

Alternatively spliced transcripts and novel pseudogenes of the *Plasmodium falciparum* resistance-associated locus *pfcr*t detected in East African malaria patients

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Objectives: Polymorphisms in the lysosomal transporter encoded by the *pfcr*t gene directly impact on *Plasmodium falciparum* susceptibility to aminoquinolines. The Lys76Thr mutation is the critical change conferring chloroquine resistance *in vitro* and *in vivo*, but always occurs with additional non-synonymous changes in the *pfcr*t coding sequence. We sought to better describe *pfcr*t polymorphisms distal to codon 76.

Methods: Small-volume samples ($\leq 500 \mu\text{L}$) of parasite-infected blood collected directly from malaria patients presenting for treatment in Sudan and Tanzania were immediately preserved for RNA extraction. The *pfcr*t locus was amplified from cDNA preparations by nested PCR and sequenced directly to derive full-length mRNA sequences.

Results: In one of two sites in Sudan, two patients were found with an unorthodox spliced form of *pfcr*t mRNA in which two exons were skipped, but it was not possible to test for the presence of the putative protein products of these aberrant transcripts. Genomic DNA sequencing from dried blood spots collected in parallel confirmed the presence of spliced *pfcr*t pseudogenes in a minority of parasite isolates. Full-length cDNA from conventionally spliced mRNA molecules in all study sites demonstrated the existence of a variety of *pfcr*t haplotypes in East Africa, and thus provides evidence of intragenic recombination.

Conclusions: The presence of pseudogenes, although unlikely to have any direct public health impact, may confound results obtained from simple genotyping methods that consider only codon 76 and the adjacent residues of *pfcr*t.

Keywords: malaria, chloroquine resistance transporters, alternative splicing

Introduction

Resistance of the malaria parasite *Plasmodium falciparum* to antimalarial drugs has posed a serious threat to public health in Africa in the past,¹ and may do so again in the future.² Molecular markers of resistance to chloroquine, pyrimethamine and sulfadoxine were identified and validated some years after the efficacy of these drugs had waned; the utility of markers was thus mainly in gaining understanding of events that had already happened. In the context of artemisinin-based combination therapy (ACT) being the mainstay of uncomplicated malaria management worldwide, malaria researchers have recently focused on the identification of candidate markers for diminished efficacy of artemisinin and of the partner drugs used in ACT.³ Such markers are needed as molecular tools for surveillance for the purpose of preventing the global spread of resistant parasites from foci of origin, and this objective has become more urgent since the recent description in Cambodia and neighbouring countries of parasites with reduced susceptibility to artemisinin compounds.^{4,5}

It is now well established that ACT selects for particular alleles of two genes, *pfmdr1* and *pfcr*, both firmly associated with resistance to the 4-aminoquinolines chloroquine and amodiaquine. In a number of African studies of ACT efficacy in symptomatic children and adults, the aminoquinoline-susceptible haplotypes of these genes were significantly more common in recurrent parasitaemia detected during follow-up after treatment with the most commonly used ACT, artemether/lumefantrine.^{6–9}

Encoding a transporter located in the parasite's digestive vacuole membrane, *pfcr* is an important determinant of susceptibility to antimalarial chemotherapy. Variant genes encoding a threonine instead of a lysine at codon 76 (the K76T mutation) are highly resistant to chloroquine *in vitro* and more likely to be present in parasites surviving chloroquine or amodiaquine treatment *in vivo*,^{10–13} but less likely to survive in ACT-treated malaria patients.^{6–9} The gene encoding the PfCRT protein comprises 13 short exons interrupted by AT-rich introns, imposing technical challenges when sequencing the gene from small-volume, non-replenishable samples. Thus, available sequence data not immediately adjacent to codon 76 have mostly been derived from cDNA obtained from culture-adapted parasite lines.^{14,15} More information on polymorphisms across the full-length *pfcr* gene is therefore needed, particularly from *in vivo* studies of treated malaria patients. In this study transcripts of the *pfcr* gene were isolated from peripheral blood collected directly from symptomatic malaria cases prior to treatment in two East African settings, and characterized by extended cDNA sequencing, in order to reconstruct full genetic haplotypes present *in vivo*. We report for the first time alternatively spliced forms of *pfcr* mRNA, and the presence of genomic intron-free *pfcr* ORF sequences, likely to represent processed—and probably transcribed—pseudogenes.

