**Plasmodium falciparum** genetic factors rather than host factors are likely to drive resistance to ACT in Ghana

Peter Hodoameda1, Nancy Odurowah Duah-Quashie2, Charles Oheneba Hagan4, Sena Matrevi2, Benjamin Abuaku2, Kwadwo Koram2 and Neils Ben Quashie1,2,3*

**Abstract**

**Background:** Artemisinin-based combination therapy (ACT) partner drugs, currently used in Ghana are lumefantrine, amodiaquine and piperaquine. *Plasmodium falciparum* isolates with reduced susceptibility to these partner drugs may affect treatment outcome. Mutations in *pfmdr1* gene is linked to reduced parasite susceptibility to amodiaquine and lumefantrine. In addition, the potency of the partner drugs in vivo depends on the metabolism by the cytochrome P450 (CYP) enzyme in the host. Mutations in the CYP2C8 and CYP3A4 genes are linked to reduced metabolism of amodiaquine and lumefantrine in vitro, respectively. This study investigated the host and parasite genetic factors affecting the susceptibility of the malaria parasite to ACT partner drugs.

**Methods:** Archived samples from 240 patients age ≤ 9 years participating in anti-malarial drug resistance survey in Ghana, and given artemether with lumefantrine (AL) or artesunate with amodiaquine (AA), were selected and analysed. Polymerase chain reaction (PCR) followed by Sanger sequencing was used to determine the polymorphisms in CYP2C8, CYP3A4 and pfmdr1 genes.

**Results:** For CYP3A4, all had wild type alleles, suggesting that the hosts are good metabolizers of lumefantrine. For CYP2C8 60% had wild type alleles, 35% heterozygous and 5% homozygous recessive alleles suggesting efficient metabolism of amodiaquine by the hosts. For pfmdr1 gene, at codon 86, 95% were wild type (N86) and 5% mutant (Y86). For codon 184, 36% were wild type (Y184) and 64% mutant (F184) while for codons 1034, 1042 and 1246, 100% (all) were wild type. The high prevalence of N86-F184-D1246 haplotype (NFD) suggest presence of parasites with reduced susceptibility to lumefantrine and not amodiaquine. Delayed clearance was observed in individuals with mutations in the pfmdr1 gene and not cytochrome 450 gene. Both synonymous and non-synonymous mutations were observed in the pfmdr1 at low prevalence.

**Conclusion:** The outcome of this study indicates that the parasite’s genetic factors rather than the host’s are likely to drive resistance to ACT in Ghana.

**Keywords:** Polymorphism, Mutation, Resistance, Cytochrome, Molecular markers, Prevalence
relatively slow-acting partner drug, for the treatment of uncomplicated malaria in disease-endemic areas [2]. The recommended first-line artemisinin combination therapy (ACT) in Ghana for treating uncomplicated malaria is artesunate with amodiaquine (AA), artemether with lumefantrine (AL) or a combination of dihydroartemisinin with piperaquine [3]. The reason for combining the drugs (ACT) is to slow down the development of resistance to anti-malarial drugs by *P. falciparum* [4]. The fast-acting drug quickly reduces parasite load whilst the slow-acting anti-malarial gradually destroys residue parasites. The potency of ACT is dependent on the efficacy of both the artemisinin component and the partner drug [4]. Reduced susceptibility of parasites to partner drugs in ACT can potentially result in resistance to artemisinin in future as parasites that escape the fast action of artemisinin or its derivatives will not be cleared by the partner drug and this could allow for growth and expansion of a drug-resistant parasite population [4].

Variations observed in effectiveness of ACT in malaria-endemic regions are dependent on parasite genetic factors [5], as well as human genetic factors [6]. For parasite genetic factors, polymorphisms which arise due to single nucleotide changes in the *pfmdr1* gene in its coding region have been linked to differential parasite susceptibility to ACT partner drugs, such as amodiaquine [7] and lumefantrine [8]. This makes *pfmdr1* an important likely candidate for initiating ACT partner drug resistance [9].

