Tafenoquine: A Breakthrough Drug for Radical Cure and Elimination of Malaria

Gokul Gopi1, Surama Manjari Behera2 and Priyamadhaba Behera1*

1All India Institute of Medical Sciences, Bhubaneshwar, India; 2Regional Medical Research Center, Bhubaneswar, India

Abstract

Forty percent of the world’s population is at risk of Plasmodium vivax infection. Relapse is a feature of malaria caused by P. vivax and P. ovale due to the presence of the parasite’s hypnozoite stage that allows it to stay dormant in the human liver. The associated morbidity and economic burden is high, as P. vivax causes severe anemia, miscarriage among pregnant women, malnutrition, and developmental delay in young children due to its chronic relapsing nature. Till recently, for more than 60 years the only licensed antimalarial with proven hypnozoitocidal activity was primaquine. The World Health Organization recommends a regimen of 3-day chloroquine plus 14 days of primaquine for radical cure. Poor adherence to the primaquine course limits its public health benefit on a large scale. Tafenoquine is an 8-aminoquinoline with slower elimination rate, hence a single dose of it is sufficient for hypnozoitocidal activity. Additionally, the schizontocidal activity of tafenoquine makes it a superior drug to the currently available antimalarials, which are mostly single stage specific. Recently, tafenoquine was approved in the USA and Australia for the radical cure of P. vivax malaria in patients aged ≥16 years who are receiving appropriate antimalarial therapy for acute P. vivax malaria, and for the prophylaxis of malaria in patients aged ≥18 years. We have reviewed the available literature of tafenoquine here, and this article explores the possibility of tafenoquine as a key tool for control and elimination of malaria.

Introduction

Globally, in 2017, an estimated 219 million cases of malaria were reported (95% confidence interval [CI]: 203–262 million), compared with 239 million cases in 2010 (95% CI: 219–285 million) and 217 million cases in 2016 (95% CI: 200–259 million). Fifteen countries within sub-Saharan Africa and India carried nearly 80% of the global malaria burden. Among those, five countries accounted for nearly 50% of all malaria cases worldwide: Nigeria (25%); Democratic Republic of the Congo (11%); Mozambique (5%); India (4%); and Uganda (4%).

An increasing number of countries are progressing to elimination, with 19 countries attaining elimination status (zero indigenous cases for 3 years or more) between 2000 and 2017. Although several countries continue to reduce their malaria burden, the rate of reduction has slowed in the highest burden countries; in fact, in some of those countries, malaria cases appear to have risen. With the current global trends being off track for the global technical strategy for malaria 2016–2030 morbidity and mortality targets for 2020, all indications are that the goals are unlikely to be achieved. To get back onto a trajectory that will ensure the achievement of global technical strategy morbidity and mortality milestones for 2025, a response is required to change the current trend in countries that are off track, while sustaining the momentum in those that are on target. This calls for intensified efforts, especially in the highest burden countries.

Until recently, Plasmodium vivax was a relatively neglected pathogen, and research and clinical efforts mainly focused on reducing the mortality associated with Plasmodium falciparum. However, P. vivax is still a major health and economic burden across Asia and Latin America, where the infection is still prevalent. About 82% of estimated vivax malaria cases in 2017 occurred in just five countries (India, Pakistan, Ethiopia, Afghanistan and Indonesia). Unlike P. falciparum, P. vivax and P. ovale have the ability to form hypnozoites that persist in the human liver for variable periods of time and can result in clinical relapses many months or years after the primary infection. Hence, the efforts focused on eradicating malaria need to address this hypnozoite reservoir of P. vivax. A clinical cure of P. vivax can be achieved by clearing the blood-borne pathogen from a patient. However, to achieve a radical cure, in addition to the clinical cure, the patient

Keywords: Tafenoquine; Malaria; Plasmodium vivax; Primaquine.

Abbreviations: CI, confidence interval; CQ, chloroquine; CYP2D6, cytochrome P450 2D6; DETECTIVE, Dose and Efficacy Trial Evaluating Chloroquine and Tafenoquine In Vivax Elimination; GATHER, Global Assessment of Tafenoquine Hemolytic Risk; G6PD, glucose 6 phosphate dehydrogenase; TQ, tafenoquine; WRAIR, Walter Reed Army Institute of Research.