Materials and methods

Study population

Peripheral blood samples for this study were collected as part of a trial of artemether/lumefantrine efficacy in both adults and children, conducted in New Halfa and Gedarif in eastern Sudan in 2006,⁹ and a pharmacokinetic

study of children aged 1–10 years treated with artemether/lumefantrine for uncomplicated *P. falciparum* infection, performed in Fukayosi, Bagamoyo district, Tanzania in 2006.^{16,17} Details of the study area, study design, patient recruitment criteria, ethical considerations and treatment outcome have been previously reported for both studies.

Sample collection and nucleic acid preparation

For RNA isolation and cDNA synthesis, in both Sudanese studies whole blood was collected in EDTA tubes prior to treatment. A 60 µL aliquot of each sample was mixed with 500 µL of TRI reagent (Sigma) and stored initially at –20°C and later at –80°C once the field work was completed. RNA was extracted by the TRI reagent protocol or the SV total RNA extraction kit (Promega) as per the manufacturers' instructions and DNA removed by digestion with deoxyribonuclease I. cDNA was synthesized employing the Superscript III kit (Invitrogen) as per the manufacturer's recommendations, using random oligonucleotide hexamers to prime the first-strand synthesis. In Fukayosi, from each patient 0.5 mL of whole blood was collected in ribonuclease-free tubes containing 0.5 mL of PBS and 1 mL of 2× Lysis Solution (Applied Biosystems), mixed by shaking and stored directly on ice, then transported to Karolinska Institutet, Sweden on dry ice and stored at –20°C. RNA isolation was performed using the ABI 6100 machine according to the Applied Biosystems protocol for isolation of total RNA from whole blood. Samples were thereafter stored at –80°C. cDNA was synthesized using random hexamers provided in the High Capacity cDNA Reverse Transcription Kit (Applied Biosystems).

For DNA extraction, pretreatment finger-prick blood samples were collected onto glass fibre membranes (Wallac, Finland), dried and stored at 4°C. DNA was extracted by the Chelex[®] method as previously described.⁹

Amplification and sequencing of the pfcr gene

Full-length cDNA and genomic DNA (gDNA) of the *pfcr* gene were amplified in two overlapping PCR amplicons. Fragment 1 [86 nt of the 5' untranslated region (UTR), plus codons 1–247] was amplified with primers CRTP1¹¹ and NG1_REV in the outer reaction and semi-nested with NG4_REV for fragment 1. Fragment 2 (codons 220–424, plus 151 nt of the 3' UTR) was amplified with primers NG5_FOR and NG3_REV for the outer reaction and NG2_FOR and NG3_REV for the nested reaction (Figure 1a). Non-specific and incomplete single-stranded DNA was removed from PCR products using EXO I exonuclease (Fermentas) prior to sequencing on both strands using the amplification primers (ABI BigDye v3 Cycle Terminator kit). In addition, two extra primers (NG6_REV and NG7_FOR; Table 1) were employed at the 5' and 3' terminals, respectively, to sequence the full length of the gene. For gDNA, additional sequencing reactions employing primers P11 (on exon), P13 (crosses intron) and P17 (on intron) were performed. Amplification and sequencing primers, and cycling conditions used, are presented in Table 1. Analysis was performed as previously described.⁹

Results

The full-length cDNA sequence of the *pfcr* gene, assembled from two overlapping amplicons together comprising codons 1–424, was successfully obtained from 26 and 35 of the isolates from eastern Sudan and Bagamoyo district, respectively, while codons 1–220 were successfully sequenced in an additional 23 and 13 isolates, respectively.

Expression of an alternatively spliced variant of pfcr

In two isolates from New Halfa, Sudan (samples 18 and 24), the PCR amplifying cDNA from the 5' segment of the *pfcr* mRNA produced two products by gel electrophoresis: the expected 827 bp