The polymorphic *pfmdr1* alleles that are mostly found in Africa are N86Y, F184Y and D1246Y. The *P. falciparum* N86-F184-D1246 haplotype (NFD haplotype) has been linked to decreased susceptibility to anti-malarial drugs, such as melfloquine and lumefantrine. The selection of the NFD haplotype has been seen in malaria treatment using AL. The different haplotype, which is the Y86-Y184-Y1246 haplotype (YYY haplotype), is associated with reduced amodiaquine susceptibility [10].

Differences in genetic make-up of humans is the principal factor that defines the level of drug availability in the blood to clear the parasites [6]. The cytochrome P450 enzyme family (CYP genes) is involved in the metabolism of different anti-malarial drugs [6]. Amodiaquine is mainly metabolized by CYP2C8 [11] while lumefantrine is metabolized mainly by CYP3A4 [12]. Different mutations in the promoter region, introns or exons can result in different alleles of the CYP450 genes in different individuals. The metabolism of a drug or a combination of drugs could be decreased, increased or unaffected depending on the allele(s) an individual possesses [13]. Elucidating the exact role these disparities in the genes coding for the enzymes involved in ACT metabolism is vital for understanding the inter-individual pharmacokinetic differences observed in persons using ACT [14].

This study investigated *P. falciparum* and host genetic factors that are likely to affect the efficacy of ACT partner drugs used in Ghana.

**Methods**

**Study design**

Blood blot filter paper used in this study were archived samples collected in 2016 from children participating in studies previously described by Abuaku et al. [15]. The study of Abuaku et al. was part of routine surveillance on the therapeutic efficacy of ACT in Ghana; the efficacy of AA and AL were studied in six sentinel sites representing the forest and savannah zones of the country. Three sites representing the two ecological zones studied AA whilst the other three sites studied AL. Additionally, archived samples from a site in the Coastal savanna ecological zone was included in the analysis. In each site, the study was a one-arm, prospective evaluation of the clinical, parasitological and haematological responses to directly observed treatment with either AA or AL among children 6 months to 9 years old with uncomplicated *falciparum* malaria. The WHO 2009 protocol on surveillance of anti-malaria drug efficacy was used for the study with primary outcomes as prevalence of day 3 parasitaemia and clinical and parasitological cure rates on day 28.

An informed consent was obtained from each parent or guardian. A medical doctor prescribed either AA or AL to the study participants who were then followed up for 28 days.

The archived samples were selected from three sentinel sites, Navrongo, Begoro and Cape Coast, which represent three distinct eco-epidemiological zones in Ghana (Fig. 1). Begoro is located in the tropical forest ecological zone, Navrongo is in the northern savanna ecological zone and Cape Coast is situated in the coastal savanna ecological zone. The samples (100 μl blood) were collected on Whatman 3 filter paper (Sigma, UK), stored in plastic bags containing silica gel, and kept at room temperature until use.

The samples from a group of participants referred hereafter as ‘cohort 1’ comprised 120 recruited patients who were given AL, and a second group, cohort 2, comprised 120 patients who received AA. Of the 240 study participants’ archived samples analysed, 60 were selected from the savannah zone, 90 from the coastal zone and 90 from the forest zone.

**DNA extraction**

Malaria parasite DNA was extracted from the archived blood blots filter papers using a QIAamp DNA minikit (Qiagen, Germany) following the manufacturer’s protocol. Conventional PCR [16] was performed to amplify the region of interest using published protocols [17–19].
PCR products were sequenced using Sanger sequencing at Macrogen, The Netherlands.

**Detection of pfmdr1 polymorphisms**

The regions of pfmdr1 gene was amplified and sequenced to determine the presence of any mutation. The amplification was carried out using the protocol described by [17]. The polymorphisms were analysed at codons 86 (asparagine to tyrosine), 184 (tyrosine to phenylalanine), 1034 (serine to cysteine), 1042 (asparagine to aspartic acid) and 1246 (aspartic acid to tyrosine). A PCR followed by Sanger sequencing was used in determining these polymorphisms. Thirty microlitres aliquot of PCR products were kept on ice and shipped for sequencing at Macrogen, The Netherlands.

