >2 categories. Variables with p values <0.25 were introduced in the multivariate logistic regression model. A manual backward stepwise approach was used to remove nonsignificant variables; only variables with p values <0.05 were retained in the final model. Interactions were sought by introducing interaction terms in the logistic regression model and testing for their significance at the 0.05 level. Moreover, because our data ranged from 1996 through 2003 and we used 2 WHO definitions for severe disease (15,16), we systematically tested interactions between year of diagnosis and all relevant associations; all analyses were adjusted for the year of diagnosis. Statistical analysis was performed by using Stata 10 (StataCorp LP, College Station, TX, USA).

**Results**

During the 1996–2003 study period, 27,085 malaria cases were reported to CNREPIA. Of these, 21,888 cases were P. falciparum malaria (the study population), among which 20,431 (93.3%) were uncomplicated, 832 (3.8%) were severe, 433 (2%) were asymptomatic, 160 (0.7%) showed hyperreactive malarial splenomegaly, and 32 (0.1%) were unspecified. Among the 21,888 P. falciparum cases, 96 deaths (case-fatality rate [CFR] of 0.4%) were related to malaria. The annual number of malaria cases reported by the network increased to ≈3,000 cases up to 1999–2000 and then incidence began to regularly decrease. The CFR of severe P. falciparum malaria was higher during 2000–2003 than before, reaching 15.3% in 2003 (Table 3).

Most patients were male (sex ratio M:F = 1.7), and the median age was 29.6 (range 0–96) years. African travelers were most numerous (44.6%), followed by European travelers (26.5%), African residents (12.9%), and European expatriates living in Africa (5.4%); "others" represented 10.6%. Most patients (97.5%) became infected with malaria in Africa: 59.2% in western Africa, 26.2% in central Africa, 11.2% in Madagascar and the Comoros Islands, and 0.9% in eastern Africa (Table 4; online Appendix Table). Others (2.5%) had returned mainly from French Guiana, Haiti, India, Sri Lanka, Thailand, and Indonesia. Almost one third of the patients (30.4%) stated they had taken an appropriate chemoprophylaxis, whereas more than half stated they had not. The median duration of stay was 32 days (interquartile range [IQR] 21–62 days). The median time from return to symptom onset was 6 days (IQR 1–12 days), and 11% of patients had their first symptom before returning to France. The median time from symptom to diagnosis was 3 days (IQR 1–6 days). Most patients went to a hospital first (69.4%) and, compared with Europeans, Africans were more likely to do so (73% vs. 62.7% p<0.001). At the time of diagnosis, 7.9% of patients had high-level parasitemia (>5% of parasitized erythrocytes). Seventy-five percent of
patients were hospitalized, and the median hospitalization length was 3 days (range 0–169 days). Laboratory results for 13.1% of patients revealed low platelet counts (<50 × 10^9 cells/L); 6% of patients had low hemoglobin levels (<8 g/dL).

**Description of Severe Cases**

A total of 832 patients had severe malaria according to WHO definition (51% recorded before the revision in 2000). From 1996 through 2003, the evolution of the number of severe cases was the inverse of that of the total number of imported *P. falciparum* cases. Since 2000, we observed a slow decrease in the total number of imported *P. falciparum* cases, whereas severe cases increased in number and proportion (Table 3).

Of the 832 patients who had severe malaria, 386 (46.4%) were European travelers, 98 (11.8%) European expatriates, 190 (22.8%) African travelers, 73 (8.8%) African residents, and 85 (10.2%) others. Sex ratio (M:F) was 2.3, and the median age was 38.2 years (range 0–92 years). Twenty-seven patients (3.2%) were >70 years of age, and 127 (15.3%) were <15 years of age. With a 15.3% CFR, eastern Africa accounted for 57.5% of imported malaria cases in our series. These populations, either immune or possessing residual immunity, are less likely to acquire severe cases of malaria (9), which may explain why the total proportion of severe cases in our series, fluctuating according to years from 2.5% to 5.3% (mean 3.8%), appears lower than those reported in previous studies (3–5, 7, 10–12). We identified 7 risk factors independently associated with severe imported *P. falciparum* malaria; 4 were the same as those associated with fatal imported malaria (older age, being a native from an area where malaria is nonendemic, infection occurring in East Africa, and absence of appropriate chemoprophylaxis), and 3 were new risk factors (first visit was to a general practitioner, time to diagnosis 4–12 days, and diagnosis during the fall–winter season (online Appendix Table). However, short stays and male gender were no longer predictive of severity after controlling for all the variables, including year of diagnosis. Table 5 shows the results of multivariate analysis. Independent factors associated with severe malaria were no different whether patients were African or European, and the year of diagnosis did not modify the effect of the associations.

