Primaquine double dose for 7 days is inferior to single-dose treatment for 14 days in preventing Plasmodium vivax recurrent episodes in Suriname

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Introduction
Worldwide, almost 2.5 billion people are at risk for a Plasmodium vivax infection.1 In the Americas, more than 70% of all malarial infections are caused by P. vivax, and morbidity is high due to frequent recurrent episodes.2 Recent data have shown a steep decline in cases of malaria since 2000, but still more than 148 million people in this region remain at risk for the disease.1,3

Background: Recurrent episodes of Plasmodium vivax are caused by dormant liver stages of the parasite, which are not eradicated by chloroquine. Therefore, effective treatment also includes the use of primaquine (PQ). However, this secondary preventive therapy is often not effective, mostly due to poor adherence to the relatively long treatment course, justifying a comparative study of the efficacy of different durations of PQ treatment.

Materials and methods: We included patients presenting with an acute and documented P. vivax infection from January 2006 to February 2008. All patients received chloroquine 25 mg/kg over a 3-day period. Subsequently, patients in group 7D received PQ 30 mg/day for 7 days, and patients in group 14D received standard PQ 15 mg/day for 14 days. All doses were given under supervision and patients were followed up for at least 6 months. The Kaplan–Meier method was used to estimate cumulative probability of recurrence up to 12 months after treatment initiation stratified by treatment group. Cox regression was used to assess possible determinants for recurrent parasitemia.

Results: Forty-seven of the 79 included patients (59.5%) were allocated to group 7D and 32 patients (40.5%) were allocated to group 14D. Recurrent parasitemia was detected in 31.9% of the cases in group 7D compared to 12.5% of the cases in group 14D (hazard ratio [HR] =3.36, 95% CI 1.11–10.16). Cumulative probability for recurrent parasitemia at 3, 6, and 12 months was 0.201 (95% CI 0.106–0.362), 0.312 (95% CI 0.190–0.485), and 0.424 (95% CI 0.274–0.615) for group 7D and 0.100 (95% CI 0.033–0.279), 0.100 (95% CI 0.033–0.279), and 0.138 (95% CI 0.054–0.327) for group 14D, respectively. When adjusted for possible confounders, differences in recurrent parasitemia remained significant between the two regimens in Cox regression analysis.

Conclusion: More than 30% of the patients receiving shorter treatment course had recurrent parasitemia, suggesting that the standard dose of 15 mg/day PQ for 14 days is more efficacious than 30 mg for 7 days in preventing P. vivax recurrent episodes. Furthermore, we suggest that P. vivax treatment in Suriname should be changed to PQ 30 mg/day for 14 days, as per Center for Disease Control and Prevention recommendation, in light of a recurrence rate of over 10%, even in group 14D.

Keywords: malaria, vivax, recurrent episode, primaquine, treatment duration, efficacy
Among the five species of plasmodium that affect humans, only \textit{P. vivax} and \textit{Plasmodium ovale} have stages (hypnozoites) that remain dormant in the liver cells for lengthy periods of time before reactivating and invading the blood (relapse) weeks to months after the primary infection.\textsuperscript{4} The objective of treating \textit{P. vivax} is to cure both the blood stage as well as the liver stage infection. Primaquine (PQ) is currently the only commercial antimalarial drug that combats hypnozoites in the liver.\textsuperscript{5}

WHO currently recommends chloroquine (CQ) as the first-line treatment for \textit{P. vivax} infection.\textsuperscript{6} In areas with CQ-resistant \textit{P. vivax}, artemisinin-based combination therapies containing piperaquine, mefloquine, or lumefantrine are the recommended treatments.\textsuperscript{6} For the prevention of relapses, PQ is so far the only approved drug. There are several PQ regimens in common use today.\textsuperscript{7} The currently recommended regimen for the cure of \textit{P. vivax} by the WHO is CQ 25 mg base/kg over a 3-day period, followed by PQ 0.25 mg–0.5 mg/kg bodyweight daily over a 14-day period.\textsuperscript{8} Usually, a simplified regimen of CQ 1500 mg over a 3-day period is given, followed by 15 mg of PQ per day for 14 days. Failure of this regimen can be due to a number of reasons, such as CQ resistance,\textsuperscript{8–10} intrinsic tolerance to PQ,\textsuperscript{11,12} reduced absorption, counterfeit drugs, and lack of compliance.\textsuperscript{13} Besides variation in drug susceptibility, factors such as rate and burden of hypnozoite carriage can also impact the disease outcome.\textsuperscript{14}