**Detection of CYP2C8 and CYP3A4 polymorphisms**

The CYP2C8 polymorphisms were analysed at codon 269 (isoleucine to phenylalanine). The CYP3A4 polymorphisms were analysed at position -392A>G of the proximal promoter region. A PCR followed by Sanger sequencing was used to determine the polymorphism in CYP2C8 as reported by Cavaco et al. [18] and CYP3A4 as described by Hodel et al. [19]. A PCR product of 120 bp and 717 bp represents a successful amplification of CYP2C8 and CYP3A4, respectively. Aliquot of the PCR products was shipped appropriately for sequencing at Macrogen, The Netherlands.

**Data analysis**

Data were organized using R software, SPSS software (version 20) and GraphPad Prism version 6. Sequence data were analysed with the BLAST program (https://blast.ncbi.nlm.nih.gov/) to determine the authenticity of the sequences. Multiple sequences were aligned with MAFFT (EMBL.EBI, Hinxton, Cambridge, UK) using the 3D7 wild-type as reference. Consensus sequence editing and single nucleotide polymorphism (SNP) detection was carried out using the CLC Main Workbench 7.9.1 (Qiagen, Aarhus, Denmark). The CYP2C8 sequences were aligned to CYP2C8 (ENSG 0000138115) as reference sequence while CYP3A4 sequences were aligned to CYP3A4 (ENSG 0000160868) as reference sequence from NCBI database. Genotype deviations from the Hardy–Weinberg equilibrium were also determined. The Hardy–Weinberg equilibrium determines whether or not the allele or genotype frequencies for a particular gene will remain constant from generation to generation in the absence of evolutionary influences such as genetic drift, inbreeding and founder effect [20]. All tests in this study were considered statistically significant when \( p \) value < 0.005.

**Results**

**Prevalence of individuals CYP3A4 and CYP2C8 genotype**

Ninety-three individuals were successfully genotyped for CYP3A4 of which all (100%) were wild type (Fig. 2). The genotypes analysed for CYP3A4 was not in Hardy–Weinberg equilibrium. Ninety-four individuals were successfully genotyped for CYP2C8 of which 60% (56/94) had wild-type alleles, 35% (33/94) heterozygous and 5% (5/94) homozygous recessive alleles (Fig. 2). The genotypes analysed for CYP2C8 was in Hardy–Weinberg equilibrium.

**Prevalence of isolates with pfmdr1 codons 86, 184, 1034, 1042, 1246 alleles and NFD haplotype**

Ninety-five P. falciparum out of 120 clinical isolates was successfully genotyped for the pfmdr1 gene. For pfmdr1 genotype at codon 86, 95% (90/95) were wild-type (N86) and 5% (5/95) mutant (Y86) (Fig. 3). For codon 184, 36% (34/95) were wild-type (Y184), 64% (61/95) mutant (F184) while for codons 1034, 1042 and 1246, all (100%) were wild-type. There were both non-synonymous and synonymous mutations observed at low frequencies in some of the samples analysed (Table 1). The pfmdr1 haplotypes observed were 57.8% (55/95) NFD, 34.7% (33/95) NYD, 6.3% (6/95) YFD, and 1% (1/95) YYD (Fig. 4).
Non-synonymous mutations that lead to change in amino acid and synonymous mutations which do not lead to a change in amino acid, were both observed at low frequencies in some of the samples analysed. It must be noted that natural selection on both synonymous and non-synonymous mutations plays an important role in shaping levels of synonymous polymorphism. In this study, six novel synonymous mutations were observed in the coastal zone and 9 in the savannah zone. Multiply synonymous mutations were observed in samples from savannah zone. With regard to the novel non-synonymous mutations, 5 were observed in samples from the coastal zone whilst 3 were seen in samples from the savannah zone.

**Discussion**

It is obvious that resistance of *P. falciparum* to ACT partner drugs may lead to the gradual evolution of strains of parasites with reduced susceptibility to artemisinin. The failure of partner drugs should therefore be of great concern to national malaria control programmes in disease-endemic areas. Since both the host and the parasite genome play a role in metabolism of ACT, a key question

![Fig. 2](image1) Prevalence of CYP3A4 and CYP2C8. (a) Genotype for CYP3A4, AA is wild-type, AG is heterozygous, and GG is homozygous recessive, (b) Genotype for CYP2C8, AA is wild-type, AT is heterozygous, and TT is homozygous recessive.

