Abstract. Several single nucleotide polymorphisms (SNPs) have been linked to antimalarial drug resistance in *Plasmodium falciparum*. However, standard polymerase chain reaction (PCR) methods to detect these polymorphisms are unable to detect SNPs in variants representing < 20% of the parasites in a mixed infection, nor can they detect polymorphisms at nearby loci. Here we use heteroduplex tracking assays (HTAs) to analyze *dhps* 540 mutations in 96 samples from Peru and *pfcrt* 76 mutations in 70 samples from China. All samples had been previously analyzed by standard PCR. We detected drug-resistant minority variants and two novel non-synonymous *pfcrt* mutations in China. In Peru, we found no drug-resistant minority variants and a synonymous mutation in *dhps*. Thus, even in regions of low malaria transmission, HTA assays are more informative than PCR with agarose gel electrophoresis.

Drug-resistant *Plasmodium falciparum* malaria is a growing worldwide problem. Molecular markers promise to be important surveillance tools. Single nucleotide polymorphisms (SNPs) in genes such as *dhfr*, *dhps*, *cytb*, and *pfcrt* have been associated with antimalarial resistance in *vitro* and in *vivo*. However, there are two problems with the current methods. First, standard polymerase chain reaction (PCR) methods, such as restriction length polymorphism (RFLP), allele-restricted PCR (AR-PCR), and real-time PCR, can misclassify patients with multicolonial infections because PCR cannot reliably detect variants representing < 20% of the parasite population in a single host (minority variants). Second, these methods are designed to target only specific mutations and consequently do not provide information about unknown or unsuspected mutations in variant alleles. Heteroduplex tracking assays (HTAs) for drug resistance SNPs, on the other hand, are sensitive to minority variants and may detect polymorphisms at positions near known resistance mutations. In this report, we describe a new HTA for *dhps* 540 that is sensitive for minority variants. We use this assay and a previously described HTA for *pfcrt* 76 to evaluate clinical isolates for novel and previously described SNPs. Our findings include the presence of two novel non-synonymous polymorphisms in *pfcrt* and one novel synonymous polymorphism in *dhps*.

Clinical samples obtained from two cohorts were evaluated by HTA. For the *pfcrt* analysis, genomic DNA was used from 70 samples that were collected in 2001 from consenting symptomatic uncomplicated malaria patients in Hainan, People’s Republic of China. For the *dhps* analysis, genomic DNA from 96 consenting patients with symptomatic uncomplicated *P. falciparum* malaria was provided by the US Naval Medical Research Center Detachment, Lima, Peru. A single replicate of the *dhps* 540 HTA was carried out on these samples.

The *pfcrt* 76 HTA was carried out in duplicate on these samples as previously described in the literature. For the *dhps* 540 HTA, PCR amplification of the *dhps* 540 gene was carried out using a Peltier thermal cycler (MJ Research, Waltham, MA) in a volume of 50 µL. The reaction consisted of 5 µL DNA, 1.25 units HotStar Taq DNA polymerase (Qiagen, Valencia, CA), 5 µL 10× PCR buffer, 1 µL 10 mmol/L dNTP mix (Promega, Madison, WI), and 300 nmol/L of forward and reverse primers, which were previously described. This reaction was amplified by preheating to 95°C for 15 minutes, followed by 35 cycles of 95°C for 30 seconds, 50°C for 30 seconds, and 72°C for 1 minute. The reaction was completed with a 10-minute hold at 72°C. The probe was radiolabeled, and HTAs were performed under the conditions noted by Ngrenngarmlert and others with some modifications. An annealing reaction consisting of 8 µL PCR product (either a control or sample DNA) was mixed with 1 µL 10× annealing buffer (1 mol/L NaCl, 100 mmol/L Tris-HCL, pH 7.5, 20 mmol/L EDTA), 2 µL 6× loading dye, 0.5 µL 0.1 mmol/L F primer, and 0.5 µL radiolabeled probe in a total volume of 12 µL. The annealing reaction and electrophoresis were carried out under the conditions previously described. All HTA gels included the following controls: a non-template control (NTC) PCR reaction and PCR reactions from the appropriate wild-type and mutant DNA stocks. Gels were analyzed as previously described.

Malaria DNA samples and controls used to develop assays were acquired from MR4 (www.mr4.org). For *pfcrt* 76 assays, *P. falciparum* strain 3D7 (MRA-102G, from strain MRA-102 deposited by DJ Carucci) was used for wild-type DNA and strain K1 (MRA-159G, from strain MRA-159 deposited by DE Kyle) for mutant DNA. Plasmids (~5,100 bp in size) containing wild-type (MRA-189) and mutant (MRA-190) *dhps* DNA were used for the *dhps* 540 assays.