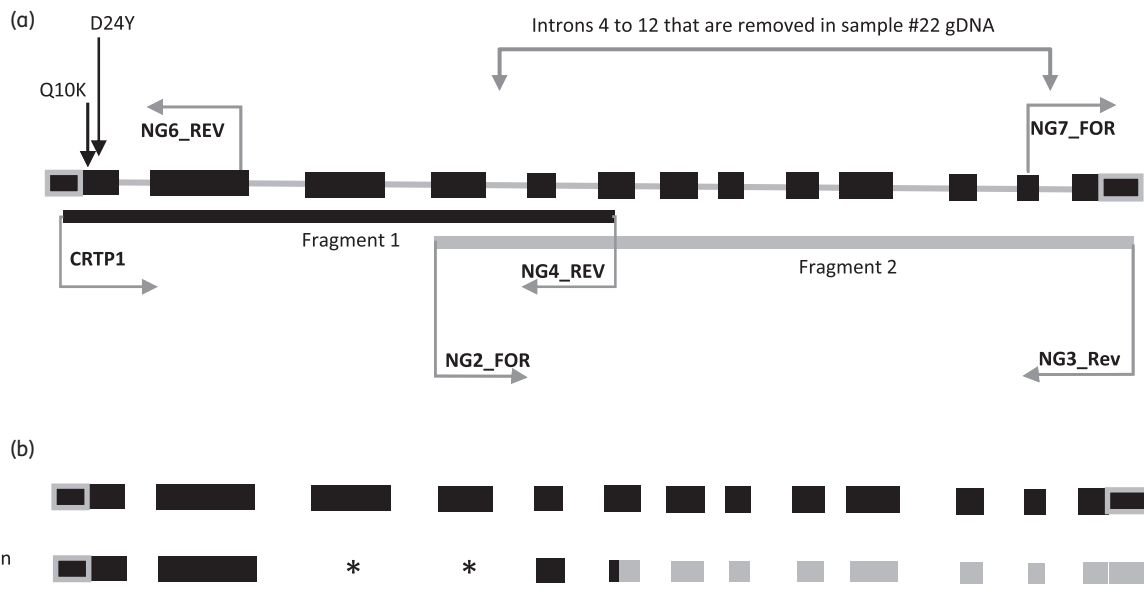


Figure 1. Amplification strategy of the *pfcr1* gene employed in this study. (a) Amplification of the PF3D7_0709000 gDNA of the *pfcr1* gene, exons (black boxes), introns (solid grey lines) and UTR (grey boxes with black fill). Fragment 1 (solid black line) and fragment 2 (solid grey line) are indicated beneath the gene map. PCR primer pairs used to amplify the whole gene (in both gDNA and cDNA) are indicated by horizontal arrows. Primers NG6_REV and NG7_FOR were used to sequence the UTR. Locations of Q10K and D24Y polymorphisms are indicated with arrows. Intron deletion region in gDNA is indicated. (b) cDNA amplified from the *pfcr1* gene. Upper transcript map is the reference PF3D7_0709000 *pfcr1* cDNA. Lower map is the variant *pfcr1* cDNA amplified in sample 24. Black rectangles represent the sequenced transcribed exons. Asterisks represent the non-transcribed (spliced) exons. Grey rectangles represent the putative mRNA downstream from codon 220; this region was not sequenced from cDNA but inferred from gDNA sequence.

Table 1. Primer sequences and PCR cycling conditions for amplifying and sequencing full-length *pfcr1*

Gene	PCR primer sequence 5' to 3'	Cycling conditions	Product size (bp)
<i>pfcr1</i> fragment 1 (exons 1–4)	outer PCR	94°C ³ min, (94°C ¹ min; 45°C ¹ min, 65°C ¹ min)×30 cycles, 65°C ¹⁰ min	873
	CRTP1: CCG TTA ATA ATA AAT ACA CGC AG NG1_REV: GGA TAC CAT AGC ATT TAA TC		
	nested PCR	94°C ³ min, (94°C ¹ min; 50°C ¹ min, 65°C ¹ min)×30 cycles, 65°C ¹⁰ min	827
	CRTP1: CCG TTA ATA ATA AAT ACA CGC AG NG4_REV: AGC ATT TAA TCT TAA AAT GTC		
<i>pfcr1</i> fragment 2 (exons 4–13)	outer PCR	94°C ³ min, (94°C ¹ min; 50°C ¹ min, 65°C ¹ min)×30 cycles, 65°C ¹⁰ min	874
	NG5_FOR: CTC GGA GCA GTT ATT ATT G NG3_Rev: ATT CCT TAT AAA GTG TAA TGC GA		
	nested PCR	94°C ³ min, (94°C ¹ min; 50°C ¹ min, 65°C ¹ min)×30 cycles, 65°C ¹⁰ min	827
	NG2_FOR: GCT CTT GTA GAA ATG AAA TTA TC NG3_REV: ATT CCT TAT AAA GTG TAA TGC GA		
5' and 3' UTR sequencing primers	NG6_REV: GAA CAT AAT CAT ACA AAT A NG7_FOR: CCG TGT AGG AAA TAT TAT CT		
Sequencing primers for confirming intron skipping	P11: TCCTTTTCCAATTGTTCACTTC P13: GACGGAGCATGGGTAAGAAG P17: CAAATGGCTTGTTCCATCAA		

product and an amplicon of ~500 bp. In sample 24 this band was more intense than the 827 bp product, while in sample 18 both bands were of moderate intensity (Figure 2a). Direct sequencing of the ~500 bp PCR product for sample 24 revealed that, for the

majority of cDNA molecules amplified, exons 3 and 4 were completely absent, resulting in a truncated cDNA sequence (521 bp; Figure 1b and Figure 2b). This transcript, if translated, encodes a putative 322 amino acid hypothetical protein, which would be