![Fig. 3](image2) Prevalence of *pfmdr1* codons 86, 184, 1034, 1042 and 1246. (a) *pfmdr1* 86, (b) *pfmdr1* 184, (c) *pfmdr1* 1034, (d) *pfmdr1* 1042 and (e) *pfmdr1* 1246. The blue bar is for wild-type allele while the red bar is for the mutant allele.

| Ecological zone | Novel synonymous mutation [frequency in parenthesis] | Novel non-synonymous mutation [frequency in parenthesis] |
|-----------------|-----------------------------------------------------|------------------------------------------------------|
| Coastal         | F106S [1], E236K [1], Y296N [1], E275K [1], E261K [1], S1217Y [1] | G293G [1], G102G [1], G284G [1], I1119I [1], L127L [1] |
| Savannah        | E236K [1], E275K [3], R299K [2], E261K [3] | L108L [1], D117D [1], T1069T [1] |
The cytochrome P450 enzyme family (CYP genes) is a key enzyme involved in the metabolism of different anti-malarial drugs [6]. Lumefantrine is metabolized to desbutyl-benflumetol mainly by CYP3A4 [12]. A change from adenine (A) to guanine (G) at position 392 of CYP3A4 gene proximal promoter region results in CYP3A4*1B allele [24]. This mutant has been reported to have poor enzyme activity [25]. From the results obtained in the current study, 93 individuals were successfully genotyped for CYP3A4 of which 100% had the wild-type gene. This observation suggests that lumefantrine is well metabolized in the participants. Again, delayed clearance observed in patients treated with AL was seen to have one or more mutations in the pfmdr1 gene of the P. falciparum clinical isolates rather than mutation in the CYP3A4 gene of the individuals (Table 2). Based on these observations it can be strongly inferred that the parasite genetic factors could be the driving force behind drug efficiency in the children treated with AL, and this could possibly be the determinant of clinical resistance to the ACT in future. However, there is the need to tread cautiously since this inference is more or less speculative as it is not backed by any pharmacokinetic studies of desbutyl-lumefantrine in the children. It must however be emphasized that similar findings have been reported [26].

The CYP2C8 is the main enzyme that metabolizes amodiaquine to desethyl amodiaquine (DEAQ) [27]. The wild-type CYP2C8*1 and the mutant CYP2C8*2 are the most predominant in Ghana [28]. A change from adenine (A) to thymine (T) at nucleotide position 895 on exon 5 results in the CYP2C8*2 mutant. CYP2C8*2 has been shown to be associated with decreased enzyme activity in vitro and reduced intrinsic clearance of amodiaquine [11]. From the results of the study, 94 individuals were successfully genotyped for CYP2C8 of which 60% (56/94) had wild-type alleles, 35% (33/94) heterozygous and 5% (5/94) homozygous recessive alleles. This result is contrary to what has been reported by Kudzi et al., 2009 [28]. The high number of individuals with wild-type CYP2C8 suggests that amodiaquine was well metabolized in the participants. However delayed clearance was observed
in individuals who reported with high parasitaemia (>100,000) on day 0 and with one or more mutation(s) in the pfmdr1 gene. These individuals had either wild-type or heterozygous CYP2C8 genotype (Table 3) suggesting ample concentration of DEAQ in their plasma. Thus it was expected that their parasites should have been easily cleared. There was no delayed clearance observed in CYP2C8*2 individuals. It is speculated that the CYP2C8 genotype of an individual may not alter the metabolism of the drug significantly, hence the plasma concentration of DEAQ may be adequate to clear the parasite. The absence of delayed clearance in CYP2C8*2 individuals can also be explained by the fact that dihydroartemisinin (DHA), which is a metabolite of artesunate, clears most of the parasites and leaves only a few supposedly ‘weakened parasite’ residues making the presence of a suboptimal concentration of DEAQ enough to clear the parasite residue in these individuals.

