Genetic Analysis of Primaquine Tolerance in a Patient with Relapsing Vivax Malaria

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Patients with Plasmodium vivax malaria are treated with primaquine to prevent relapse infections. We report primaquine failure in a patient with 3 relapses without any possibility of reinfection. Using whole genome sequencing of the relapsing parasite isolates, we identified single nucleotide variants as candidate molecular markers of resistance.

Of the 5 species of Plasmodium that cause human malaria, P. vivax has the broadest geographic distribution with 2.85 billion persons at risk throughout the world (1). Scientists are becoming increasingly aware of the potential severity of P. vivax infections and their effects on public health (2). A major challenge is the treatment of the dormant stages, hypnozoites, in the liver. Activation of hypnozoites from this reservoir causes subsequent blood-stage infections, or relapses, weeks to years after the primary infection.

Primaquine (PQ) remains the only approved agent to eliminate hypnozoites. Treatment failure, defined by the occurrence of relapses despite PQ therapy, is often ascribed to inadequate dosing, poor adherence, or reinfection (3). However, several cases of PQ tolerance without these confounding factors are reported (4,5). The mechanism underlying PQ tolerance is not understood, although host and parasite genetic factors are implicated. We describe the genetic analysis of parasite and host markers in a patient with 3 P. vivax malaria relapses in a malaria-nonendemic setting where reinfection was not possible.

The Case

The patient is a 38-year-old man from northeast Africa. In December 2008, he experienced a febrile illness in Sudan that was diagnosed as vivax malaria. He was treated with chloroquine (CQ) but did not receive PQ. The patient recovered and moved to Canada in mid-January 2009. One month after his primary infection, he sought treatment at a hospital in Canada with fever, chills, and malaise. P. vivax malaria was diagnosed by microscopy and real-time polymerase chain reaction. He was treated with CQ (600 mg base immediately, 300 mg base at 6, 24, and 48 h), followed by 14 d of PQ (30 mg by mouth daily). His estimated weight was 60 kg. The patient’s symptoms resolved, and smears were negative for Plasmodium on day 16. The patient experienced a second episode of symptomatic P. vivax malaria 3 months later. He was treated with CQ as before, followed by 28 days of PQ (30 mg by mouth daily). Smears were negative 2 days later. Nearly 30 months later, the patient had a third episode of P. vivax malaria. He had not traveled outside North America since his arrival in Canada. He was treated with CQ for 3 d, then PQ for 14 d (30 mg by mouth daily). Smears on days 2 and 9 after CQ treatment were negative. The importance of adherence was emphasized at each clinic visit, and the patient affirms that he took the full course of PQ treatment at the same time every day.

To identify mutations in parasite genes that are potentially associated with primaquine tolerance, we performed whole genome sequencing on P. vivax DNA obtained from patient samples at each relapse (EAC01–03). In total, 55,517 high-confidence single nucleotide variants (SNVs) were genotyped (online Technical Appendix, wwwnc.cdc.gov/EID/article/19/5/12-1852-Techapp1.pdf). The 3 parasite isolates were genetically related, but not identical, and they have been proposed to be meiotic siblings (A.T. Bright et al., unpub. data).

In addition, the 3 strains contained SNVs in genes homologous to known P. falciparum drug-resistance genes, including pvdhps, pvmdr, and pvmrp (6–8). Variants compared to the P. vivax reference strain Sall, presumed to be primaquine sensitive, were found at 27 of 39 sites within 5 known and putative drug resistance genes (Table). All 3 isolates possessed a double mutant antifolate-resistant genotype in pvdhfr (6). The SNVs within the putative drug-resistance genes in each of the patient’s 3 samples were identical except at amino acid positions 976 and 1393 of the pvmdrl gene. The parasite genomes were also

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DOI: http://dx.doi.org/10.3201/eid1905.121852

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compared to the Brazilian strain of *P. vivax*, which was obtained from a patient who had multiple malaria episodes in a malaria-nonendemic country despite primaquine treatment (9). Comparison of the genotypes at the 5 genes demonstrated similar profiles. All strains exhibit intermediate to high levels of antifolate resistance on the basis of mutant genotypes identified in *pvdhfr* and *pvdhps*. In addition, the parasite strains obtained in this study share variant alleles with Brazilian in 2 multidrug resistance–associated transporters, *pvmdr1* and *pvmdr2*. Host pharmacogenetics may also contribute to PQ failure by affecting drug metabolism. Genetic polymorphisms in the *CYP* gene family are associated with poor or intermediate metabolism of many drugs used to treat tropical infections (10) and several of these enzymes are specifically implicated in the metabolism of PQ (11) and other antimalarial drugs (12). We, therefore, determined whether the patient carried alleles that might also explain the failure of treatment. Based on allele frequencies in northeastern African populations, polymorphisms within 4 of the 60 *CYP* genes were selected for genotyping: *CYP1A2*^*1C*, *CYP2B6*^*6*, *CYP3A4*^*1B*, and *CYP2D6*^*4* (online Technical Appendix). The patient was homozygous for the wild-type allele at all 4 loci.