**Factors Associated with Severe Cases**

The CFR was 3.8% during the study period and changed over time (decrease from 1996 to 2000, then an increase since; p<0.001 by standard χ^2 test or trends test). Many factors were associated with an increased risk of severe malaria in univariate analysis, including older age, male gender, European origin, infection in East Africa, short stays (<21 days), inappropriate chemoprophylaxis, initial visit to a general practitioner, time to diagnosis, and diagnosis during the fall–winter season (online Appendix Table). However, short stays and male gender were no longer predictive of severity after controlling for all the variables, including year of diagnosis. Table 5 shows the results of multivariate analysis. Independent factors associated with severe malaria were no different whether patients were African or European, and the year of diagnosis did not modify the effect of the associations.

**Discussion**

With a mean of >4,000 cases per year during the study period, France is the country reporting the highest number of imported malaria cases; >80% of cases are caused by *P. falciparum*, the species responsible for almost all severe cases and death in travelers. African travelers are particularly at risk of acquiring malaria when visiting friends and relatives. Together with African residents who declare malaria during a stay in France, African travelers accounted for 57.5% of imported malaria cases in our series. These populations, either immune or possessing residual immunity, are less likely to acquire severe cases of malaria (9), which may explain why the total proportion of severe cases in our series, fluctuating according to years from 2.5% to 5.3% (mean 3.8%), appears lower than those reported in previous studies (3–5, 7, 10–12).
Severe Imported Malaria, France

European patients (2,3). Severe malaria was particularly frequent among nonimmigrants, as previously reported in a smaller series (7–9,12). These results are consistent with the hypothesis of persistent acquired immunity, even after several years of nonexposure (12), which may partly protect African travelers from severe malaria (9,12).

Genetic factors may also partly explain the relative protection of African travelers compared with Europeans, as noted by Lewis et al. (8). Five countries, Côte d’Ivoire, Cameroon, Senegal, Mali, and the Comoros Islands, accounted for the largest numbers of malaria cases; case severity rates varied between 1.4% and 4.8%. Travelers returning from the Comoros Islands were almost exclusively migrants who were visiting friends and relatives, which may explain the particularly low proportion of severe cases observed for this country. Conversely, countries in eastern Africa such as Djibouti, Kenya, Mozambique, and Tanzania accounted for fewer cases but displayed a disproportionately high number of severe cases (10.6%–16.7%). This observation confirmed what has been reported for CFR (2).

As suggested by a preliminary study (17), a relationship between \textit{P. falciparum} drug-resistance level and severity of imported cases may exist. Because countries in eastern Africa usually harbor high proportions of drug-resistant \textit{P. falciparum} strains (20,21), further specific studies are required to assess this point. The risk of severe malaria was higher when antimalarial chemoprophylaxis was absent or inappropriate. These results are in agreement with previous observations (5,6,8) and advocate strengthening the message of prevention through the use of antimalarial chemoprophylaxis. Even when this prophylaxis fails to prevent malaria because of lack of observance, drug resistance, or pharmacologic hazards, antimalarial