When Chi square test was used to determine the association between CYP2C8/CYP3A4 and pfmdr1 genotypes and day 3 positivity, there was no significant difference. For the few cases of delayed parasite clearance using AA, the lack of association between the wild-type enzyme and the cases indicate that the host gene-type of the enzyme could not be responsible for the delayed parasite clearance. Therefore, this observation suggests that the parasite genetic factor among others could be responsible for the delayed clearance rather than the host genetic factors.

| Sample number | CYP3A4 genotype | pfmdr1 mutation(s) |
|---------------|-----------------|-------------------|
| 1             | AA              | F184              |
| 2             | AA              | F184              |
| 3             | AA              | F184              |
| 4             | AA              | F184              |
| 5             | AA              | F184              |
| 6             | AA              | F184              |
| 7             | AA              | F184              |
| 8             | AA              | Y86, F184         |
| 9             | AA              | F184              |
| 10            | AA              | F184              |
| 11            | AA              | F184              |
| 12            | AA              | F184              |
| 13            | AA              | F184              |
| 14            | AA              | F184              |
| 15            | AA              | F184              |
| 16            | AA              | F184              |

Sixteen of the samples with delayed clearance had the CYP3A4 wild-type individuals for lumefantrine metabolism and parasite pfmdr1 mutation(s) as indicated

| Sample number | CYP2C8 genotype | pfmdr1 mutation(s) |
|---------------|-----------------|-------------------|
| 1             | AA              | F184              |
| 2             | AA              | F184              |
| 3             | AT              | F184              |
| 4             | AT              | Y86               |
| 5             | AT              | Y86, F184         |

There were similar numbers of both non-synonymous and synonymous mutations observed at low frequencies in the coastal and forest ecological zones (Table 1). The synonymous mutations may not have any significant effect on the susceptibility of the parasite to the anti-malarial drugs since it does not lead to change in amino acids. However, the novel non-synonymous mutations observed in this study may suggest the possible emergence of new mutations that may lead to reduced parasite susceptibility to ACT in Ghana sooner than later.

**Conclusion**
Observations made in this study give enough grounds to conclude that parasite genetic factors rather than the host’s is more likely to drive resistance to ACT, especially AL in Ghana. All individuals successfully genotyped for CYP3A4 were wild-type, suggesting that lumefantrine is well metabolized in the participants. The high percentage of CYP2C8 wild-type individuals also suggests that amodiaquine is metabolized efficiently. High prevalence of N86, F184 and D1246 suggests AL is less efficacious than AA. The outcome of this study conveys a warning that malaria parasites are becoming resistant to anti-malarial drugs. Prompt monitoring of ACT is required. There is the need to find other anti-malarial drugs that could be used as ACT partner drugs.

**Supplementary information**
Supplementary information accompanies this paper at https://doi.org/10.1186/s12936-020-03320-7.

**Abbreviations**
ACT: Artemisinin-based combination therapy; AA: Artesunate + amodiaquine; AL: Artemether + lumefantrine; CYP: Cytochrome; DEAQ: Desethyl amodiaquine; DHA: Dihydroartemisinin; pfcr: Plasmodium falciparum chloroquine resistance transporter; pfmdr1: Plasmodium falciparum multidrug resistance
1. IRB: Institutional Review Board; NMIMR: Noguchi Memorial Institute for Medical Research; ABC: ATP-binding cassette; PCR: Polymerase chain reaction; WHO: World Health Organization; NFD: N86-F184-D1246 haplotype; YYY: Y86-Y184-Y1246 haplotype.

Acknowledgements
The authors wish to acknowledge the contributions of Dr. Edmund Tetteh Narrey of the Centre for Tropical Clinical Pharmacology and Therapeutics (CTCPT), Mr. Samuel Ahorhorlu (CTCPT), Dr. William Kudzi (CTCPT), Mr. Philip Agyeeman (NMIMR), Miss Selassie Bruku (NMIMR). Nana Aba Enulson (NMIMR), Mr. Kofi Ntimamo (NMIMR).