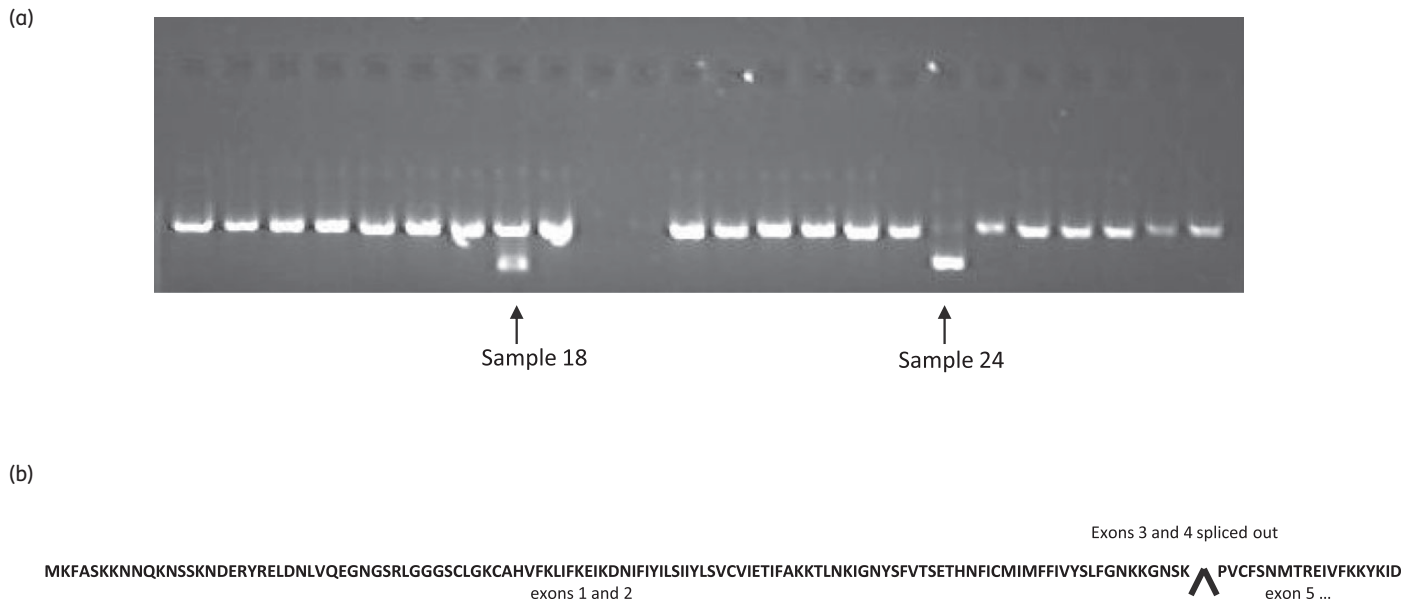


Figure 2. (a) Ethidium-bromide-stained agarose gels of RT-PCR products of N-terminus of *pfcr* mRNA. Lane 1, Hyper Ladder IV (100 bp); lanes 2–24, clinical samples. Arrows indicate samples 18 and 24 in which two RT-PCR products were observed, the expected 827 bp and the 521 bp truncated product. (b) Predicted amino acid sequence of abnormally spliced *pfcr* mRNA from patient 24. The amino acid sequence of exons 1 and 2 is shown. The arrow indicates the splice site at the end of exon 2, followed by the sequence at the beginning of exon 5.

102 amino acids shorter than the canonical *PfCRT* encoded by the 3D7 clone reference (PF3D7_0709000, www.PlasmoDB.org). It is interesting to note that exons 2 and 5 are both type 0 exons, which means that splicing occurs exactly on codon triplet boundaries (i.e. at the third nucleotide), and so the removal of both exons 3 and 4 does not change the reading phase of the codons. Figure 1(b) illustrates the predicted structure of the full-length spliced cDNA. The sequence of *pfcr* mRNA in all remaining patient samples from Gedarif and Fukayosi displayed the expected spliced form. This suggests that the splice variant is not always present.

