Molecular surveillance of *Plasmodium falciparum* drug resistance in the Republic of Congo: four and nine years after the introduction of artemisinin-based combination therapy

Felix Koukouikila-Koussounda1†, Sankarganesh Jeyaraj2†, Christian N. Nguetse2, Charles Nchotebah Nkonganyi2, Kossiwa Clarisse Kokou2, Mandingha K. Etoka-Beka1, Francine Ntoumi1,2 and Thirumalaisamy P. Velavan1,2,3,4*

**Abstract**

**Background:** Resistance to anti-malarial drugs hinders efforts on malaria elimination and eradication. Following the global spread of chloroquine-resistant parasites, the Republic of Congo adopted artemisinin-based combination therapy (ACT) in 2006 as a first-line treatment for uncomplicated malaria. To assess the impacts after implementation of ACT, a molecular surveillance for anti-malarial drug resistance was conducted in Congo 4 and 9 years after the introduction of ACT.

**Methods:** Blood samples of 431 febrile children aged 1–10 years were utilized from two previous studies conducted in 2010 (N = 311) and 2015 (N = 120). All samples were screened for malaria parasites using nested PCR. Direct sequencing was used to determine the frequency distribution of genetic variants in the anti-malarial drug-resistant *Plasmodium falciparum* genes (*Pfcrt, Pfmdr1, Pfatp6, Pfk13*) in malaria-positive isolates.

**Results:** One-hundred and nineteen (N = 70 from 2010 and N = 49 from 2015) samples were positive for *P. falciparum*. A relative decrease in the proportion of chloroquine-resistant haplotype (CVIET) from 100% in 2005, 1 year before the introduction and implementation of ACT in 2006, to 98% in 2010 to 71% in 2015 was observed. Regarding the multidrug transporter gene, a considerable reduction in the frequency of the mutations N86Y (from 73 to 27%) and D1246Y (from 22 to 0%) was observed. However, the prevalence of the Y184F mutation remained stable (49% in 2010 compared to 54% in 2015). Isolates carrying the *Pfatp6* H243Y was 25% in 2010 and this frequency was reduced to null in 2015. None of the parasites harboured the *Pfk13* mutations associated with prolonged artemisinin clearance in Southeast Asia. Nevertheless, 13 new *Pfk13* variants are reported among the investigated isolates.

**Conclusion:** The implementation of ACT has led to the decline in prevalence of chloroquine-resistant parasites in the Republic of Congo. However, the constant prevalence of the *PfMDR1* Y184F mutation, associated with lumefantrine susceptibility, indicate a selective drug pressure still exists. Taken together, this study could serve as the basis for...
Background
In the Republic of Congo, malaria remains one of the leading causes of morbidity and mortality. Chloroquine and sulfadoxine–pyrimethamine were extensively used for several decades as first-line treatment for uncomplicated malaria, but parasites rapidly developed resistance to both these drugs [1–5]. As a result, in 2006, Congo changed its national drug policy and switched to artesunate + amodiaquine (ASAQ) and artemether + lumefantrine (AL) combinations as first-line and second-line drugs, respectively, for the treatment of acute uncomplicated malaria.

Chloroquine resistance (CQR) is caused by mutations in two genes, Plasmodium falciparum chloroquine resistance transporter (Pfcrt) and multidrug resistance transporter-1 (Pfmdr1), both located on the digestive food vacuole of the parasite [6]. These two genes also reduce the susceptibility to other quinolone anti-malarial agents such as amodiaquine, lumefantrine and mefloquine [7–9]. Many studies reported that in the absence of drug pressure, chloroquine-resistant strains have been replaced by sensitive ones [10, 11].

Many P. falciparum candidate genes are believed to be associated with artemisinin resistance. The endoplasmic and sarcoplasmic reticulum Ca²⁺-ATPase orthologue of P. falciparum (Pfatp6) has been postulated to be the target of artemisinin [12]. The H243Y, L263E, E431K, A623E, and S769N mutations have been shown to influence the functional activities in Pfatp6 [13–15]. In 2014, Ariey et al. showed that mutations (C580Y, R539T, Y493H, M476I) in the Pfk13 gene were associated with delayed parasite clearance [16]. As chloroquine-resistant P. falciparum strains spread worldwide from Southeast Asia, a similar devastating effect can also occur in case of artemisinin-resistant isolates. However, it has been shown that artemisinin-resistant parasites could also emerge independently in Southeast Asia, as Pfk13 mutations occur unexpectedly in many sites within the region [17].

