In Africa, control programs that target primarily *Plasmodium falciparum* are inadequate for eliminating malaria. To learn more about prevalence and genetic variability of *P. malariae* in Africa, we examined blood samples from 663 asymptomatic and 245 symptomatic persons from western Kenya during June–August of 2014 and 2015. *P. malariae* accounted for 5.3% (35/663) of asymptomatic infections and 3.3% (8/245) of clinical cases. Among asymptomatic persons, 71% (32/45) of *P. malariae* infections detected by PCR were undetected by microscopy. The low sensitivity of microscopy probably results from the significantly lower parasitemia of *P. malariae*. Analyses of *P. malariae* circumsporozoite protein gene sequences revealed high genetic diversity among *P. malariae* in Africa, but no clear differentiation among geographic populations was observed. Our findings suggest that *P. malariae* should be included in the malaria elimination strategy in Africa and highlight the need for sensitive and field-applicable methods to identify *P. malariae* in malaria-endemic areas.

Over the past decade, malaria control strategies in Africa have reduced the number of malaria cases and deaths. Nevertheless, non-*Plasmodium falciparum* malaria still presents a major challenge for malaria elimination (1,2). Global malaria elimination programs focus primarily on *P. falciparum*. Recent research efforts and control programs have drawn resources to *P. vivax*. Compared with the distribution of *P. falciparum* and *P. vivax*, the distribution of *P. malariae* is relatively sparse and variable. *P. malariae* is endemic to West and Central Africa (3), South America (15), Asia (16,17), and the western Pacific region (18,19). Knowledge of genetic variation among isolates from these geographic areas is still lacking. One study indicated a remarkably low level of sequence diversity at the *msp1* locus in *P. malariae* from Brazil (20). Similarly, the lack of variation at the *dhfr* and *dhps* loci has been shown for *P. malariae* from Asia and the western Pacific region (21,22). These findings suggested that antimalarial drugs might be imposing selective pressure on the genetic diversity of *P. malariae*. The circumsporozoite protein (*csp*) gene, which is known to be critical for plasmodia sporozoite motility and hepatocyte invasion (23), has been shown to be variable in length and is a sequence of the tandemly

**Plasmodium malariae** Prevalence and *csp* Gene Diversity, Kenya, 2014 and 2015

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isolates 7G8 (MR4-MRA-926) and 30, and the gene region and used together with the Pakchong (MR4-MRA-32) P. ovale (MR4-MRA-180). As to determine the blood smears were prepared for microscopic examination of error in parasite prevalence with 0.05 type I error. Thick and thin symptoms and were determined to be positive for These patients had fever or malaria-related signs or symptoms. A slide was considered negative when no parasites were observed after counting >100 microscopic fields. At the time of sample collection, all slides were read by 2 microscopists. If counts were discordant, the slides were examined by a third microscopist. The density of parasitemia was expressed as the number of asexual parasites per microliter of blood, assuming a leukocyte count of 8,000 cells/µL, according to World Health Organization guidelines.

We extracted parasite DNA from half of a dried blood spot by using the Saponin/Chlex method (32). The final extracted volume was 200 µL. For all samples, nested amplification of the 18S rRNA gene region of plasmodia (P. falciparum, P. vivax, P. malariae, and P. ovale) was used for parasite detection and species identification. As positive controls for all amplifications, we used DNA from P. falciparum isolates 7G8 (MR4-MRA-926) and HB3 (MR4-MRA-155), P. vivax Pakchong (MR4-MRA-342G) and Nicaragua (MR4-MRA-340G), P. malariae (MR4-MRA-179), and P. ovale (MR4-MRA-180). As negative controls, we used water and noninfected samples to ensure lack of contamination. Reaction was performed in a Bio-Rad MyCycler thermal cycler according to the published protocol (33) (details in online Technical Appendix 1, https://wwwnc.cdc.gov/EID/article/23/4/16-1245-Techapp1.pdf).

In addition, the amount of parasite DNA was estimated by using the SYBR Green (Thermo Scientific, Foster City, CA, USA) qPCR detection method with Plasmodium species-specific primers that targeted the 18S rRNA genes (34,35). Reactions were performed in a CFX96 Touch Real-Time PCR Detection System (Bio-Rad, Foster City, CA, USA). To confirm specific amplifications of the target sequence, we performed melting curve analyses for each amplified sample. To measure reproducibility of the cycle threshold (Ct), we calculated the mean value and standard deviations from triplicates in 2 independent assays. The parasite gene copy number in a sample was quantified by using the equation (30) GCN sample = e$^{\Delta Ct}$, where GCN stands for gene copy number; ΔCt, the difference in Ct between the negative control and the sample; e, exponential function; and E, amplification efficiency (online Technical Appendix 1).

**CSP Sequencing and Phylogenetic Analyses**
Four internal primers were designed specifically on the P. malariae csp gene region and used together with the repeated peptide units in P. falciparum (24,25), P. vivax (26,27), and P. malariae isolates from Central Africa (28). The vast antigenic variation observed in P. falciparum as a result of immune selection pressure can influence the capacity of mosquito transmission and the effectiveness of malaria vaccine (29). In this study, we sought to determine the prevalence of infection and age distribution of persons with asymptomatic and symptomatic P. malariae infection in western Kenya, the genetic affinity between P. malariae isolates from East Africa and other regions, and the level of csp gene diversity among P. malariae and the significance of this diversity.