Lastly, we examined whether the patient metabolized PQ to carboxy-primaquine (CPQ), the main PQ metabolite found in plasma. Drug levels were measured with a stereoselective bioanalytical LC-MS/MS method (W. Hanpithakpong et al., unpub. data). A plasma sample was collected on day 12 of treatment of the 3rd relapse, at 12–15 h post-dose. The total PQ and CPQ concentrations were 90 ng/mL and 1,042 ng/mL, respectively. The measured PQ concentration was similar to simulated maximum concentrations at steady-state in healthy male volunteers (96 ng/mL) and patients with vivax malaria (88 ng/mL). Concentration-time profiles for CPQ could not

### Table. Genetic polymorphisms in drug-resistance genes from relapsing isolates of *Plasmodium vivax* *

<table>
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<tr>
<th>Gene</th>
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<th>Polymorphism</th>
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*UTR, untranslated region; ND, not determined.
be simulated because of limited published information. These data demonstrate appropriate absorption of PQ and metabolism into CPQ.

Conclusions

Although this case highlights the challenges in managing patients with *P. vivax* who relapse after high doses of PQ, it also provides a unique opportunity to clarify the mechanisms underlying PQ tolerance. The multiple relapses in this patient result from previously acquired hypnozoites that likely possessed a genetic profile rendering them tolerant to PQ. Genotyping did not identify any mutations within 4 of the *CYP* loci potentially responsible for the antiparasite effect of PQ and plasma measurements demonstrated adequate levels of PQ and CPQ. However, this study presents a limited screen of polymorphisms in the *CYP2D6* gene (13), and we cannot exclude the possibility that other alleles contribute to PQ tolerance.

Parasite genotype data demonstrate that the 3 isolates contain mutations in several putative drug-resistance genes. All 3 isolates are resistant to antifolates and harbor mutations in the ABC transporter genes that are implicated in resistance to numerous antimalarial drugs. Of particular interest are the mutations in the *pvmrp1* gene that encodes a putative multidrug resistance-associated protein. Studies from *P. falciparum* implicate PIMRP1 in glutathione efflux, consistent with the predicted mode of action of PQ in disrupting mitochondrial function (14). Furthermore, gene knockouts of *pfmrp1* have increased sensitivity to several antimalarial drugs, including PQ, which suggests this protein may play a role in transporting antimalarial agents out of the parasite (15).

This case study demonstrates the feasibility of using molecular tools to better understand therapeutic responses to PQ. Genetic analysis of SNVs in putative resistance genes may identify molecular markers of parasite resistance or correlate with known variations in PQ sensitivity of strains from different geographic areas. Clarification of the role of genetic factors involved in PQ efficacy cannot be readily addressed in populations in which endemic transmission occurs because relapses cannot be distinguished from reinfections. Genetic studies of relapses that occur in nontransmission settings provide a unique opportunity to answer questions about this human pathogen.

Acknowledgments

We are grateful to Lilly Miedzinski for detailed clinical information about the patient’s clinical features and course of illness, to J. Kevin Baird for scientific advice and guidance, and to Michael Good for helpful comments on the manuscript.

This work was supported by Alberta Health Services. E.A.W. and A.T.B were supported by National Institutes of Health Grant R21-AI085374-01A1. A.T.B. was supported in part by the UCSD Genetics Training Program through an institutional training grant from the National Institute of General Medical Sciences (T32 GM008666). J.T. and N.J.W. are part of the Wellcome-Trust-Mahidol University-Oxford Tropical Medicine Research Programme supported by the Wellcome Trust of Great Britain.

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References

Genetic Characterization of Primaquine Tolerance


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Genetic Analysis of Primaquine Tolerance in a Patient with Relapsing Vivax Malaria

Technical Appendix

Supplemental Materials and Methods

Ethics Statement

This study was approved by the Health Research Ethics Board of the University of Alberta. The patient provided written informed consent. The consent form states in English that blood samples collected from the patient may be used to genetically characterize the parasites and CYP genes from the patient, and that samples may be shared with other researchers for the purpose of investigating the basis of primaquine (PQ) resistance.