Intron deletion in genomic copies of *pfcr*

In order to investigate whether the observed splicing reflects rearrangements at the genomic level, and to address whether *pfcr* introns are structured differently in these samples, DNA was extracted from glass fibre membranes by the Chelex method, which does not preserve RNA, from samples 18 and 24, and from three additional randomly selected samples for comparison, 22, 5 and 75 from New Halfa. These were subjected to amplification and sequencing, which was successful only for the 3' region of the *pfcr* gene, coding for the protein carboxy-terminus. Interestingly, amplification produced two PCR products of unexpected sizes: a 600 bp product and an ~450 bp product in samples 18 and 24, and also in one of the comparators, 22.

For each of these isolates, the two PCR products were excised from agarose gels following electrophoresis, purified and sequenced. Four primers (NG2_FOR, P11, P13 and P17) located in this region were employed for sequencing these products. Direct sequencing of the 600 bp PCR product revealed complete removal of introns, with the analysed partial genomic sequence found to be identical to the predicted mRNA sequence. Figure 3 shows the partial *PfCRT* amino acid sequence encoded by an intronless gDNA

segment amplified from samples 18 and 24. Interpretable sequence data were not obtained from the 450 bp amplicon in any of these isolates. Thus some *P. falciparum* isolates from New Halfa harbour additional genome copies of *pfcr* that lack introns. We were not able to demonstrate that these unusual genomic *pfcr* loci were the origin of the unusual transcripts identified in the previous cDNA analysis, and these may be inactive, processed pseudogenes that are not expressed. Alternatively, these may be transcribed by blood stage parasites, giving rise to the unconventionally spliced transcripts observed.

Haplotypes deduced from *pfcr* transcript sequences

cDNA sequence data were used to construct extended haplotypes for the *pfcr* locus in each *P. falciparum* isolate that exhibited the wild-type splicing pattern. Two new polymorphisms were observed in a single isolate from Sudan: the non-synonymous nucleotide change C28A generating a Gln to Lys mutation at codon 10 (Q10K), and the nucleotide change G70T resulting in an Asp to Tyr mutation at codon 24 (D24Y). This codon 24 polymorphism was also observed in three isolates from Tanzania. Using the single-letter amino acid nomenclature, the following non-synonymous *pfcr* mutations were observed in addition to codons 10 and 24: M74I, N75E, K76T, A220S, Q271E, N326S and R371I. (In the following paragraphs, only these variant positions are considered.)

Among the 23 Sudanese isolates where only the 5'-region of the cDNA was fully sequenced, the wild-type haplotype MNKA (at codons 74–76 and 220) was present in 4 (17.4%), and the remaining 19 isolates carried the mutant IETS haplotype, associated with chloroquine resistance in Africa. Of the 26 isolates for which full-length transcript sequences were obtained, chloroquine-resistant haplotypes dominated, with five distinct