Authors’ contributions
PH, NDQ, and NBQ conceived and designed the study. PH carried out the molecular genetic studies and led the drafting of the manuscript. NDQ, and NBQ also contributed to drafting of the manuscript. OCH, SM, BA, and KK gave technical support and contributed significantly to drafting of the manuscript. All authors read and approved the final manuscript.

Funding
This research work was funded with grant offered to Peter Hodoameda by West African Centre for Cell Biology of Infectious Pathogens as part of his MPHil Fellowship.

Availability of data and materials
All data generated or analysed during this study are included in this published article.

Ethics approval and consent to participate
Ethical clearance was obtained from the Noguchi Memorial Institute for Medical Research (NMIMR) Institutional Review Board (IRB) and the Ghana Health Service IRB number IRB00001276. Informed consent was obtained from the guardians of the participants. This was after the purpose of the study has been explained to them in their native language and given an opportunity to ask questions that is of concern to them.

Consent for publication
Not applicable.

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

Author details
1 West African Center for Cell Biology of Infectious Pathogens, University of Ghana, P. O. Box LG54, Legon, Ghana. 2 Epidemiology Department, Noguchi Memorial Institute for Medical Research, College of Health Sciences, University of Ghana, P. O. Box LG581, Legon, Ghana. 3 Centre for Tropical Clinical Pharmacology and Therapeutics, University of Ghana Medical School, P. O. Box GP4236, Accra, Ghana. 4 University of Cape Coast Medical School, Cape Coast, Ghana.