There is a variance in recurrent episodes geographically and across \textit{P. vivax} strains with the standard regimen.\textsuperscript{3,15} In Colombia, Alvarez et al found a 17.6% recurrence rate when using the standard regimen.\textsuperscript{7} Therapy against the more tropic Chessel strain, frequently found in South-East Asia and Oceania, achieved only a 70%–80% efficacy,\textsuperscript{16,17} suggesting tolerance to the standard PQ therapy in those regions. In support of this theory, studies where a higher total dose was given for 14 days showed a higher efficacy than the standard treatment; therefore, a regimen of 30 mg daily (which corresponds to 0.5 mg/kg/day) for 14 days in those regions is now widely recommended.\textsuperscript{16,18,19}

Unsupervised treatment has a higher rate of relapse than supervised treatment,\textsuperscript{11} suggesting that possible nonadherence leads to reduced effectiveness. Therefore, adherence to the treatment regimen is important in determining the efficacy of the treatment. Clyde et al have shown that 60 mg/day for 7 days is equally effective as 30 mg/day for 14 days,\textsuperscript{20} thereby suggesting that it is the total dose of PQ administered, rather than the duration of treatment, that determines efficacy.\textsuperscript{16,21,22}

Shorter regimens would theoretically improve user compliance and acceptability if the total dose equals the dose proposed by the Center for Disease Control or the WHO.\textsuperscript{13,21} In some countries, such as Brazil, such a shorter, higher dose regimen is being implemented; PQ 0.5 mg/day for 7 days is being used as the first-line treatment for \textit{P. vivax}.\textsuperscript{21} However, because of a higher rate of relapses due to subtherapeutic PQ doses in overweight patients,\textsuperscript{23} weight-adjusted PQ doses are recommended for patients weighing more than 70 kg in Brazil.\textsuperscript{24}

The objective of the present study was to compare the efficacy of a total dose of 3.5 mg/kg in a 7-day regimen of 0.5 mg/kg/day (a maximum of 30 mg per day) and the standard regimen of 0.25 mg/kg/day (maximum of 15 mg per day) for 14 days, with a follow-up period up to a year. Restriction fragment length polymorphism analysis was used after polymerase chain reaction to attempt differentiation between possible new infections and true relapses.\textsuperscript{25}

Materials and methods

The Suriname Ministry of Health’s Ethics Committee (CMWO) granted ethical approval for this study. The study was an open, randomized “en bloque;” prospective clinical controlled trial. The en-blocque method was used for practical reasons, which involved allocating patients from same households to the same group to prevent confusion in treatment among the patients (first 25 patients were allocated to group 1; next two blocks of 25 to group 2, and the following 25 to group 1 again). Initially, patients were recruited from Kwamala Samutu, an Amerindian village in the south of Suriname with high endemicity of \textit{P. vivax} malaria. A sharp decline in the incidence of \textit{P. vivax} in that region and the rest of the interior of Suriname, due to effective prevention strategies, necessitated additional recruitment of patients from the Urban Malaria Center in the capital, Paramaribo. Patients recruited from the Malaria Center usually traveled to regions within the interior often for work or vacation. Since Suriname is a relatively small country and with continuous transmission of \textit{P. vivax} within the interior of Suriname, it is unlikely that \textit{P. vivax} strains will not be equally distributed.

Inclusion criteria were a confirmed \textit{P. vivax} infection (a positive thick blood smear), fever, or a history of fever (≥37.5°C axillary temperature). Patients had to be over 6 months of age, and signed informed consent had to be obtained from all patients, or parents or legal guardians of those under 18 years old. They had to be willing to attend follow-up for at least 6 months, maximum of 12 months. Reasons for exclusion included pregnancy, a history of antimalarial drug intake during the previous 2 weeks, concomitant infection with other plasmodium species,
symptoms of severe malaria, other severe or chronic disease, hypersensitivity to antimalarial treatment, intolerance to the prescribed antimalarial treatment (vomiting), known positive glucose-6-phosphate dehydrogenase deficiency status, and consent withdrawal.