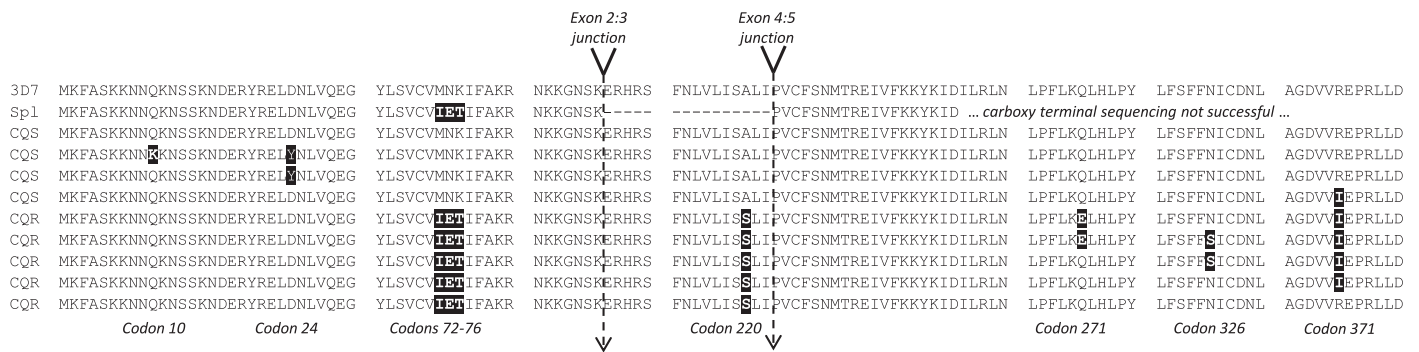


Figure 3. Clustal alignment of predicted amino acid sequences of nine distinct *pfcr1* haplotypes identified by sequencing cDNA generated directly from peripheral blood parasite mRNA. Contributing data were derived from two New Halfa isolates with identical splice variant transcripts (Sp1), 26 full-length cDNA sequences from New Halfa and Gedaref, Sudan, and 35 full-length cDNA sequences from Fukayosi, Tanzania. Haplotype labels indicate predicted chloroquine resistance status (CQS, susceptible; CQR, resistant). Vertical gaps in the alignment indicate invariant sequence tracts not included. Labels under the alignment refer to variant positions (shaded).

full-length haplotypes (Figure 3). A single isolate displayed the reference (wild-type) haplotype MNKAQNR at codons 74–76, 220, 271, 326 and 371. Importantly, two individuals in Sudan harboured parasites with hybrid haplotypes, IETSQNR and IETSQNI, indicative of intragenic recombination within the *pfcr1* locus in this population.

Among the 13 isolates from Fukayosi, Tanzania, for which only the 5'-region was sequenced, the wild-type MNKA at codons 74–76 and 220 was the predominant allele, being present in 7 isolates (53.8%), with 3 (23.1%) carrying the IETS haplotype. Three patients (23.1%) had a mixture of both the IET and MNK, all of whom carried the mutant 220S. A total of 35 isolates were successfully sequenced for the full-length transcript (Figure 3). Two hybrid haplotypes, IETSQNR and MNKAQNI at codons 74–76, 220, 271, 326 and 371, were detected in a single Tanzanian isolate each. Thus in both Sudan and Tanzania, recombinant *pfcr1* alleles comprising sequence elements of both chloroquine-resistant and wild-type *pfcr1* were observed. The prevalence of each haplotype observed in both Sudan and Tanzania is presented in Figure 4.

Discussion

The CRT membrane transporter is an important modulator of parasite response to antimalarial chemotherapy, and yet few studies have addressed the structure or abundance of *pfcr1* transcripts in natural *P. falciparum* infections. Here, we sequenced cDNA amplified from *pfcr1* mRNA present in *P. falciparum*-infected peripheral blood samples taken prior to treatment from malaria patients in eastern Sudan and coastal Tanzania. This approach provided evidence that, *in vivo*, peripheral blood parasites express alternatively spliced *pfcr1* mRNA, the first such demonstration either *in vivo* or *in vitro*, and that naturally occurring parasite genomes can harbour additional *pfcr1* copies in the form of spliced pseudogenes. Although these extra gene copies may be the source of the unorthodox *pfcr1* mRNA molecules we detected, additional studies are needed to confirm a direct relationship. Further, we determined extended *pfcr1* haplotypes, identifying new polymorphisms at codons 10 and 24, and providing the first evidence that chloroquine-resistant and chloroquine-susceptible forms of *pfcr1* recombine in natural parasite populations.

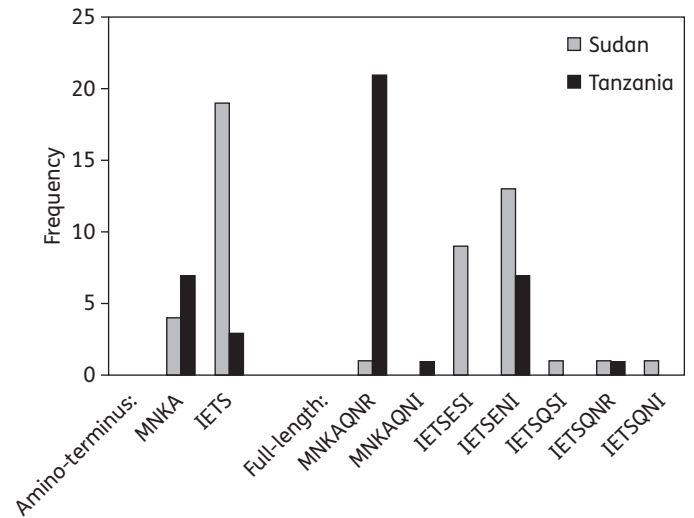


Figure 4. Frequency of *pfcr1* haplotypes in cDNA from isolates in eastern Sudan ($n=49$) and Tanzania ($n=40$). Number of observations of each haplotype defined at codons 74–76 and 220 (amino-terminus and full-length data) and 271, 326 and 371 (full-length data only). Data from isolates with multiple genotypes at the *pfcr1* locus were not included.