Received: January 27, 2019; Revised: May 08, 2019; Accepted: May 18, 2019

*Correspondence to: Priyamadhaba Behera, Bioinformatics Room, Department of Community and Family Medicine, AIIMS, Bhubaneswar - 751019, Odisha, India. Tel: +91 9910830997; E-mail: priya.madhaba@gmail.com

has to be cleared of the hypnozoite reservoir so as to prevent any future relapses. For more than 60 years, primaquine (PQ) has been the only licensed drug for P. vivax hypnozoite eradication.\textsuperscript{3} The standard regimen for a radical cure treatment of P. vivax infection includes the treatment with artemunate based combination therapy (ACT; including its derivatives) or chloroquine (CQ) for 3 days, along with PQ for 14 days started from day-1 of the treatment. The most significant obstacle to achieving an adequate clinical effectiveness of this regimen is compliance.\textsuperscript{4} Patients often feel asymptomatic within a few days after onset of the treatment, resulting in poor compliance for completing the 14-day course of PQ.

Consider a population as a whole, and this small pool of patients who still carry the dormant parasites act as the pathogen reserve, aiding in the transmission of parasite to other healthy individuals. Level-headed use and forestalling resistance are the key issues with PQ use. PQ has been utilized for over 60 years with foreseen high efficacy in relieving P. vivax infection. Its adequacy has been challenged with accessibility, prescribing practices, and adherence.

Dread of hemolytic potential in glucose 6 phosphate dehydrogenase (G6PD)-deficient people fundamentally diminishes eligibility of PQ to be used as a mass administered drug in public health programs. As indicated by the World Health Organization, antimalarial drug resistance emerges because of spontaneous mutation, and thus is unpredictable. Though the mechanism of resistance to CQ has been widely studied and understood, for 8-aminoquinolone compounds like PQ, such a mechanism has not been identified yet.\textsuperscript{5} There has been recently emerging reports mentioning PQ resistance from various parts of the world. Unfortunately, PQ resistance is regularly confused as treatment failure (relapse occurrence), even after the full course of treatment and at the right therapeutic dose.\textsuperscript{5} Along these lines, defining or confirming genuine PQ resistance is disputable.

Baird and Hoffman\textsuperscript{7} performed and audit and published instances of PQ resistance from over more than 30 years past; they showed that an extensive number PQ treatment failure was seen with patients who took 15 mg/day for 5 days. The standard recommended adult dose of PQ is 15 mg/day for 14 days (210 mg in total). In any case, new studies from a few nations have indicated that this regimen is no longer viable and thus call for higher doses of PQ.\textsuperscript{5,7} Poor compliance, need for increasing drug doses, and rising concerns regarding drug resistance among the malarial parasites have been pushing the scientific community and researchers to search for a drug that could address these concerns and would accelerate the global movement towards eradicating malaria. Tafenoquine (TQ) is a relatively new 8-aminoquinolone, belonging to the same class as PQ with a long elimination half-life (14–28 days versus 4–6 hour for PQ); this long half-life allows infrequent dosing.

**Pharmacokinetics and pharmacodynamics**

TQ is unique in the sense that it has a very long half-life and, like PQ, is active against the preerythrocytic form (liver), the erythrocytic form (sexual), and the gametocytes of the *Plasmodium* species, that includes *P. vivax* and *P. falciparum*.\textsuperscript{11–15} TQ exhibits extensive protein binding of over 99.5% and in a healthy adult has a volume of distribution of ~2,470 L with an interindividual variability of 24.1%. Though its full excretion profile is unknown, metabolism of TQ is slow. With an apparent oral clearance of \( \approx 3 \) L/h, it has an average terminal half-life of \( \approx 15 \) days. While the effect of renal or hepatic impairment on the pharmacokinetics of TQ is still unknown, the pharmacokinetic profile of TQ remains unaffected by age, sex, ethnicity, and bodyweight. Compared with the fasting state, co-administration of TQ with a high-calorie, high-fat meal increased the total exposure to drug by up to 41% and the time to reach peak plasma concentration was increased by approximately one-third.\textsuperscript{14}

Drug interaction studies in healthy volunteers concluded that the pharmacokinetics of TQ were not affected to a clinically relevant extent upon coadministration with CQ.\textsuperscript{16} dihydroartemisinin—
Fig. 1. Important milestones related to Tafenoquine (TQ).
piperazine or artemether–lumefantrine. Correspondingly, TQ did not show a clinically significant effect on the pharmacokinetics of coadministered dihydroartemisinin, piperazine, artemether, or lumefantrine. 

Though the exact mechanism(s) of action responsible for its antiplasmodial activities remain unknown, certain studies involving various protozoan parasites, including P. falciparum, have demonstrated TQ to interfere with mitochondrial functions resulting in an apoptotic-like death of the organism. TQ may also exert its effect by inhibiting hematin polymerization. In addition to its antiplasmodial activities, TQ also causes red cell shrinkage and erytopysis or suicidal erythrocyte death, a process similar to apoptosis in nucleated cells. The activity of TQ targeting the pre-erythrocytic (liver) stages of plasmodium species prevents the development of relapses in P. vivax malaria. A study from Thailand focusing on the transmission blocking potential of TQ, evaluated the efficacy of TQ against the sporogonic stage of the P. vivax parasite after letting mosquitoes feed on gametocytemic blood containing TQ. TQ reduced the transmission of parasite to the mosquito at doses of ≥25 mg/kg.