Long-term monitoring of parasite sensitivity to previously withdrawn anti-malarial drugs, such as CQ, can provide useful surveillance information if these drugs target similar resistance markers to current or candidate ACT partner drugs [18]. Therefore, the aim of this study was to determine the prevalence of polymorphisms in the anti-malarial resistance genes, including Pfcrt, Pfmdr1, Pfatp6, and Pfk13 in field isolates from the Republic of Congo collected from two cross-sectional studies conducted in 2010 and 2015, approximately four and nine years after the adoption of ACT.

Methods
Study site and sample collection
The study was conducted in the Southern and Northern districts in Brazzaville, the capital of the Republic of Congo. Malaria occurs holo-endemically and the transmission rates in the country are high and perennial, the majority of malaria cases being caused by P. falciparum. A total of 431 children aged 1–9 years were recruited in two phases: from April to June 2010 [Group A (N = 311)] and from September 2014 to February 2015 [Group B (N = 120)] from previous studies [19, 20] were utilized in this study. The 2010 cohort was from three districts of Makenkékané health division in southern region of Brazzaville, Republic of Congo. This cohort comprised of children aged 1–9 years and are permanent residents of the study area. The second cohort (September 2014 to February 2015) was collected from children aged from 1 to 10 years presenting with fever at the paediatric ward of the MNG hospital in Talangai, a northern Brazzaville region in Republic of Congo. This study was conducted in febrile children presenting with fever at the MNG hospital. Children who were positive for malaria by microscopy were included. Febrile children with other pathologies such as severe diarrhoea, pulmonary infection and HIV were excluded. At the time of enrolment 4 mL of whole blood were collected in heparinized tubes from all children and thick and thin blood smears were performed. In the case of malaria (positive thick and thin blood smears with axillary temperature ≥37.5 °C), collected blood samples were stored at −80 °C for molecular analyses, parasite density determined and children were treated with AS + AQ or AL. Ethical approval was given by the Institutional Ethics Committee for Research on Health Sciences of the Republic of Congo. Written informed consent was obtained from parents or guardians of children.

Plasmodium species identification, and Pfcrt, Pfmdr1, Pfatp6, Pfk13 genotyping
Genomic DNA was extracted from 200 μL of peripheral whole blood samples using QIAamp DNA Blood Mini Kit.
(Qiagen, Hilden, Germany). The *Plasmodium* 18S rRNA gene was amplified by nested PCR to detect positive samples and further differentiate the species as described elsewhere [21, 22]. The primer pairs and thermocycling conditions are summarized in Table 1. Fragments of the *Pfcrt* (410 bp containing the polymorphisms at codons 72–76), *Pfmdr1* [(580 bp including the polymorphisms at codons 86 and 184) and (864 bp containing the polymorphisms at codons 1034–1246)], *Pfatp6* (800 bp carrying the polymorphisms at codons 243–769) and *Pfk13* (849 bp comprising the main polymorphisms associated with delayed clearance in Southeast Asia) were amplified by nested PCR as described elsewhere [11, 23, 24]. The primer sequences and PCR conditions are described in Table 2. The PCR products were visualized through electrophoresis on 1.2% agarose gel stained with SYBR Green I in 1× Tris-electrophoresis buffer (90 mM Tris–acetate, pH 8.0, 90 mM boric acid, 2.5 mM EDTA). Thereafter, PCR products were purified using Exo-SAP-IT (USB, Affymetrix, USA) and directly used as templates for DNA sequencing using the BigDye terminator v. 1.1 cycle sequencing kit (Applied Biosystems, Foster City, USA).