The sensitivity of the *dhps* 540 HTA to detect minority variants was tested using artificial mixtures of control DNA (wild-type and mutant plasmids) in duplicate as described previously in the literature. The plasmid mixtures were maintained at a total concentration of 1 × 10^7 ng/µL DNA (roughly equivalent to 0.1 ng genomic DNA based on a 22.8-MB genome). Bands were only counted as positive if they were visible to the eye. The HTA probe formed heteroduplexes with different mobilities when annealed to *P. falciparum* DNA amplicons from wild-type parasites and from parasites containing the *dhps* 540 mutation.
resistance mutations of interest. It was possible to accurately and reproducibly detect drug-resistant minority variants comprising as little as 1% of the total population (Table 1).

The pfcrt76 HTA successfully detected DNA in 55/70 (79%) of the P. falciparum isolates from Hainan, China. Because of depleted DNA stocks, repeat evaluation with more sample DNA was not possible. Twenty-two (40%), 28 (51%), and 5 (9%) samples contained pure wild-type, pure mutant, and mixed parasites, respectively, giving a prevalence similar to previous reports. However, among the mixed samples, three samples contained minority-variant drug-resistant parasites that would have been missed by conventional genotyping. However, among the mixed samples, three samples contained minority-variant drug-resistant parasites that would have been missed by conventional genotyping. These represented 11.3%, 3.4%, and 14.4% of the parasite DNA. Thus, 60% of samples would be classified as resistant (pure or mixed mutant) by HTA, whereas only 55% would be expected by ARPCR.

Among these samples, two contained a novel heteroduplex band that migrated more slowly than the typical mutant heteroduplex. Both were mixed infections, also containing a heteroduplex that migrated to the typical mutant location (Figure 1A, Lanes K and L). The novel variant (the higher band in the figure) represented on average 14.4% and 41.7% of the parasite DNA, respectively. To confirm the presence of the variants in the clinical samples, the PCR product was cloned using the TOPO TA Cloning kit (Invitrogen, Carlsbad, CA). The pfcrt-plasmid construct was amplified by colony PCR under the conditions noted for the assay, and the products were sequenced by HTA. The plasmid from colonies containing each variant was purified using the Promega Wizard Miniprep kit (Promega), and the pfcrt insert from the plasmid was sequenced at the UNC Automated Sequencing Facility. Both PCR reactions from each of these two samples and both PCRs from two wild-type samples were evaluated to give four independent sequences per variant. The wild-type sequence matched the 3D7 consensus sequence (CVMNK haplotype), the Chinese consensus (Genbank accession no. XM_001348968.1), the Chinese wild-type variant (containing the S90G polymorphism; Genbank accession no. FJ424266), and the minority mutant variant (containing the NS88 polymorphism; Genbank accession no. FJ424267). Sequence review confirmed this as a novel SNP.

The dhps540 HTA successfully detected 96/96 (100%) of the P. falciparum isolates from Peru. None of these samples contained drug-resistant minority variants. One sample did contain a wild-type minority variant (Figure 1B, Lane P) representing 8% of the parasite DNA. However, the HTA identified four that contained a heteroduplex that migrated more slowly than the typical wild-type variant (Figure 1B, Lane M). Three of the four samples that contained this novel variant were pure. The fourth contained a mixture of novel, representing 7% of the DNA, and typical wild-type variants. Two samples containing pure wild-type, mutant, and pure novel variants were each cloned and sequenced. The wild-type sequence matched the 3D7 consensus sequence. The mutant sequence contained the A to G SNP in the first position of the codon associated with the dhps540E mutation. The novel variant contained a SNP (A to G) in the third position of the
codon and was synonymous to the wild-type (Genbank accession no. FJ424265). Sequence review (PlasmoDB GeneID: PF08_0095) showed this polymorphism to be novel.

This report suggests that surveillance for known drug resistance mutations by HTA may have two advantages: 1) the assays are sensitive to minority variants that may change estimates of the prevalence of a mutation in the population and allow for earlier detection of the emergence of drug resistance,2,3 and 2) it may provide insights into genetic variability in the region closely associated with these loci. This is because HTAs are typically sensitive to SNPs in the region to which the probe binds (typically a 150- to 200-bp region).2,3 We show here that these advantages are preserved over geographically distinct regions and at different genetic loci. The non-synonymous mutations found in pfcrt are particularly interesting because they may represent continued selection caused by drug pressure, because chloroquine is still used to treat P. vivax in this region. However, HTA also has disadvantages. Not all mutations change the motility of the heteroduplex in unique ways. A complete picture of in-host diversity may only be possible using high-throughput sequencing technologies.

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