Scientific and ethical clearance was given by the institutional scientific and ethical review boards of the Kenya Medical Research Institute and the University of California Irvine. Written informed consent/assent for study participation was obtained from all consenting heads of households, parents/guardians (for minors <18 years of age), and each person who was willing to participate in the study.

**Materials and Methods**

**Study Areas and Participants**
During June–August of 2014 and 2015, blood samples were collected from persons in 4 villages at the Lake Victoria basin (elevation =1,000 m) of western Kenya (Figure 1). These villages represent parts of the Lake Victoria area previously shown by nested and quantitative PCR (qPCR) methods to have high, stable rates of malaria transmission and prevalence (10%–40%) among children 5–14 years of age (30,31).

Community samples were collected from nonfebrile schoolchildren in 7 public primary schools (70–100 children/school, 2 schools/village except Kombewa). An equal number of boys and girls 6–15 years of age were randomly selected from each school. To determine P. malariae prevalence in the adult population, we randomly selected 63 persons (32 male and 31 female) >15 years of age from 18 households in Kombewa. We examined a total of 663 persons (32 male and 31 female) >15 years of age from 18 households in Kombewa. We considered fever or malaria-related symptoms.

Clinical samples were collected from 113 male and 132 female patients, <1 to 76 years of age, in 3 district hospitals. This sample size provided an estimation of 4% margin of error in parasite prevalence with 0.05 type I error. These patients had fever or malaria-related signs or symptoms and were determined to be positive for Plasmodium spp. by microscopy at the time of sampling. Thick and thin blood smears were prepared for microscopic examination to determine the Plasmodium species, and ≈50 µL blood was blotted onto Whatman 3MM filter (Sigma Aldrich, St. Louis, MO, USA) papers. Filter papers were air dried and stored in zip-sealed plastic bags with silica gel absorbent at room temperature until DNA extraction.

**Microscopy and PCR of Plasmodium spp.**
We examined slides under microscopes at 100× magnification and counted the number of parasites per 200 leukocytes. A slide was considered negative when no parasites were observed after counting >100 microscopic fields. At the time of sample collection, all slides were read by 2 microscopists. If counts were discordant, the slides were examined by a third microscopist. The density of parasitemia was expressed as the number of asexual parasites per microliter of blood, assuming a leukocyte count of 8,000 cells/µL, according to World Health Organization guidelines.

We extracted parasite DNA from half of a dried blood spot by using the Saponin/Chlex method (32). The final extracted volume was 200 µL. For all samples, nested amplification of the 18S rRNA gene region of plasmodia (P. falciparum, P. vivax, P. malariae, and P. ovale) was used for parasite detection and species identification. As positive controls for all amplifications, we used DNA from P. falciparum isolates 7G8 (MR4-MRA-926) and HB3 (MR4-MRA-155), P. vivax Pakchong (MR4-MRA-342G) and Nicaragua (MR4-MRA-340G), P. malariae (MR4-MRA-179), and P. ovale (MR4-MRA-180). As negative controls, we used water and noninfected samples to ensure lack of contamination. Reaction was performed in a Bio-Rad MyCycler thermal cycler according to the published protocol (33) (details in online Technical Appendix 1, https://wwwnc.cdc.gov/EID/article/23/4/16-1245-Techapp1.pdf).

In addition, the amount of parasite DNA was estimated by using the SYBR Green (Thermo Scientific, Foster City, CA, USA) qPCR detection method with Plasmodium species-specific primers that targeted the 18S rRNA genes (34,35). Reactions were performed in a CFX96 Touch Real-Time PCR Detection System (Bio-Rad, Foster City, CA, USA). To confirm specific amplifications of the target sequence, we performed melting curve analyses for each amplified sample. To measure reproducibility of the cycle threshold (Ct), we calculated the mean value and standard deviations from triplicates in 2 independent assays. The parasite gene copy number in a sample was quantified by using the equation (30) GCN sample = e$^{\Delta Ct}$, where GCN stands for gene copy number; ΔCt, the difference in Ct between the negative control and the sample; e, exponential function; and E, amplification efficiency (online Technical Appendix 1).
published primers (28; online Technical Appendix 1 Table) to unambiguously amplify the 3 segments, the N terminal, the central repeat, and the C-terminal regions of the csp gene. A total of 37 P. malariae isolates were amplified and sequenced. All resulted sequences were verified by comparing them with those in the GenBank database by using BLAST (https://blast.ncbi.nlm.nih.gov/Blast.cgi). Sequences were translated into protein sequences and analyzed together with all csp protein sequences available in GenBank of P. malariae from East Africa (Kenya and Uganda), West Africa (Cameroon), Central Africa (Côte d’Ivoire), and South America (Venezuela) and of P. brasilianum from South America (Brazil and Venezuela). It is noteworthy that although P. malariae and P. brasilianum coexist in Brazil, no csp sequence for P. malariae is available. Because of the potential for alignment errors associated with gaps in the nucleotide sequences, we used translated amino acid sequences with unambiguous indels in phylogenetic analyses. Sequence diversity, including measures of evolutionary distances and average pairwise divergence, were estimated and compared among geographic regions (online Technical Appendix 1).