Like PQ, TQ can induce hemolysis in G6PD-deficient individuals. Drug-induced hemolysis in healthy volunteers with moderately decreased G6PD enzyme activity (40–60% of normal) and who were heterozygotes for the Mahidol487A G6PD deficiency variant showed a linear correlation with increasing single dose of TQ, from 100 mg through 300 mg. However, the hemolytic risk of a single dose TQ 300 mg did not appear to be greater than that with a 14-day course of PQ at 15 mg/day. Among the heterozygous healthy female volunteers with G6PD enzyme activities of 61–80% or >80% of normal, the greatest drop in hemoglobin for TQ was experienced during 300 mg daily dosing. The other coprimary endpoint (the coprimary endpoints) versus 1 of 85 participants (1.2%; 95% CI: 0.2 to 6.4) who received PQ plus CQ. The other coprimary endpoint (the coprimary endpoints) versus 1 of 85 participants (1.2%; 95% CI: 0.2 to 6.4) who received PQ plus CQ.

Recent reports have featured a potential pharmacogenetic impact on the adequacy of PQ in people who are normally deficient in cytochrome P450 2D6 (known as CYP2D6) activity. Both in mice and people, this exploration has given reliable proof that metabolic enactment of PQ by CYP2D6 is required for its activity against hypnozoites, perhaps by means of a toxic metabolite. Further research in mice suggested possible extension of this CYP2D liability to other members of the 8-aminquinolone class, including TQ.

Indications and adverse reactions

The efficacy of TQ has been demonstrated successfully in prophylaxis as well as radical cure (prevention of relapse) of P. vivax malaria. Three key randomized, double-blind, placebo-controlled and/or active referenced, multinational studies have investigated the efficacy of TQ coadministered with CQ as a radical cure for P. vivax malaria; these are the Dose and Efficacy Trial Evaluating CQ and TQ In Vivax Elimination (known as DETECTIVE; NCT01376167) parts 1 and 2 and the Global Assessment of TQ Hemolytic Risk (known as GATHER; NCT02216123).

Another four randomized, double-blinded, placebo-controlled and/or active referenced studies have evaluated the prophylactic efficacy of TQ (200 mg for 3 days followed by weekly 200 mg maintenance doses); one phase III trial conducted on healthy Australian soldiers (nonimmune subjects) deployed in a malaria endemic zone of Timor Leste and three phase II trials conducted on inhabitants of African regions endemic for P. falciparum malaria trials (NCT02488980, NCT02491606, and NCT02488990).

Though TQ was generally well tolerated in the clinical trials, some of the most common adverse events reported were headache, dizziness, nausea, vomiting, and decreased hemoglobin. In the DETECTIVE part 2 trial, the most common (incidence ≥5%) adverse reaction reported prior to day 29 among the TQ plus CQ recipients was dizziness (8% vs. 3% with CQ alone) followed by nausea (6% vs. 7%), vomiting (6% vs. 5%), decreased hemoglobin (5% vs. 2%) and headache (5% vs. 7%). Other adverse reactions, like neuropsychiatric disorders (anxiety, insomnia, abnormal dreams), abnormal blood biochemical panel (raised blood creatinine, increased blood methemoglobin, increased alanine aminotransferase), and eye disorders (photophobia, vortex keratopathy) were reported in ≤3% of clinical trial subjects who received a single dose of 300 mg TQ. The concerns with ophthalmic safety of single-dose 300 mg TQ was further analyzed in a dedicated ophthalmologic study involving nearly 300 healthy volunteers (NCT02658435). There were no reports of vortex keratopathy or retinal abnormalities associated with TQ in this study.

Schmidt et al. have shown that the tissue schizontocidal activity of PQ is a function of total dose rather than of duration of administration, but the toxicity of PQ limits the amount patients may be given in any period of time. Additionally, the fact that the therapeutic dose and the toxic dose are close, leads to serious problems associated with the use of PQ. Clinically important side-effects of PQ include gastrointestinal disturbances, methemoglobinemia, acute intravascular hemolysis in individuals deficient in G6PD enzyme, and possible immunosuppression through inhibition of lymphocyte proliferation.