| Genus and species | Primer name | Primer sequence (5′-3′) | PCR conditions | Reference |
|-------------------|-------------|-------------------------|----------------|-----------|
| *Plasmodium* genus | *rPLU* 1    | TCAAAGATTAAGCCATGCAAGTGA | 95 °C for 5 min; 25× [94 °C for 1 min; 58 °C for 1 min; and 72 °C for 2 min]; 72 °C for 5 min | Snounou et al. [22] |
|                   | *rPLU* 5    | CCTGTTGTTGCCTTAAACTTC   |                |           |
| *P. falciparum*    | *rFAL1*     | TTAACCTGTTTGGAAAACAAATATT | 95 °C for 5 min; 30× [94 °C for 1 min; 56 °C for 1 min; and 62 °C for 1 min] |           |
|                   | *rFAL2*     | ACACAATGAACTCAATCATGACTACCCGTC |                |           |
| *P. malariae*      | *rMAL1*     | AAATTCGACATGCTATACAAAAATTATACAAA | | Fuehrer et al. [21] |
|                   | *rMAL2*     |                          |                |           |
| *P. vivax*         | *rVIV1*     | GCAGTTCACACGCGAAGGAAAGAAG | 35 cycles of 94 °C for 30 s; 56 °C for 30 s; and 62 °C for 1 min | Mekonnen et al. [11] |
|                   | *rVIV2*     | TCTTACCTCTTACTTAAACCATGATC-TAC |                |           |
| *P. ovale*         | *rOVA1WC*   | TGTAGATTCATACACCGACTG | 34 cycles of 94 °C for 1 min and 72 °C for 2 min | Fuehrer et al. [21] |
|                   | *rOVA2WC*   | TAGTACCTGTGAAGGCCTT |                |           |

| Loci              | Primer name | Primer sequence (5′-3′) | PCR conditions | Reference |
|-------------------|-------------|-------------------------|----------------|-----------|
| *Pfcrt* SNPs at codons 72 and 76 | *OF* P1 | 5′-CCGTTAATAAATAACACCGC-3′ | 35 cycles of 94 °C for 30 s; 56 °C for 30 s; and 62 °C for 1 min | Mekonnen et al. [11] |
|                   | *OR* P2    | 5′-CAGATATATACACCATGCTCC-3′ |                |           |
|                   | *NF* P3    | 5′-AGGTCTCTGTTCTGGT-3′ | 30 cycles of 94 °C for 30 s; 56 °C for 30 s; and 65 °C for 1 min |           |
|                   | *NR* P4    | 5′-CAAAACTATAGTCCACTTGTTT-3′ |                |           |
| *Pfmdr1* SNPs at codons 86 and 184 | *OF* P5 | 5′-AGGTTCAGAAAAAGGTTGAAAC-3′ | 30 cycles of 94 °C for 30 s; 55 °C for 30 s; and 65 °C for 1 min |           |
|                   | *OR* P6    | 5′-ATGACACCCAACAAATAAT-3′ |                |           |
|                   | *NF* P7    | 5′-ACAAAAAGAAGTACCGCTGAAT-3′ | 30 cycles of 94 °C for 30 s; 60 °C for 30 s; and 65 °C for 1 min |           |
|                   | *NR* P8    | 5′-AAACCGCAAAGTAAACATAAAA-GTC-3′ |                |           |
| *Pfmdr1* SNPs at codons 1034, 1024, and 1246 | *OF* P9 | 5′-CTCTACTACCTGGTCTTGGATC-3′ | 34 cycles of 94 °C for 30 s; 55 °C for 1 min and 72 °C for 2 min |           |
|                   | *OR* P10   | 5′-GACATATATACACCATGCTCC-3′ |                |           |
|                   | *NF* P11   | 5′-CAGATATGAAATGTCTTAAA-3′ | 29 cycles of 94 °C for 30 s; 60 °C for 30 s; and 65 °C for 1 min |           |
|                   | *NR* 12    | 5′-CAGAATACCCCAATGCTTGA-3′ |                |           |
| *Pfatp6* at codon 243, 263 and 431 | *NF* P13 | 5′-TCATACCTACCCGCTTATGGT-3′ | 35 cycles of 94 °C for 45 s; 55 °C for 1 min and 72 °C for 1 min | Zakeri et al. [24] |
|                   | *NR* P14   | 5′-TCCTCTTACACTCACCAATGCTCC-3′ |                |           |
| *Pfatp6* SNPs at codon 623 and 769 | *NF* P15 | 5′-TGAGAGATGCCCCCCCTGAAATGC-3′ | 30 cycles of 95 °C for 30 s; 58 °C for 2 min; and 72 °C for 2 min | Ariey et al. [16] |
|                   | *NR* P16   | 5′-TGCTACATATGCTGTTGGTCT-3′ |                |           |
| *Pfk13*            | *OF* P17   | 5′-CGGATACTGGACATGGCAGAA-3′ | 40 cycles of 95 °C for 30 s; 60 °C for 30 s; and 65 °C for 1 min |           |
|                   | *OR* P18   | 5′-CGGAACTCTGTTGGTCAACAGC-3′ |                |           |
|                   | *NF* P19   | 5′-GCCAAGCTGCCATTGATTG-3′ | 30 cycles of 95 °C for 30 s; 58 °C for 2 min; and 72 °C for 2 min |           |
|                   | *NR* P20   | 5′-GCCTTGTGTAAGAGCAGA-3′ |                |           |

OF outer forward, OR outer reverse, NF nested forward, NR nested reverse.
on an ABI 3130XL DNA sequencer. Single nucleotide polymorphisms (SNPs) were identified by assembling the sequences with each reference sequence using Codon code Aligner 4.0 software and were reconfirmed visually from their respective electropherograms.