The unusual variant transcript of *pfcr1* described here was detected in clinical isolates from only one of two study sites in eastern Sudan, and did not occur among the sequenced Tanzanian isolates. Due to the complete removal of exons 3 and 4 (amino acids 121–222), this transcript encodes a significantly altered putative CRT. This shorter protein will not include the transmembrane domains 3–5, while having a drastically truncated transmembrane domain 6 (Figure 5). Also, this change obliterates three polymorphic amino acid positions (T152A, S163R and the canonical S220A), each probably relevant to CRT function, as they have been associated with different responses of the parasite to several drugs, including chloroquine.^{10,18} However, we have no evidence to date that such a CRT protein is expressed from the alternatively spliced mRNA, and all isolates expressing this variant transcript also expressed the normally

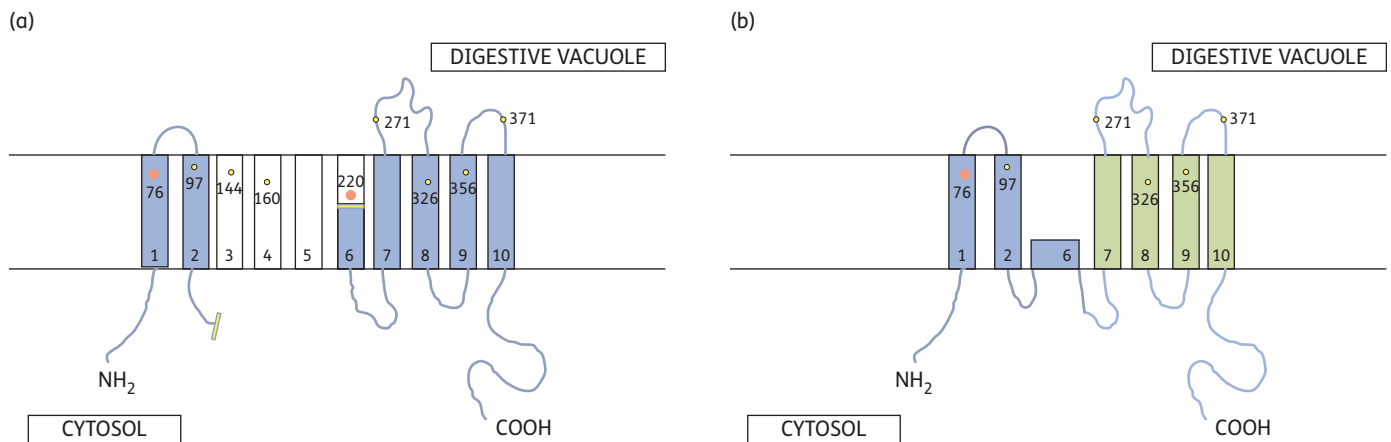


Figure 5. A possible linear arrangement of transmembrane regions in the full-length CRT protein (a) and putative truncated CRT protein encoded by alternative-spliced *pfcr* mRNA from Sudanese malaria patients (b). Transmembrane regions are depicted as rectangles; blue denotes presence in putative truncated form of CRT and white denotes presence only in full-length CRT. Freeform blue line denotes extramembrane polypeptide tracts in the cytosol or inside the digestive vacuole lumen. Transverse yellow bars denote site of splicing event and loss of contiguous nucleotide/amino acid sequence. Polymorphic residues are depicted by yellow or orange (codons 76 and 220) dots. Amino terminal polymorphisms at codons 10 and 24 are not shown.

spliced, full-length *pfcr* transcript. As exons 2 and 5 are spliced precisely in-frame with codon boundaries (known as type 0 splicing), the downstream sequence frame is maintained, such that the alternative splicing event can generate mRNA molecules that encode in-frame putative polypeptides. This also raised the question as to whether the alternative spliced format of *pfcr* and the full-length mRNA were both being expressed from the established *pfcr* locus on chromosome 7 (exemplified by PF3D7_0709000).