Statistical Analyses
A 1-tailed t-test was used to test for the significance of differences in parasite gene copy number between P. malariae from symptomatic and asymptomatic patients and between P. malariae and P. falciparum in co-infected samples. In addition, we calculated the Pearson correlation coefficient ($r^2$) for parasite gene copy number and age by using R (https://www.r-project.org/).

Results

P. malariae Prevalence and Patient Age Distribution
Among the 663 samples from asymptomatic persons, P. malariae was detected by PCR in 35 (5.3% prevalence). Among these, 29 were mixed infections (with P. falciparum) and 6 were P. malariae monoinfections (Figure 1; Table 1). P. malariae was found to be most prevalent in Kombewa (14.3%, 19/133 cases), followed by Kendu Bay (5.3%, 8/150 cases). Prevalence of P. falciparum prevalence was relatively high at these 2 sites (44% and 59%, respectively; Table 1). In Kombewa, 13 of 19 P. malariae cases were detected in younger persons (<15 years of age), which was significantly higher than the number of cases detected in older persons (6 cases, $p = 0.04$; Table 1). Although such a comparison between age groups cannot be made for the other sites, a similar pattern was observed for symptomatic patients.

Among the 245 samples from symptomatic patients, 8 (3.3%) P. malariae cases were detected; 6 were mixed
infections with *P. falciparum* and 2 were *P. malariae* monoinfestations (Table 2). When the samples were stratified by patient age, all *P. malariae* infections in symptomatic persons were in infants or very young children of <5 years of age (8/135, 5.9% infection rate). Although *P. falciparum* infection was highest among patients >5 to ≤15 years of age, no *P. malariae* was detected in persons in this and older age groups despite smaller samples in these groups. No significant difference was detected between male and female patients.

**Comparisons of Diagnostic Approaches and Parasitemia**

Compared with microscopy, nested PCR revealed a significantly higher number of *P. malariae* infections in the community (Table 3). All samples that were *P. malariae* positive by microscopy were identified as positive by PCR and qPCR. Across the study sites, nested PCR–based prevalence ranged from 0 to 12.2% (average 4.8%), >2-fold higher than by microscopy (0 to 3.8%, average 1.9%; Table 3). The discrepancy between the 2 methods was also reflected by the difference in *P. falciparum* prevalence; 10% more positive infections were detected by nested PCR than by microscopy. Nevertheless, such a discrepancy was not as substantial as that for *P. malariae*.

Although the number of *P. malariae*–positive clinical samples detected in this study was low, these samples indicated an overlapping range of parasite gene copy number (geometric mean 6.4×10^3/µL, range 4.3×10^1 to 1.2×10^7/µL; Figure 2) with that of the samples from asymptomatic persons (geometric mean 4.8×10^3/µL, range 0.5×10^3 to 9.4×10^7/µL) without differing significantly (p>0.05). Similar results were observed in the level of *P. malariae* parasitemia, for which samples from symptomatic and asymptomatic persons did not differ significantly (Figure 2). Parasite gene copy number and *P. malariae* parasitemia were significantly positively correlated with each other (r^2 = 0.77, p<0.01; online Technical Appendix 1 Figure 1).

Parasite gene copy number and parasitemia for *P. falciparum* were generally higher than those for *P. malariae* (Figure 3, panel A). Among the 35 mixed infections, 28 (80%) gene copy numbers were higher for *P. falciparum* than for *P. malariae* (online Technical Appendix 1 Figure 2). Among these samples overall, the amount of *P. falciparum*

---

### Table 1. Prevalence of *Plasmodium malariae* and *P. falciparum* among asymptomatic persons in the community, Kenya, June–August 2014 and 2015*

<table>
<thead>
<tr>
<th>Site, patient age, y</th>
<th>No. tested</th>
<th>Total</th>
<th><em>P. malariae</em></th>
<th><em>P. falciparum</em></th>
<th>Mixed†</th>
</tr>
</thead>
<tbody>
<tr>
<td>Kombewa</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>≤15</td>
<td>63</td>
<td>41 (65.1)</td>
<td>0</td>
<td>28 (44.4)</td>
<td>13 (20.6)</td>
</tr>
<tr>
<td>&gt;15</td>
<td>70</td>
<td>35 (50)</td>
<td>2 (2.9)</td>
<td>29 (41.4)</td>
<td>4 (5.7)</td>
</tr>
<tr>
<td>Chulaimbo</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>≤15</td>
<td>190</td>
<td>76 (40)</td>
<td>2 (1.1)</td>
<td>71 (37.4)</td>
<td>3 (1.6)</td>
</tr>
<tr>
<td>&gt;15</td>
<td>150</td>
<td>97 (64.7)</td>
<td>0</td>
<td>89 (59.3)</td>
<td>8 (6)</td>
</tr>
<tr>
<td>Kendu Bay</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>≤15</td>
<td>190</td>
<td>57 (30)</td>
<td>2 (1.1)</td>
<td>54 (28.4)</td>
<td>1 (0.5)</td>
</tr>
<tr>
<td>Port Victoria</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>≤15</td>
<td>663</td>
<td>306 (46.2)</td>
<td>6 (0.9)</td>
<td>271 (40.9)</td>
<td>29 (4.4)</td>
</tr>
</tbody>
</table>