Results

**Plasmodium species identification**

Using nested PCR for the species identification, 22.5% (70/311) and 40.8% (49/120) samples collected in 2010 and 2015, respectively, were positive for *P. falciparum*. The other species namely *Plasmodium malariae*, *Plasmodium vivax* and *Plasmodium ovale* were not detected in the study samples. The positive samples for *P. falciparum* malaria infection were screened for genetic variants in the *Pfcrt*, *Pfmdr1*, *Pfatp6*, and *Pfk13* associated with antimalarial drug resistance.

**Pfcrt polymorphisms**

The success rate of *Pfcrt* amplicon sequencing was equally 57% with the samples from the 2010 cohort (40/70) and the 2015 cohort (28/49). The mutation frequencies are shown in Table 3. A triple mutation M74I, N75E and K76T was observed, each occurring at a frequency of 98% in 2010 and 71% in 2015. All the mutants carried the Cysteine–Valine–Isoleucine–Glutamate–Threonine (CVIET) haplotype while the wild type strains had the Cysteine–Valine–Methionine–Asparagine–Lysine (CVMNK) haplotype.

**Pfmdr1 polymorphisms**

Forty-one samples (59%) from 2010 and 26 samples (53%) from 2015 were sequenced for the detection of *Pfmdr1* polymorphisms. The frequency of the *Pfmdr1* gene mutations at positions 86 and 1246 are shown in Table 3. The frequency of N86Y mutant alleles was 73% in 2010 and 27% in 2015. The D1246Y mutation in 2010 occurred at a frequency of 22 and 0% in 2015. Neither the S1034C mutation nor the N1042D mutation in the *Pfmdr1* gene was observed in the sequenced samples. The prevalence of the Y184F variant was 49% in 2010 and 54% in 2015. Other mutations occurred at quite low frequencies are represented in Table 3.

**Pfatp6 polymorphisms**

Eight samples (11%) from 2010 and 16 samples (33%) from 2015 were successfully sequenced to identify mutations in the *Pfatp6* gene. Out of all the five key point mutations that were investigated, only H243Y and E431K mutations were observed. Both mutations occurred in 2010 at a frequency of 25% (2/8) and 12% (1/8), respectively, while in 2015, only the E431K mutation was observed occurring at a frequency of 12% (2/16). Other mutations found at positions 291 (I291V), 402 (L402V), 569 (N569K), 630 (A630S), 632 (G632E), 639 (G639D), 646 (F646F), and 747 (H747Y) are shown in Table 4. In addition to these SNPs, a variation of asparagine-tandem repeat region (codon 457–465) was observed in one sample.

**Table 3: Prevalence of *Pfcrt* and *Pfmdr1* polymorphisms**

| Locus | Mutation | Cohort 2010 (%) | Cohort 2015 (%) |
|-------|----------|----------------|----------------|
| *Pfcrt* | M74I, N75E and K76T | 39/40 (98) | 20/28 (71) |
| *Pfmdr1* | N86Y | 30/41 (73) | 7/26 (27) |
| | S113R | 1/41 (2) | 0/26 (0) |
| | Y184F | 20/41 (49) | 14/26 (54) |
| | T1069T | 8/41 (20) | 3/26 (12) |
| | G1113G | 0/41 (0) | 1/26 (4) |
| | D1127D | 1/41 (2) | 0/26 (0) |
| | T1157T | 2/41 (5) | 2/26 (8) |
| | V1225A | 1/41 (2) | 0/26 (0) |
| | D1246Y | 9/41 (22) | 0/26 (0) |

Values in parentheses represent the percentage and is rounded
*N. falciparum* positive samples