\begin{table}
\centering
\caption{Distribution of imported \textit{Plasmodium falciparum} malaria cases by country of acquisition, France, 1996–2003}
\begin{tabular}{llll}
\hline
Country & \textit{P. falciparum} malaria cases & No. (%) severe cases \\
\hline
Comoros & 2,017 & 28 (1.4) \\
Cameroon & 2,707 & 76 (2.8) \\
Congo & 885 & 25 (2.8) \\
Guinea & 823 & 24 (2.9) \\
Central African Republic & 728 & 25 (3.4) \\
Côte d’Ivoire & 4,623 & 160 (3.5) \\
Togo & 604 & 21 (3.5) \\
Ghana & 194 & 7 (3.6) \\
Benin & 1,012 & 39 (3.8) \\
Mali & 2,124 & 83 (3.9) \\
Gabon & 671 & 32 (4.8) \\
Senegal & 2,234 & 108 (4.8) \\
Mauritania & 96 & 5 (5.2) \\
Burkina Faso & 740 & 41 (5.5) \\
Madagascar & 432 & 34 (7.8) \\
Niger & 152 & 12 (7.9) \\
Tanzania & 38 & 4 (10.5) \\
Guinea-Bissau & 50 & 6 (12) \\
Nigeria & 123 & 15 (12.2) \\
Mozambique & 29 & 4 (13.8) \\
Kenya & 101 & 16 (15.8) \\
Equatorial Guinea & 31 & 5 (16.1) \\
Djibouti & 12 & 2 (16.7) \\
Cape Verde & 4 & 1 (25) \\
Other & 1,458 & 59 (4.1) \\
Total & 21,888 & 832 (3.8) \\
\hline
\end{tabular}
\end{table}

\begin{table}
\centering
\caption{Factors independently associated with severe malaria among patients treated for \textit{Plasmodium falciparum} malaria in hospitals, France, 1996–2003*}
\begin{tabular}{llll}
\hline
Variables & Odds ratio (95% confidence interval) & p value \\
\hline
Age group, y & & & \\
<15 & 1 & & <0.0001 \\
16–30 & 0.9 (0.7–1.2) & & \\
31–45 & 1.06 (0.8–1.3) & & \\
46–60 & 1.8 (1.4–2.3) & & \\
>60 & 2.7 (2.0–3.6) & & \\
Origin and residence & & & \\
African travelers & 1 & & <0.0001 \\
African residents & 1.5 (1.1–1.9) & & \\
European travelers & 3.2 (2.6–3.8) & & \\
European expatriates & 3.7 (2.9–4.9) & & \\
Others & 1.9 (1.5–2.6) & & \\
Region of malaria acquisition & & & \\
Western Africa & 1 & & <0.0001 \\
Central Africa & 0.8 (0.7–0.9) & & \\
Eastern Africa & 2.6 (1.7–4.1) & & \\
Austral Africa & 1.1 (0.6–2.2) & & \\
Madagascar and Comoros Islands & 0.7 (0.5–0.9) & & \\
Others & 0.9 (0.6–1.5) & & \\
Chemoprophylaxis & & & \\
Appropriate drugs† & 1 & & 0.001 \\
No chemoprophylaxis & 1.3 (1.1–1.5) & & \\
Inappropriate drugs‡ & 1.5 (1.2–1.9) & & \\
Place of first visit & & & \\
Hospital & 1 & & <0.0001 \\
General practitioner & 1.4 (1.2–1.7) & & <0.0001 \\
Time between onset and diagnosis, d & & & \\
<1 & 1 & & <0.0001 \\
2–3 & 0.9 (0.8–1.2) & & \\
4–6 & 1.6 (1.3–1.9) & & \\
7–12 & 1.5 (1.1–1.8) & & \\
>12 & 0.7 (0.5–0.9) & & \\
Symptom onset & & & \\
After return to France & 1 & & \\
Before return to France & 1.2 (1.01–1.5) & & 0.03 \\
Season of diagnosis & & & \\
Spring–summer & 1 & & <0.0001 \\
Fall–winter & 1.3 (1.2–1.5) & & \\
\hline
\end{tabular}
\end{table}

* N = 21,888.
† Appropriate chemoprophylactic drugs were mefloquine, atovaquone-proguanil, doxycycline, and chloroquine-proguanil.
‡ According to national recommendations, inappropriate chemoprophylactic drugs were chloroquine, proguanil, pyrimethamine, and sulfadoxine-pyrimethamine.
chemoprophylaxis may confer a degree of protection against the severe form of malaria.

Management of patients who had uncomplicated malaria was not standardized among the different hospitals of our network. Depending on local procedures or on individual evaluations, patients were hospitalized, usually for the duration of their treatment, or were treated on an outpatient basis. Nevertheless, each patient with severe malaria needed to be hospitalized in an intensive care unit (exceptions to this rule depend on local medical practice). The odds of severe malaria developing were increased by 40% when the patient’s initial visit was to a general practitioner rather than to a hospital. This effect remained after controlling for time to diagnosis and suggests that this association was not due to a simple delay of diagnosis.