*According to nested PCR of the 18S rRNA gene.
†*P. malariae* and *P. falciparum.*

---

### Table 2. Prevalence of *Plasmodium malariae* and *P. falciparum* among symptomatic persons, Kenya, June–August 2014 and 2015*

<table>
<thead>
<tr>
<th>Site, patient age, y</th>
<th>No. tested</th>
<th>Total</th>
<th><em>P. malariae</em></th>
<th><em>P. falciparum</em></th>
<th>Mixed†</th>
</tr>
</thead>
<tbody>
<tr>
<td>Chulaimbo</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>≤5</td>
<td>27</td>
<td>18 (66.7)</td>
<td>2 (7.4)</td>
<td>15 (55.6)</td>
<td>0</td>
</tr>
<tr>
<td>&gt;5 to ≤15</td>
<td>4</td>
<td>3 (75)</td>
<td>0</td>
<td>3 (75)</td>
<td>0</td>
</tr>
<tr>
<td>&gt;15</td>
<td>13</td>
<td>3 (23.1)</td>
<td>0</td>
<td>3 (23.1)</td>
<td>0</td>
</tr>
<tr>
<td>Kendu Bay</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>≤5</td>
<td>44</td>
<td>38 (86.4)</td>
<td>0</td>
<td>35 (79.5)</td>
<td>3 (6.8)</td>
</tr>
<tr>
<td>&gt;5 to ≤15</td>
<td>34</td>
<td>31 (91.2)</td>
<td>0</td>
<td>31 (91.2)</td>
<td>0</td>
</tr>
<tr>
<td>&gt;15</td>
<td>24</td>
<td>23 (95.8)</td>
<td>0</td>
<td>23 (95.8)</td>
<td>0</td>
</tr>
<tr>
<td>Port Victoria</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>≤5</td>
<td>64</td>
<td>54 (84.4)</td>
<td>0</td>
<td>51 (79.7)</td>
<td>3 (4.7)</td>
</tr>
<tr>
<td>&gt;5 to ≤15</td>
<td>22</td>
<td>20 (90.9)</td>
<td>0</td>
<td>20 (90.9)</td>
<td>0</td>
</tr>
<tr>
<td>&gt;15</td>
<td>13</td>
<td>9 (69.2)</td>
<td>0</td>
<td>9 (69.2)</td>
<td>0</td>
</tr>
<tr>
<td>Total</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>≤5</td>
<td>135</td>
<td>110 (81.5)</td>
<td>2 (1.5)</td>
<td>101 (74.8)</td>
<td>6 (4.4)</td>
</tr>
<tr>
<td>&gt;5 to ≤15</td>
<td>60</td>
<td>54 (90)</td>
<td>0</td>
<td>54 (90)</td>
<td>0</td>
</tr>
<tr>
<td>&gt;15</td>
<td>50</td>
<td>35 (70)</td>
<td>0</td>
<td>35 (70)</td>
<td>0</td>
</tr>
</tbody>
</table>

*According to nested PCR of the 18S rRNA gene.
†*P. malariae* and *P. falciparum.*
DNA (geometric mean $1.6 \times 10^{2}$ /µL, range $1 \times 10^{1}$ to $5.5 \times 10^{3}$ /µL) was significantly higher than the amount of *P. malariae* DNA (geometric mean $4.7 \times 10^{1}$ /µL, range $0.4 \times 10^{1}$ to $1.1 \times 10^{3}$ /µL; *p* = 0.003), consistent with the difference in parasitemia according to microscopy (*P. malariae* geometric mean $3.2 \times 10^{2}$ parasites/µL vs. *P. falciparum* geometric mean $1.1 \times 10^{3}$ parasites/µL; online Technical Appendix 1 Figure 2).

When all *P. malariae* samples were pooled, the parasite gene copy number did not correlate significantly with patient age ($r^2 = 0.07$; online Technical Appendix 1 Figure 3). Neither *P. malariae* prevalence rate nor parasite gene copy number differed significantly according to patient sex.

### Genetic Relatedness and csp Divergence of *P. malariae*

The *csp* alignment comprised 530 aa, of which 34 (6.4%) were polymorphic among the studied parasites of different taxa (online Technical Appendix 2, https://wwwnc.cdc.gov/EID/article/23/4/16-1245-Techapp2.pdf). To avoid polymorphism caused by PCR error, we sequenced each isolate at least twice in both directions. Substantial length variation was observed in the central repeat region, where the number of NAAG (the repeat codon unit in which n denotes the number of repeats) in *P. malariae* ranged from 49 to 85 units. These tandem repeats could be rapidly evolving.

![Figure 2](image1.png)  
Figure 2. Parasite gene copy numbers (per microliter) detected by SYBR Green (Thermo Scientific, Foster City, CA, USA) quantitative PCR and parasitemia (parasites per microliter) determined by microscopy of *Plasmodium malariae* samples from asymptomatic and symptomatic persons. Median, first quartile, and fourth quartile of the data are shown for each sample category (horizontal lines). No significant difference was observed between asymptomatic and symptomatic persons in terms of *P. malariae* parasite gene copy number and parasitemia. Squares represent samples with gene copy number measured by quantitative PCR; circles, samples with parasitemia estimated by microscopy; closed squares and circles, *P. malariae* samples from asymptomatic persons; open squares and circles, *P. malariae* samples from symptomatic patients. NS, not significant.