**Table 4: Prevalence of *Pfatp6* and *Pfk13* gene polymorphisms**

| Locus | Mutation | Cohort 2010 (%) | Cohort 2015 (%) |
|-------|----------|----------------|----------------|
| *Pfatp6* | H243Y | 2/8 (25) | 0/16 (0) |
| | L291V | 0/8 (0) | 1/16 (6) |
| | L402V | 1/8 (12) | 2/16 (12) |
| | E431K | 1/8 (12) | 2/16 (12) |
| | N569K | 5/8 (63) | 2/26 (12) |
| | A630S | 3/8 (38) | 0/16 (0) |
| | G632E | 0/8 (0) | 1/16 (6) |
| | G639D | 0/8 (0) | 1/16 (6) |
| | F646F | 1/8 (12) | 0/16 (0) |
| | H747Y | 0/8 (0) | 1/16 (6) |

**Pfk13**

| Locus | Mutation | Cohort 2010 (%) | Cohort 2015 (%) |
|-------|----------|----------------|----------------|
| D464N | 1/41 (2) | 0/25 (0) |
| S466I | 0/41 (0) | 1/25 (4) |
| Q467H | 0/41 (0) | 1/25 (4) |
| N489N | 0/41 (0) | 1/25 (4) |
| G496G | 0/41 (0) | 1/25 (4) |
| R539R | 1/41 (2) | 0/25 (0) |
| S549P | 1/41 (2) | 0/25 (0) |
| I552M | 0/41 (0) | 1/25 (4) |
| L598L | 1/41 (2) | 0/25 (0) |
| E602E | 1/41 (2) | 0/25 (0) |
| N609N | 0/41 (0) | 1/25 (4) |
| F628L | 0/41 (0) | 1/25 (4) |
| K669K | 1/41 (2) | 0/25 (0) |

Values in parentheses represent the percentage
*N. falciparum* positive samples
sample from 2010, with an insertion of two asparagine residues after the 465th codon.

**Pfk13 polymorphisms**

Regarding the Kelch13 propeller domain, 41 out of 70 samples (59%) for 2010 cohort and 25 out of 49 samples (51%) for 2015 cohort were sequenced and used for analysis. The identified mutations in the Kelch13 propeller domain are shown in Table 4. The Pfk13 variants M476I, Y493H, R539T, I543T, and C580Y reported to occur in Southeast Asia were not observed in the Republic of Congo. Thirteen new substitutions were identified in the Pfk13 propeller domain. Six Pfk13 non-synonymous substitutions were observed in the propeller blades. The Pfk13 non-synonymous variants I552M and S549P observed in this study were previously reported in Central African Republic and in India, respectively.

**Discussion**

Malaria is still considered as one of the major health problems in the Republic of Congo and uncomplicated malaria occurs in young children due to *P. falciparum*. Malaria incidence is 0.9 malaria episode/year/child in Brazzaville and malaria transmission is perennial both in southern and northern Brazzaville, with *P. falciparum* being the most predominant species. In this study, *P. falciparum* was the only species detected in all the malaria-positive samples. Several factors, including both malaria parasite (e.g., pyrogenic threshold, multiplication rate, cytoadhesion) and host genetics (e.g., immunity, tolerance, pregnancy, co-morbidities) can influence the prevalence of the disease [25]. The wide spread of CQ-resistant parasites prompted the WHO to recommend ACT for the management of the disease in endemic regions. In 2006, Congo changed its national policy introducing ACT for the treatment of uncomplicated malaria. The first report describing the prevalence of polymorphisms in PfCRT conferring CQR showed that all the *P. falciparum* isolates were carrying the PfCRT mutant alleles [2, 5]. The PfCRT CVIET haplotype has been shown to be the most prevalent in Africa in contrast to the SVMNT haplotype found predominantly in Southeast Asia [23, 26]. Also, the occurrence of the PfCRT M74I and N75E mutations has been linked to the K76T mutation as reported by Severini et al. who noted a 100% linkage between these alleles [27]. This explains the occurrence at the same frequency of these three mutations in this study. Studies conducted in Malawi and Tanzania reported a successful restoration of CQ-sensitive strains after the implementation of ACT for the treatment of uncomplicated falciparum malaria [28–30]. The return of these CQ-sensitive strains as observed in other studies support the hypothesis that the removal of drug pressure has led to a re-expansion of a heterogeneous susceptible parasites that might have prevailed when CQ was used.

Likewise, a significant decrease in the prevalence of the PfCRT haplotypes associated with CQR was noted in this study from 2005 in republic of Congo [2], 1 year before the treatment policy was changed. However, their frequency did not decrease further after 9 years since the implementation of ACT, suggesting a possible use of CQ in the region by the population owing to the fact that it is less expensive than ACT.