The main safety concern with 8-aminoquinolines like PQ and TQ is drug-induced hemolysis in patients with G6PD enzyme deficiency. The DETECTIVE trials excluded patients with <70% of normal G6PD activity. The GATHER study assessed the hemolytic potential of TQ (300 mg as single dose), which excluded male patients with <70% of normal G6PD activity and female patients with <40% of normal G6PD activity. Clinically relevant hemolysis was defined as an overall drop in hemoglobin to ≤6.0 g/dL or a decrease in hemoglobin of ≥30% or ≥3 g/dL from baseline at any visit after the first dose of study medication. Only 4 of the 166 participants (2.4%; 95% CI: 0.9 to 6.0) who received TQ plus CQ experienced clinically significant hemolysis up to day 180 (one of the coprimary endpoints) versus 1 of 85 participants (1.2%; 95% CI: 0.2 to 6.4) who received PQ plus CQ. The other coprimary endpoint (prespecified) was the proportion of females who experienced clinically relevant hemolysis up to day 180 who also had moderate G6PD deficiency (G6PD enzyme activity between 40–70%). However, in that event the single patient who was enrolled did not receive TQ.

Public health importance

The current global strategies for eliminating malaria include measures for prevention of malaria, early detection of the disease with early initiation of appropriate treatment, and active surveillance systems. Some of the challenges faced by P. vivax malaria-endemic countries include limited access to effective drugs treating liver stages of the parasite (schizonts and hypnozoites), emergence of drug resistance, and misperception of P. vivax malaria as nonlethal. On average, as shown from experience, the elimination of P. vivax foci can be achieved but not in <3 years, compared with the elimination of P. falciparum, which can be achieved in 1 year. One of the important steps toward malaria elimination is to achieve
TQ is a promising new drug for advancement of the goal towards global malaria elimination. Currently, TQ is approved in the USA and Australia for the radical cure of *P. vivax* malaria in patients aged ≥16 years who are receiving appropriate antimalarial therapy for acute *P. vivax* malaria and for the prophylaxis of malaria in patients aged ≥18 years. Similar efficacy to primaquine with a single dosing regimen, with better compliance, along with the recent development of small portable devices to quantitatively analyze G6PD activity, makes TQ a promising breakthrough drug for radical cure of malaria. Further research about the availability of a low-cost, portable and efficient point-of-care testing platform for detection of G6PD deficiency, as well as more detailed risk versus benefit data regarding the use of TQ, will be vital before its widespread use in public health care.

**Future research directions**

PQ has been in consistent use since 1952 for preventing relapse in patients with *P. vivax* and *P. ovale* malaria. Hemolysis is the major concerning side effect of PQ in individuals who are deficient in G6PD enzyme or its activity. TQ seems to address the problem of compliance due to its single dosing regimen and seems to be an ideal candidate to replace PQ; however, it has similar hemolysis potential as PQ. The available data related to hemolysis and pharmacologic profile of PQ use in community can be utilized for the use of TQ since both the drugs are from the same family and have similar pharmacological properties.

Recently, there have been advancements in the technological community to address the issue of hemolysis and different tools and sensors have been developed to assess the G6PD enzyme activity. Though with emerging technologies and feedback from the use of these devices, the scientific community has been driven to reduce the size and complexity of these devices; yet, they still, by far, are not portable and the operation of these devices requires trained technicians, which largely limits use in community health programs. Recently, a new device—STANDARD™ G6PD—has been developed by a south Korean company (SD BIOSENSOR) with support from PATH. This is a handheld device that delivers results in 2 minutes and provides a quantitative measure of G6PD activity, including in heterozygous women, by using a capillary blood sample and a cartridge similar in format to a glucose meter. According to the company’s description, “It provides a quantitative measurement of both G6PD levels and total hemoglobin, enabling health workers to determine if radical cure with an 8-aminoquinoline-based drug is appropriate for patients.”

The product is currently Conformité Européenne-marked to conform with the European Union In Vitro Diagnostic Medical Devices Directive (98/79/EC), with ongoing clinical evaluation through studies in Brazil, Ethiopia and India and a full clinical evaluation report is expected by mid-2019. Further research about the availability of a low-cost, portable and efficient point-of-care testing platform for detection of G6PD deficiency, better availability and affordability, as well as more detailed risk versus benefit data regarding the use of TQ will be vital before its use in public health care for malaria control.

**Conclusions**

TQ is a promising new drug for advancement of the goal towards...

**Conflict of interest**

The authors declare they have no conflict of interests.

**Author contributions**

Development of concept (PB, SMB and GG), manuscript writing (GG and PB), review of manuscript (SMB, GG and PB).

**References**


DOI: 10.14218/ERHM.2019.00004 | Volume 4 Issue 2, June 2019
Gopi G. et al: Role of tafenoquine in elimination of malaria

Explor Res Hypothesis Med