![Figure 3](image2.png)  
Figure 3. *Plasmodium malariae* and *P. falciparum* parasite gene copy numbers (per microliter) and parasitemia (parasites per microliter) in co-infected samples. Median, first quartile, and fourth quartile of the data are shown for each sample category (horizontal lines). Parasite gene copy number and parasitemia were lower in *P. malariae*-positive than in *P. falciparum*-positive samples. Squares represent samples with gene copy number measured by quantitative PCR; circles, samples with parasitemia estimated by microscopy; red, *P. malariae* samples; blue, *P. falciparum* samples.

---

### Table 3. Methods used to diagnose *Plasmodium* infections in asymptomatic populations, Kenya, June–August 2014 and 2015*

<table>
<thead>
<tr>
<th>Site, method</th>
<th>No. tested</th>
<th>Total</th>
<th><em>P. falciparum</em></th>
<th><em>P. vivax</em></th>
<th><em>P. malariae</em></th>
<th><em>P. ovale</em></th>
<th><em>P. falciparum/malariae</em></th>
</tr>
</thead>
<tbody>
<tr>
<td>Kombewa</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Microscopy</td>
<td>133</td>
<td>54 (41.2)</td>
<td>49 (37.4)</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>5 (3.8)</td>
</tr>
<tr>
<td>PCR</td>
<td>133</td>
<td>70 (53.4)</td>
<td>54 (41.2)</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>16 (12.2)</td>
</tr>
<tr>
<td>Chulaimbo</td>
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<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Microscopy</td>
<td>190</td>
<td>46 (24.2)</td>
<td>42 (22.1)</td>
<td>0</td>
<td>1 (0.5)</td>
<td>0</td>
<td>3 (1.6)</td>
</tr>
<tr>
<td>PCR</td>
<td>190</td>
<td>76 (40.1)</td>
<td>71 (37.4)</td>
<td>0</td>
<td>2 (1.1)</td>
<td>0</td>
<td>3 (1.6)</td>
</tr>
<tr>
<td>Kendu Bay</td>
<td></td>
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</tr>
<tr>
<td>Microscopy</td>
<td>150</td>
<td>78 (52)</td>
<td>75 (50)</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>3 (2)</td>
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<tr>
<td>PCR</td>
<td>150</td>
<td>97 (64.6)</td>
<td>89 (59.3)</td>
<td>0</td>
<td>0</td>
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<td>8 (5.3)</td>
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</tr>
<tr>
<td>Microscopy</td>
<td>190</td>
<td>36 (18.5)</td>
<td>35 (18.5)</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>1 (0.5)</td>
</tr>
<tr>
<td>PCR</td>
<td>190</td>
<td>57 (30)</td>
<td>54 (28.4)</td>
<td>0</td>
<td>2 (1.1)</td>
<td>0</td>
<td>1 (0.5)</td>
</tr>
<tr>
<td>All sites</td>
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<td></td>
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<td>201 (30.4)</td>
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<td>1 (0.2)</td>
<td>0</td>
<td>12 (1.8)</td>
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<td>268 (40.5)</td>
<td>0</td>
<td>4 (0.6)</td>
<td>0</td>
<td>28 (4.2)</td>
</tr>
</tbody>
</table>
through a different mechanism and may influence genetic relationships among the samples. To examine such effect, we constructed phylogenetic trees with 2 sets of data: the entire sequence (530 aa) and partial sequences without the central repeat region (225 aa).

Maximum-likelihood analyses of the entire csp gene showed a clear distinction between isolates from South America and those from the other geographic regions (Figure 4, panel A). P. brasilianum and P. malariae from Venezuela formed a monophyletic group (bootstrap >95%) closely associated with P. brasilianum from Brazil. Sequences of P. malariae from Venezuela were almost identical to those of P. brasilianum from the same area. Closely related to the clade from South America was a large monophyletic group that contained P. malariae from East, Central, and West Africa and from China (bootstrap >90%). The isolates from these regions were divided into 2 subclades: I and II (Figure 4, panel A). Subclade I comprised a mix of P. malariae isolates from Kenya, Cameroon, and Côte d’Ivoire. Subclade II comprised a mix of P. malariae isolates from Kenya, Cameroon, Uganda, and China. Sequences without the central repeat region indicated consistently the distinctiveness between P. brasilianum from Brazil and P. malariae, but the P. malariae samples from different geographic regions were poorly resolved (Figure 4, panel B). The P. malariae isolate from China was nested within the African subclade, suggestive of an African origin (Figure 4, panel C). No clear microgeographic structure was detected, although sample size at the population level was small.