Mutations in the Pfmdr1 and changes in its copy number have been associated with the development of decreased parasite sensitivity to several anti-malarial drugs. The major SNPs in the Pfmdr1 gene are N86Y, Y184F and D1246Y. The wild-type N86 has been shown to be responsible for the increased tolerance to drugs such as artemether [31] and 86Y associated with delayed parasite clearance with parenteral artesunate [32]. The mutant allele 86Y has been shown to associate with increased susceptibility to mefloquine in in vitro experiments [33]. Many studies that evaluated the influence of Pfmdr1 mutations in vitro have reported that, 184F or 1042N alleles decrease artemether and lumefantrine sensitivity in Thai isolates [34], Pfmdr1 86Y and PfCRT76T in Kenyan isolates [35]. A yet another recent study from Burkina Faso reports on influence of Pfmdr1 86Y and PfCRT76T mutations in AL and ASAQ treatment [36]. The N86Y and Y184F mutants are prevalent in Asian and African continents while S1034C, N1042F and D1246Y alleles are common in South American countries [37]. It is interesting to note that 86Y mutant allele prevalence is decreased in 2015 when compared to 2010 cohort samples. Pfmdr1 86Y and 1246Y alleles are linked to CQR in Africa and the 86Y allele is associated with decreased quinine sensitivity in African clinical isolates. The fact that Pfmdr1 gene mutations are not the principal actor in CQR [38, 39] might explain the decrease in the frequency of the Pfmdr1 mutations linked to CQR observed between 2010 and 2015. Both Pfmdr1 and PfCRT genes are present on the membrane of the digestive vacuole in the malarial parasite and is believed to regulate the chemical solutes from the drug across the membrane. Few studies have reported that certain Pfmdr1 and PfCRT alleles are in linkage and substantiated their functional relatedness [40, 41].

The occurrence of key mutations in the Pfput6 gene linked to artemisinin resistance before the use of artemisinin and its derivatives in malaria treatment has been a major subject of controversy. The main question raised is whether these mutations occur as a result of some natural events or through prior exposure of parasites to drug pressure [14, 24]. One possible confounding factor could be the use of herbal plants in the treatment of malaria,
which might have similar chemical constituents to arte-
minisin, and which is common practice in many malaria-
endemic countries. Only the Pfatp6 H243Y and E431K
mutations were identified among the five key mutations
investigated. The variation in the asparagine-tandem
repeat region had been previously reported by Tanabe
et al. [42] but this had no role in artemisinin resistance.
Despite the fact that none of the driving mutations for
artemisinin resistance in the Pfk13 propeller domain
reported by Ariey et al. [16] was identified, in the present
study non-synonymous variants I552M and S549P are
detected, which were previously reported in isolates from
the Central African Republic and India, respectively.

Conclusions
The new and unreported mutations identified in this
study could help in long-term surveillance of the develop-
ment of artemisinin resistance in the Republic of
Congo. This study on molecular surveillance will support
to predict sensitivity to different anti-malarials used in
Republic of Congo and will additionally provide informa-
tion on fixation of any multidrug resistance alleles in this
population. The results observed in this study could serve
as the basis for epidemiological studies monitoring the
distribution of molecular markers of anti-malarial drug
resistance in the Republic of Congo.

Abbreviations
Pfmdr1: Plasmodium falciparum multidrug resistance 1; Pfatp6: Plasmodium
falciparum Ca2+ ATPase; Pfk13: Plasmodium falciparum Kelch-13 propeller
domain; ACT: artemisinin-based combination therapies; WHO: World Health
Organization.

Authors’ contributions
TPV and FN designed, supervised the experiments, performed data analysis
and wrote the manuscript. FFK, SJ, CNN, CNN, and KCK performed the experi-
ments. MKEB contributed to the study design and sampling procedures. All
authors read and approved the final manuscript.

Author details
1 Fondation Congolaise pour la Recherche Médicale (FCRM), Brazzaville,
Republic of Congo. 2 Institute of Tropical Medicine, University of Tübingen,
Tübingen, Germany. 3 Duy Tan University, Da Nang, Vietnam. 4 Vietnamese-
German Centre for Medical Research, Hanoi, Vietnam.

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Competing interests
The authors declare that they have no competing interests.

Availability of data and materials
All relevant data are within the paper.

Consent for publication
All authors read and approved to its final submission.

Ethics approval and consent to participate
Written informed consent was obtained from parents or guardians of all
participating children. The study was approved by the Institutional Ethics
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