Alternative splicing is an important mechanism by which precursor mRNAs are transcribed into several forms of mature mRNAs,^{19,20} leading to multiple polypeptides encoded by a single gene, and the generation of plurifunctional variant protein isoforms.²¹ The first report of a spliced variant mRNA in *P. falciparum* was the 41-3 gene encoding a variable antigen involved in immune regulation and sought as a vaccine candidate.²² Further examples are the guanylyl cyclase gene involved in the signalling pathway of male gametogenesis,²³ the erythrocyte-binding protein MAEBL²⁴ and a CDK-related protein kinase 6.²⁵ Recent transcriptomic studies have identified further variable splicing events for various *P. falciparum* genes in laboratory-cultured clones such as 3D7 and NF54,^{26,27} but there have been no reports of alternative splicing in the *pfcr* gene in laboratory lines or *in vivo*. Interestingly, Otto *et al.*²⁸ described four new splicing events on chromosome 7, but none of these encompassed the *pfcr* locus.

In attempting to understand the genomic origin of the variant cDNA sequences we observed, which corresponded to the 5' half of the normal *pfcr* mRNA, the 3' half of an additional genomic *pfcr* locus in which all introns were missing was identified in parasite gDNA amplified from dried filter-paper blood spots. To our knowledge, this is the first description of pseudogenes derived from drug resistance-associated functional genes in *Plasmodium*. A plausible explanation for our observations, which we were unable to confirm with the material available, is that the intronless *pfcr* locus we have described is a classic pseudogene that has arisen in this parasite population, and is transcribed, as *P. falciparum* pseudogenes frequently are.²⁸ As such it may

have no significant effect on parasite responses to antimalarial drugs, despite being expressed in peripheral blood parasites, although an effect on regulation of transcription from the native *pfcr* locus cannot be ruled out. The unusual cDNA we sequenced encompassed the region of *pfcr* around the codon 76 polymorphism, suggesting the same may be true for the additional *pfcr* pseudogene locus; this is therefore very likely to contribute to genotyping analysis of parasite DNA from peripheral blood. It remains possible that such loci are widespread, and have been part of the 'signal' obtained whenever field samples are genotyped at this locus, whether by PCR-RFLP, qPCR or direct sequencing.^{11,15,29} It is very important that this possible source of confounding of genotyping studies is investigated thoroughly. Interestingly, the pseudogene encodes the mutant 76Thr allele, and so represents an evolutionary event that occurred since chloroquine and related aminoquinolones came into use in the 1940s. It is fascinating to speculate that, as the codon 76 mutation becomes less prevalent due to the withdrawal of chloroquine in Africa, these pseudogenes will remain as genomic 'fossils', which may nevertheless provide a source of the mutant sequence that could be reactivated by recombination should chloroquine pressure be resumed in the future. In this context, it should be noted that evidence suggests that considerable variability was present in the *pfcr* locus before the era of chloroquine exposure.³⁰

CRT codons 72–76 are currently under opposite selection pressure by artemisinins and partner drugs currently employed in ACT.^{8,31} This pressure for parasite survival may select specific adaptive changes in gene structure, as well as changes in the regulation and expression of *pfcr* and other key loci such as *pfmdr1*.^{32–34} This adaptation may include the loss of introns, as reported for fission yeast and mice,³⁵ and the potential gain of new functions. In fact, evidence is gathering that a pseudogene can actually have critical functions, and in some cancers can regulate the 'parent' locus.^{36,37} Finally, our data describe two new polymorphic codons, for amino acids 10 and 24, and provide the first evidence of recombination among chloroquine-resistant and chloroquine-susceptible haplotypes of *pfcr* in both Sudan

and Tanzania. Such events may indicate that *pfcr* genes are still evolving in African parasite populations, generating further heterogeneity in the structure of the CRT protein, and this diversity is likely to be under active selection by ongoing ACT pressure across the continent.

Conclusions

In this study we describe aberrant spliced *pfcr* transcripts and confirm the presence of *pfcr* pseudogenes in a small number of patient parasite isolates. We also provide evidence from both Sudan and Tanzania that hybrid forms of *pfcr* with sequence characteristics of both chloroquine-resistant and wild-type alleles are present in natural populations, probably due to meiotic recombination. A weakness of our study is that the unusual transcripts and alleles of *pfcr* were detected in non-replenishable parasite RNA and DNA, and so confirmatory analyses were not possible. Similarly, no phenotypic analysis of any of the parasite isolates was carried out. However, as the variant alleles of *pfcr* were identified in more than one individual, and from both mRNA and gDNA sequences, we are confident that these parasites were present in New Halfa. Further studies of *pfcr* cDNA sequences from patient peripheral blood are needed to explore this phenomenon further.

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Transparency declarations

None to declare.

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