Among the 3 geographic regions, the level of csp sequence divergence in P. malariae was higher in isolates from East Africa than from West Africa, as reflected by a higher number of polymorphic sites and a greater extent of csp length variation despite difference in sample size (Figure 5, panels A and B). These variations were located at the 3’ N terminal through the central repeat region, where the largest degree of mismatch was observed (Figure 5, panel B). To the contrary, the level of sequence polymorphism was lowest in isolates from South America (Figure 5, panel A), but the greatest range of difference in tandem repeat units where remarkable mismatch was observed was toward the end of the central region. Despite the small sample size, the number of tandem repeats was generally lower in P. brasilianum than P. malariae (Figure 5, panel C).

Discussion

In Kenya, areas along the shoreline of Lake Victoria and coastal regions are malaria hot spots, where intense and stable
plasmodia transmission occurs throughout the year (31). For achieving the ultimate goal of eliminating malaria in Kenya, existing control programs that primarily target *P. falciparum* are inadequate. The use of rapid diagnostic tests or microscopy as first-line diagnostic methods can lead to gross underestimation of the actual prevalence of *P. malariae* (4–6). Our findings indicated that *P. malariae* accounted for ≈3% of clinical cases and ≈5% of asymptomatic infections in this malaria-endemic region. The prevalence of asymptomatic *P. malariae* infections was comparable to that recently reported for nearby islands of Lake Victoria (1.7%–3.96%) on the basis of PCR (36,37). These asymptomatic *P. malariae* infections are concerning because they are parasite reservoirs that can sustain long-term transmission. For instance, in the Colombian Amazon region, *P. malariae* was thought to account for ≤1% of all malaria infections (38,39); however, a recent study revealed that 43.6% (294/675) of clinical cases were caused by *P. malariae* (10) and suggested that these parasites have been circulating in the community undetected. Underestimation or lack of awareness of its occurrence could thus lead to increased transmission. The infectiousness of *P. malariae* for *Anopheles* mosquitoes in malaria-endemic areas remains unclear and merits further investigation.

We found that *P. malariae* infections were more common among infants and children than adults. A similar pattern has been found for Senegal, West Africa, where 91% (265/290 cases) of clinical *P. malariae* cases occurred in children <15 years of age and the mean incidence density was highest for those 5–9 years of age (3). These findings indicate that children are vulnerable to *P. malariae* infection and contrast with those reported for Papua, Indonesia, where *P. malariae* infection was higher among older (median 21 years of age) than younger persons (9). It is possible that our study sites in western Kenya, as well as in West Africa, are high-transmission areas where *P. falciparum* malaria prevalence can be ≈60% during the rainy season (30,40). Cumulative exposure to the parasites over time may enable gradual acquisition of immunity in adults. Nevertheless, our community samples were mostly obtained from schoolchildren 6–15 years of age. Underrepresentation of adult populations may underestimate the overall malaria prevalence in the study area. Although young children are more vulnerable to *P. malariae* infections, the level of *P. malariae* parasitemia does not seem to be associated with age. Chronic nephrotic syndromes attributed to *P. malariae* have been reported (41,42) and shown to be associated with significant illness from anemia in young children (8,9). However, the lack of hematologic data from our study participants limits further investigation.

Our data indicate that ≈50% of *P. malariae*–positive samples detected by PCR were undetected by microscopy. Such a low sensitivity of microscopy could be attributed to a significantly lower *P. malariae* than *P. falciparum* parasitemia, according to qPCRs. Because most *P. malariae*–positive samples had mixed infections, microscopists could have recorded only the dominant *P. falciparum* and overlooked the sparse *P. malariae* trophozoites. Also, the ring forms of *P. falciparum* and *P. malariae* are morphologically more similar to each other than to *P. vivax* and *P. ovale* (43). Misdiagnosis of parasite species by microscopy is possible (8).

### Figure 5

Comparison of circumsporozoite protein (*csp*) gene sequence divergence among *Plasmodium* isolates from different geographic regions. A) Pairwise genetic distance plot of all amino acid positions of the *csp* gene. The matrix-normalized distances based on the standard point accepted mutation (Dayhoff–PAM) model that account for the probability of change from 1 amino acid to another were calculated. Samples were analyzed as a whole and partitioned by geographic regions as indicated by colors. B) Dot plot showing matching scores, a proxy of sequence similarity, between pairwise samples calculated based on the standard Dayhoff–PAM matrix. The greatest mismatch was detected at amino acid positions 110–310, representing the 3′ N terminal through the central repeat regions. C) Variation in the number of tandem repeats in the central region of the *csp* gene. The greatest length variation was observed in the isolates from South America despite the fact that both *P. malariae* (PM) and *P. brasilianum* (PB) were included. *P. malariae* from East Africa was more variable in the number of repeats than isolates from Central/West Africa, despite difference in sample size. Median, first quartile, and fourth quartile of the data are shown for each sample category (horizontal lines). Red represents samples from South America; yellow, Central/West Africa; blue, East Africa. Circles represent *P. brasilianum*; squares, *P. malariae*. 