Sample Collection

Whole blood samples from the first (EAC01) and second (EAC02) malaria infections were collected in EDTA tubes and stored at −20°C. For the third infection (EAC03), the red blood cell pellet was stored at −80°C. Plasma was collected on day 12 of PQ treatment.

Genotyping of Cytochrome P450 (CYP) Alleles

Human DNA extraction was performed by using the PSS GC12 instrument (Precision System Science Co. Ltd) and eluted into a 100-µL volume. Four alleles were selected for genotyping analysis based on analysis of the literature taking into account the country of origin of the patient and alleles hypothesized to play a role in drug metabolism: CYP1A2*1C (Dandara, Basvi, Bapiro, Sayi, & Hasler, 2004), CYP2B6*6 (Penzak et al., 2007; Wang & Tompkins, 2008), CYP3A4*1B (Ferreira et al., 2008; Garsa, McLeod, & Marsh, 2005; Kedmi, Maayan, Cohen, Hauzi, & Rund, 2007) and CYP2D6*4 (Xie, Kim, Wood, & Stein, 2001).

Regions within the CYP1A2 and CYP3A4 genes were amplified by PCR followed by DNA sequencing according to published methods (Nakajima et al., 1999; Paganotti et al., 2011). The CYP2B6 allele was characterized by PCR-RFLP using the enzyme BsrI (Ebeshi, Bolaji, & Masimirembwa, 2011). For CYP2D6*4, PCR was performed with the primers: 5′-
CAAGAAGTCGCTGGAGCAGT-3’ (forward) and 5’-GAGGGTCGTCGTACTCGAAG-3’ (reverse) and the following PCR conditions: 94°C for 3 min, 30 cycles of 94°C for 30s, 60°C for 30s, 72°C for 30s, and a final extension step at 72°C for 10 min. PCR products were digested with EcoRII and MvaI analyzed by RFLP. These enzymes will digest the PCR product when there is a guanine at position 1934 but not when there is an adenine, which corresponds to the CYP2D6*4 allele. The presence of the mutant allele was further confirmed by direct sequencing of the PCR product.

**Parasite Genotyping**

For the three parasite samples (EAC01-EAC03), bulk genomic DNA was isolated from frozen whole blood samples using the DNeasy Blood and Tissue kit (Qiagen) as per the manufacturer’s instructions. Whole genome capture of parasite DNA for the three samples was performed as described previously (Bright et al., 2012). Captured DNA was paired-end sequenced on an Illumina HiSeq 2000 for 101 bp per read plus one 7-bp index read using Illumina v.3 chemistry. Data for each sample sequenced in this study is available in the NCBI Sequence Read Archive [SRA057904]. Fastq files obtained from sequencing were aligned to the Sal1 reference using BWA (v. 0.5.9) (Li & Durbin, 2009). Aligned reads were cleaned and analyzed by using Picard (v. 1.51) and GATK (v. 1.6+) (DePristo et al., 2011). 55,517 high confidence SNVs were genotyped in all three samples using GATK (Bright et al., submitted). Heterozygous SNV calls were excluded from downstream analysis.

**Measurement of Drug Levels**

Plasma concentrations of PQ and its major metabolite, carboxy-primaquine (CPQ), were measured with a newly developed stereoselective bioanalytical method (Hanpithakpong et al., manuscript in preparation). In summary, the method used solid-phase extraction followed by liquid chromatography coupled to tandem mass spectrometry. Triplicates of 3 quality control samples were analyzed in the same batch to ensure that accuracy and precision were acceptable according to United States Food and Drug Administration (FDA) standards (FDA Guidance for Industry–Bioanalytical Method Validation). Measured drug concentrations were compared to simulated concentration-time profiles based on literature values for pharmacokinetic parameters in healthy male volunteers (Binh et al., 2009; Cuong et al., 2006; Elmes, Bennett, Abdalla, Carthew, & Edstein, 2006; Fletcher et al., 1981; Mihaly et al., 1985; Mihaly, Ward, Edwards,
Orme, & Breckenridge, 1984) and male patients with vivax malaria (Bangchang, Songsaeng, Thanavibul, Choroenlarp, & Karbwang, 1994.; Bhatia et al., 1986; Kim et al., 2004).

References