Received: 27 April 2020 Accepted: 4 July 2020 Published online: 15 July 2020

References
1. WHO. World malaria report 2018. Geneva: World Health Organization; 2018. www.who.int/malaria. Accessed 15 Nov 2019
2. WHO. World malaria report 2018. Geneva: World Health Organization; 2019. https://www.who.int/malaria/publications/world_malaria_report_2014/mmr-2014-no-profiles.pdf. Accessed 23 Apr 2019
3. Ministry of Health. Antimalarial drug policy for Ghana. Accra: Ministry of Health, 2009.
4. White NJ, Nosten F. Artemisinin-based combination treatment of falciparum malaria. Am J Trop Med Hyg. 2007;77(6 Suppl):181–92.
5. Ojui M, Augureau M-J, Paloque L, Benot-Vical F. Plasmodium falciparum resistance to artemisinin-based combination therapies: a sword of Damocles in the path toward malaria elimination. Parasite. 2018;25:24.
6. Zanger UM, Schwab M. Cytochrome P450 enzymes in drug metabolism: regulation of gene expression, enzyme activities, and impact of genetic variation. Pharmacol Ther. 2013;138:103–41.
7. Sá JM, Tito O, Hayton K, Reyes S, Fay MP, Ringwald P, et al. Geographic patterns of Plasmodium falciparum drug resistance distinguished by differential responses to amodiaquine and chloroquine. Proc Natl Acad Sci USA. 2009;106:18883–9.
8. Siwath C, Ferreira PE, Bustamante LY, Dahlstrom S, Mårtensson A, Björkman A, et al. The role of pfmdr1 in Plasmodium falciparum tolerance to artemether-lumefantrine in Africa. Trop Med Int Health. 2007;12:736–42.
9. Chen N, Chavchich M, Peters JM, Kyle DE, Gatton ML, Cheng Q. Deamplification of pfmdr1-containing amplicon on chromosome 5 in Plasmodium falciparum is associated with reduced resistance to artemisinic acid in vitro. Antimicrob Agents Chemother. 2010;54:3395–401.
10. Helgren G, Hamrin J, Sjövå Mårtensson A, Gill JP, Björkman A. Selection of pfmdr1 mutations after amodiaquine monotherapy and amodiaquine plus artesunate in combination therapy in East Africa. Infect Genet Evol. 2007;7:562–9.
11. Parikh S, Ouedraogo JB, Goldstein JA, Rosenthal PJ, Koetz DL. Amodiaquine metabolism is impaired by common polymorphisms in CYP2C8: implications for malaria treatment in Africa. Clin Pharmacol Ther. 2007;82:197–203.
12. Lefevere G, Thomsen MS. Clinical pharmacokinetics of artemether and lumefantrine (Riamet). Clin Drug Investig. 1999;18:467–80.
13. Wu AHB. Drug metabolizing enzyme activities versus genetic variations for drug of clinical pharmacogenomic relevance. Clin Proteomics. 2011;8:1–9.
14. Ingelman-Sundberg M, Rodriguez-Antonio C. Pharmacogenetics of drug metabolizing enzymes: implications for a safer and more effective drug therapy. Philos Trans R Soc B Biol Sci. 2005;360:1563–70.
15. Abuaiu B, Duah N, Quaye L, Quashe N, Malm K, Plange CB. Therapeutic efficacy of artesunate—amodiaquine and artemether—lumefantrine combinations in the treatment of uncomplicated malaria in two ecological zones in Ghana. Malar J. 2016;15:6.
16. Lorenz TC. Polymerase chain reaction: basic protocol plus troubleshooting and optimization strategies. J Vis Exp. 2012;63:1–14.
17. Vinayak S, Alam T, Sem R, Shah NK, Susanti AI, Lim P, et al. Multiple genetic backgrounds of the amplified Plasmodium falciparum multidrug resistance (pfmdr1) gene and selective sweep of 184F mutation in Cambodia. J Infect Dis. 2010;201:1551–60.
18. Cavaco I, Stromberg-Norlklit J, Kaneko A, Misellem MI, Dahoma M, Ribeiro VL, et al. CYP2C8 polymorphism frequencies among malaria patients in Zanzibar. Eur J Clin Pharmacol. 2005;61:15–8.
19. Hodel EM, Ley SD, Qi W, Avery F, Gerton B, Beck H. A microarray-based system for the simultaneous analysis of single nucleotide polymorphisms in human genes involved in the metabolism of anti-malarial drugs. Malar J. 2009;8:285.
20. Wigginton JE, Cutler DJ, Abecasis GR. A note on exact tests of Hardy Weinberg equilibrium. Am J Hum Genet. 2005;76:887–93.
21. Ferreira PE, Helgren G, Veiga MI, Uhlén P, Kaneko A, Gill JP. PFMDR1: Mechanisms of transport modulation by functional polymorphisms. PLoS ONE. 2011;6:e23875.
22. Bopp S, Magistrado PW, Wong S, Schaffner SF, Mukherjee A, Lim P, et al. Plasmons II-III copy number accounts for bimodal piperaquine resistance among Cambodian Plasmodium falciparum. Nat Commun. 2018;9:1769.
23. Duah NO, Mateevi SA, De Souza DK, Binnah MD, Tandakloe MM, Opoku VS, et al. Increased pfmdr1 gene copy number and the decline in pfcr1 and pfmdr1 resistance alleles in Ghanian Plasmodium falciparum isolates after the change of anti-malarial drug treatment policy. Malar J. 2013;12:377.
24. Lamba JK, Lin YS, Schuetz EG, Thummel KE. Genetic contribution to variation in human CYP3A-mediated metabolism. Adv Drug Deliv Rev. 2012;64:1271–94.
25. Mutagonda RF, Kamuhabwa AA, Minzi OMS, Massawe SN, Asghar M, Homann MW, et al. Effect of pharmacogenetics on plasma lumefantrine pharmacokinetics and malaria treatment outcome in pregnant women. Malar J. 2017;16:267.
26. Kciak K, Rodrigues AS, Rosário V, Gill JP, Lopes D. The drug transporter ABCB1 c.3435C>T SNP influences artemether-lumefantrine treatment outcome. Malar J. 2017;16:383.
27. Li X-Q, Björkman A, Andersson TB, Ridderström M, Masimirembwa CM. Amodiaquine clearance and its metabolism to N-desethylamodiaquine is mediated by CYP2C8: a new high affinity and turnover enzyme-specific probe substrate. J Pharmacol Exp Ther. 2002;300:399–407.

28. Kudzi W, Dodoo ANO, Mills JJ. Characterisation of CYP2C8, CYP2C9 and CYP2C19 polymorphisms in a Ghanaian population. BMC Med Genet. 2009;10:124.

Publisher’s Note
Springer Nature remains neutral with regard to jurisdictional claims in published maps and institutional affiliations.