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In Africa, the standard treatment for *P. malariae* monoinfection is chloroquine, and for *P. falciparum* and mixed plasmodial infections it is artemisinin combination therapy (31). The combination treatment regime should cure *P. malariae* infections even in cases of misdiagnosis. However, *P. malariae* increases production of *P. falciparum* gametocytes in mixed infections, and these gametocytes can persist without proper antimalarial treatment or monitoring (44). Therefore, we highlight the need for sensitive methods to improve *P. malariae* diagnosis and provide accurate epidemiologic data for specific and effective management guidelines. Although PCR is a better diagnostic method, it uses a relatively small amount of blood from filter papers and could still underestimate *P. malariae* infections in samples with exceptionally low levels of parasitemia. More accurate prevalence data may be obtained from ultrasensitive PCR that targets multicopy regions of the parasite genome (45) or reverse transcription PCR of parasite RNA extracted from whole blood (46).

Sequences of the *csp* gene were shown to be highly polymorphic among *P. malariae* isolates from western Kenya. The most polymorphic region was in the central repeat region, where mutations and length differences were detected (24,28). Among the isolates from different geographic areas, *P. malariae* from East and Central/West Africa were genetically closely related and exhibited a comparable level of sequence variation. This variation could be attributed to positive selection, frequent recombination, and gene flow among the parasites, as follows. First, compared with *msp1*, *dhfr*, and *dhps* of *P. malariae* (20–22), the *csp* gene revealed a remarkably higher level of sequence diversity. It is possible that selection of *csp* genetic variants may confer immunogenic advantages to the pathogen during host invasion (28,47). Second, intense transmission and large vector populations in our study area might enhance frequent heterologous recombination of the parasite genome during reproduction in the mosquitoes and increase genetic diversity within populations (24,25). Third, recurrent gene flow between the parasite populations across countries, via human migration or dispersal of vector mosquitoes, promotes the spread of these genetic variants, leading to a lack of differentiation according to geographic region. Future study using other variable markers, such as microsatellites, on expanded population samples could validate our findings.

In summary, underestimation of the actual prevalence of asymptomatic infections hinders progress toward malaria elimination in Africa. The low parasitemia of *P. malariae* infections influences diagnostic sensitivity by microscopy. A more sensitive tool is needed to identify asymptomatic *P. malariae* and to improve control strategies, particularly among infants and children who are vulnerable to *P. malariae* infection.

**Acknowledgments**

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Dr. Lo is a researcher focused on molecular epidemiology and evolution of pathogens. She is interested in exploring the effects of host–parasite interactions on parasite genomic and genetic structure.

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Technical Appendix 1

Detailed description of PCR-based diagnostic assays and phylogenetic analyses of csp sequences

Microscopy

Slides were examined under microscopes 100× objective. Parasites were counted against 200 leukocytes. A slide was considered negative when no parasites were observed after counting over 100 microscopic fields. All slides were read in duplicate by two microscopists at the time of sample collection. In the case of discordance, the slides were examined by a third microscopist. The density of parasitemia was expressed as the number of asexual parasite per microliter of blood, assuming a leukocyte count of 8000 cells per microliter according to the WHO guidelines.

Nested PCR assay of Plasmodium species

Parasite DNA was extracted from half of a 50ul-dried blood spot by the Saponin/Chelex method (1). The final extracted volume was 200μl. A nested amplification of the 18S rRNA gene region of Plasmodium (P. falciparum, P. vivax, P. malariae and P. ovale) was used for parasite detection and species identification in all samples. DNA from P. falciparum isolates 7G8 (MR4-MRA-926) and HB3 (MR4-MRA-155), P. vivax Pakchong (MR4-MRA-342G) and Nicaragua (MR4-MRA-340G), P. malariae (MR4-MRA-179), as well as P. ovale (MR4-MRA-180) were used as positive controls in all amplifications. Water and uninfected samples were used as negative controls to ensure lack of contamination. Amplification was conducted in a 20ul reaction mixture containing 2ul of genomic DNA, 10ul of 2×DreamTaq™ Green PCR Master Mix (Fermentas), and 0.3uM primers. Reaction was performed in a BIORAD MyCycler thermal cycler following the published protocol (2). The amplified products were resolved electrophoretically on a 2% agarose gel in 0.5×Tris-borate (TBE) buffer and visualized under UV light.
Quantitative real-time PCR assay of Plasmodium species

Parasite DNA amount was estimated using the SYBR Green qPCR detection method with *Plasmodium* species-specific primers that targeted the 18S rRNA genes (3–5). Amplification was conducted in a 20µL reaction mixture containing 2µL of genomic DNA, 10µL 2×SYBR Green qPCR Master Mix (Thermo Scientific, USA), and 0.5µM primer. Reaction was performed in CFX96 Touch™ Real-Time PCR Detection System (Bio-Rad), with an initial denaturation at 95°C for 3 min, followed by 45 cycles at 94°C for 30 sec, 55°C for 30 sec, and 68°C for 1 min with a final 95°C for 10 sec. This was then followed by a melting curve step of temperature that ranged from 65°C to 95°C with 0.5°C increment to determine the melting temperature of each amplified product. Melting curve analyses were performed for each amplified sample to confirm specific amplifications of the target sequence. For the measure of reproducibility of the threshold cycle number (*C*ₜ), the mean value and standard deviations were calculated from triplicates in two independent assays. A cut-off threshold of 0.02 fluorescence units that robustly represented the threshold cycle at the log-linear phase of the amplification and above the background noise was set to determine *C*ₜ value for each assay. Samples yielding *C*ₜ values higher than 40 (as indicated in the negative controls) were considered negative for *Plasmodium* species. The parasite gene copy number (GCN) in a sample was quantified based on the threshold cycle using the follow equation (6): \[ \text{GCN}_{\text{sample}} = e^{[\text{E} \times \Delta \text{Ct}_{\text{sample}}]} \], where GCN stands for gene copy number, \( \Delta \text{Ct} \) for the difference in threshold cycle between the negative control and the sample, and E for amplification efficiency. The amplification efficiency of primers was assessed on 10-fold serial dilutions ranging from 10⁵ to 10¹ copies/µl of the control plasmids. DNA from *P. falciparum* isolates 7G8 (MR4-MRA-926) and HB3 (MR4-MRA-155), and *P. malariae* (MR4-MRA-179) isolate were used as positive controls. Water and uninfected samples were used as negative controls in all amplifications.