Time between onset of symptoms and diagnosis of 4–12 days was associated with an increased risk of severe malaria. Shorter delays of diagnosis enabled prompt treatment of malaria episodes and probably prevented their potential evolution towards severity. Diagnosis >12 days postsymptom onset usually reflected controlled parasitemia and uncomplicated malaria. These data emphasized once again the need for early diagnosis and prompt therapy. The higher severity found during the fall–winter season has been related previously to a potential mismanagement of malaria patients initially misdiagnosed with influenza (22,23). However, delay in diagnosis does not totally explain higher severity because the association between seasons and severity was not reduced after time to diagnosis was controlled for in multivariate analysis. Moreover, the influenza season in France is shorter than the fall–winter season considered in our analysis. Multivariate analysis in another study (E. Seringe et al., unpub. data) showed that malaria episodes around the end of the transmission season in areas where malaria is endemic were significantly associated with an increased risk of death (OR for each additional month away from the end of the malaria season 0.75, 95% CI 0.64–0.87; p<0.001). If one considers the geographic origin of the largest number of imported malaria cases, the end of the transmission season matched the French winter season. An additional factor, symptom onset before return to France, was marginally independently associated with severe malaria (OR 1.20, 95% CI 1.01–1.50; p = 0.03). Repatriations for medical reasons, including malaria infections, may partially explain this result.

This study has several limitations. Our surveillance network accounted for only 50%–55% of total malaria cases (uncomplicated, severe, or fatal) imported to France. Two annual exhaustive studies (National Quality Control Survey [24]) and F. Legros (unpub. data) suggest that our corresponding sites had a correct representation of patients. Thus, it is unlikely that factors associated with severity in imported malaria would be different among cases not seen in our network. From 2000 through 2003, the CFR in patients infected with P. falciparum and the percentage of severe cases have increased from 0.3% to 0.7% (p<0.0001) and from 2.5% to 4.5% (p = 0.3) respectively, whereas the CFR of severe cases remained stable (from 14.3% to 15.3%). These figures may be caused by fluctuations over time but may also be partially explained by the revision of WHO criteria for severe malaria in 2000. This revision may have led to a better categorization of severe cases, judging by the increased odds of death associated with severity after the 2000 reclassification (OR 1.84, 95% CI 1.20–2.90). However, no interaction was identified between year of diagnosis and relevant associations and all analyses were adjusted for the year of diagnosis.

Biological data were not introduced in multivariate analysis, first, because of a large number of missing data (59% and 60% of missing data in severe cases for platelet counts and hemoglobinemia, respectively) and, second, because parasitemia (10% of missing data), hemoglobin, and platelet counts are directly or indirectly part of the definition of severe malaria. Thus, parasitemia, although strongly associated with severe forms of malaria in univariate analysis, was not introduced in the final multivariate model because it is in the causal pathway between several factors and disease. For instance, delay to diagnosis leads to high parasitemia, which itself leads to severe forms of malaria. Adjusting for high parasitemia in the final model would make the relationship between delay to diagnosis and severe malaria disappear.

Treatment of severe malaria may have varied according to the physicians. Detailed French guidelines for the management of severe P. falciparum malaria were available (25). These guidelines strongly recommended a quinine-loading dose but did not recommend exchange transfusion. Intravenous artesunate was not used in France during the study period.

The clinical course of P. falciparum malaria is unpredictable and may result in severe illness and death. Although the acquisition of P. falciparum malaria among travelers to countries where malaria is endemic can never be completely avoided, our data suggest that severe malaria may largely be prevented. Pretravel health advice promoting the compliant use of antimalarial chemoprophylaxis for every traveler, with a particular focus on nonimmune travelers and elderly persons, is essential. In addition, increased vigilance of travelers in reporting symptoms and of physicians in providing prompt diagnosis and treatment is required to reduce any delay in patient management.

Acknowledgments

We thank all staff, clinicians, and biologists at each French National Reference Center for Imported and Autochthonous
Malaria Epidemiology and the French National Reference Center for Imported Malaria Network corresponding site.

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