**CSP sequencing and phylogenetic analyses**

Four internal primers were designed specifically on the *P. malariae*csp gene region and used together with the published primers (7) (Technical Appendix Table) to unambiguously amplify the three segments, the N-terminal, the central repeat, and the C-terminal regions of the *csp* gene. A total of 37 *P. malariae* isolates were amplified and sequenced. Amplification was conducted in a 20µL reaction mixture containing 2µL of genomic DNA, 10µL 2××DreamTaq™ Green PCR Master Mix (Fermentas) and 0.5µL of 10 µM primers. PCR cycles included an initial
denaturing step at 95°C for 3 min, 40 cycles of 95°C for 30 sec, 48-50°C for 30 sec, and 72°C for 2 min, followed by an additional extension at 72°C for 5 min in a Bio-Rad MyCycler Thermal Cycler. PCR products were visualized on 1% agarose gel and then purified and sequenced from both ends with BigDye Terminator v3.1 Cycle Sequencing Kit on an ABI 3130xl sequencer.

All obtained sequences were blasted against NCBI GenBank database for verification. They were translated into protein sequences and analyzed together with all available csp protein sequences of *P. malariae* as well as *P. brasilianum* retrieved from the GenBank database. Due to potential alignment errors associated with gaps in the nucleotide sequences, translated amino acid sequences with unambiguous indels were used in phylogenetic analyses. Sequences were aligned with MUSCLE v3.7 (8) using default settings followed by manual editing in Sequence Alignment Editor v1.d1 (9). A phylogenetic tree was reconstructed using the maximum likelihood method implemented in the PhyML program v3.0 (10). The WAG (Whelan And Goldman) substitution model, which assumes an estimated proportion of invariant sites and four gamma-distributed rate categories to account for rate heterogeneity across sites, was selected. Resulted trees were visualized in FigTree v1.4.2.

Sequence diversity including measures of evolutionary distances and average pairwise divergence were estimated using SSE v1.2 (11). The matrix-normalized distances based on the standard PAM model (12) that accounts for the probability of change from one amino acid to another were calculated. In addition, a similarity scan was performed between and within sequences using the standard PAM-Dayhoff matrix to normalize a matching score. Regions that meet or exceed the set criteria of the number of matches were plotted on a dot plot graph. *Plasmodium malariae* of East Africa (Kenya and Uganda), Central/West Africa (Cameroon and Cote d’Ivoire), and South America (Brazil and Venezuela) were compared for level of sequence diversity among gene regions.

**References**


Technical Appendix Table. Primer sequences and PCR conditions of the circumsporozoite protein (csp) gene

<table>
<thead>
<tr>
<th>Primer name</th>
<th>Sequence</th>
<th>Expected size (bp)</th>
<th>Annealing temperature</th>
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<tbody>
<tr>
<td>csp-F*</td>
<td>ATGAAGAAGTTATCCTGGCTTAGCAATATCC</td>
<td>280</td>
<td>50°C</td>
</tr>
<tr>
<td>csp-280R</td>
<td>CGGGGGGGGTTTTTCATTATTTATTC</td>
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<tr>
<td>csp-280F</td>
<td>GCTGTTGAAATCCATTGAACCAACC</td>
<td>700-800</td>
<td>48°C</td>
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<tr>
<td>csp-1070R</td>
<td>CCGCTTTTATTATCCCTTTTTTCGC</td>
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<td></td>
</tr>
<tr>
<td>csp-1070F</td>
<td>GCGAAAAATAAGGATTAAGGATGG</td>
<td>400</td>
<td>50°C</td>
</tr>
<tr>
<td>csp-R*</td>
<td>TTAGTGAAGAGTATTAAAGCTAAAAC</td>
<td></td>
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</tr>
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*Primer published in (7).

Technical Appendix Figure 1. Scatter plot showing the significant correlation of parasite gene copy number measured by quantitative real-time PCR and parasitemia by microscopy of *Plasmodium malariae* isolates.
Technical Appendix Figure 2. Scatter plots showing (A) parasite gene copy number of *P. malariae* and *P. falciparum* ranked from low to high *P. malariae* parasite gene copy number and (B) parasitemia of *P. malariae* and *P. falciparum* ranked from low to high *P. malariae* parasitemia of co-infected samples.
Technical Appendix Figure 3. Scatter plot showing the non-significant correlation of *P. malariae* gene copy number against age among samples.

$ r^2 = 0.07$

$P > 0.05$