Patients in group 14D received CQ at 25 mg/kg over a period of 3 days, followed by the standard regimen of PQ for *P. vivax*, 0.25 mg/kg/day (maximum of 15 mg/day) for 14 days, as recommended by the WHO. Patients in group 7D received the same regimen of CQ as in group 14D, followed by PQ 0.5 mg/kg/day (maximum of 30 mg/day) for 7 days. PQ doses other than the standard 15 mg dosage were prepared in capsule form by the Diakonessenhuis Hospital Pharmacy of Paramaribo, Suriname. All patients received direct observed treatment. Quality verification of the PQ tablets (Micro Labs Ltd®, Bangalore, India) used during the study was performed on random PQ samples with the GPHF-MiniLab® system.26

At the time of enrolment (day 0), blood was drawn from each patient for thick and thin blood smears, for dried blood spots on filter paper, and for a Binax NOW® Malaria rapid test (Binax, Inc., Portland, ME, USA). A complete clinical evaluation including medical history and physical examination was performed at inclusion. If indicated, patients also had to undergo a pregnancy test. Follow-up blood samples were obtained for thin and thick blood smears on days 7 and 14 and on the first workday of each month consecutively until 1 year after infection. Patients with recurrent parasitemia were treated with the standard regimen of CQ and PQ and were referred to the malaria program for further treatment management. The clinical evaluation, performed at enrolment, was repeated at each follow-up period. Patients who did not attend their scheduled appointment were actively followed up by one of the 12 fieldworkers involved in this study. Kwamala Samutu has a health clinic (medical post) in the village, facilitating the follow-up possibility of the locally recruited patients. In Paramaribo, the fieldworkers were instructed to actively follow up the patients not attending their follow-up schedule. If patients had recurrent parasitemia with *P. vivax* at 1 month, they were considered to have a possible CQ resistant strain27 and therefore not included in the recurrence analysis. Patients using other antimalarial treatment during the follow-up period were excluded as protocol violation.

Recrudescence analysis was performed on paired filter paper strips (Whatman 3 mm) blotted with finger-prick blood from patients with recurring parasitemia. DNA was extracted from the blood spots on each filter using an adapted Saponin-Chelex protocol.28 Primary master PCR amplification was performed for the two genes *pvmsp1F2*, coding for merozoite surface protein1, and *pvmsp3a*, coding for merozoite surface protein 3 alpha (both known to harbor repeat number and sequence polymorphisms), followed by a nested PCR.25,29 Detailed restriction patterns of the genes *pvmsp3a* and *pvmsp1F2* were attained by digesting with two restriction enzymes: AluI/HhaI (*pvmsp3a*) and AluI/MnlI (*pvmsp1F2*). Recurrent infections displaying deviations in one or more of the four restriction patterns were considered heterologous infections, while identical restriction profiles (or mixed patterns containing the initial and another genotype) were designated as homologous infections.

Descriptive statistics, chi-square and Mann–Whitney tests, were used to compare baseline characteristics between the two treatment groups. Kaplan–Meier method was used to estimate cumulative probability of recurrence up to 12 months after treatment initiation stratified by treatment group. Cox proportional hazards regression models were used to assess possible determinants for recurrent *P. vivax* parasitemia. Weight was dichotomized into ≤60 kg versus >60 kg, as treatments are officially weight based, yet simplification of this regimen was based on a person weighing 60 kg. Weight was forcibly entered into the final model to assess as a possible confounder. Analyses were done using SPSS package version 22.0 (SPSS Inc., Chicago, IL, USA) and STATA version 11.2 (College Station, TX, USA).

**Results**

**Patient characteristics**

A total of 85 patients were enrolled from January 2006 till February 2008. Among them, 79 could be included for further analysis: 47 patients in group 7D (59.5%) and 32 patients in group 14D (40.5%). Approximately, 77.0% (47/61) of the patients from recruitment center Paramaribo were allocated to group 7D and 23.0% to group 14D. All patients (n=18) from Kwamala Samutu were allocated to group 14D. Patient characteristics are shown in Table 1. In both groups there were more males than females, 68.1% in group 7D and 63.3% in group 14D (p=0.805). Median age (interquartile range) in groups 7D and 14D was 30.0 years (21.0–37.0) and 21.0 years (13.0–25.0), respectively, (p=0.002). Weight differed between both the groups; median weight in group 7D was 61.5 kg (54.3–70.8) and in group 14D was 59.0 kg (41.0–66.0) (p=0.086).

**Recurrent parasitemia**

In group 7D, 31.9% (n=15) had recurrent parasitemia compared to 12.5% in group 14D (n=4) (p=0.032). Cumulative probability for recurrent parasitemia at 3, 6, and 12 months...
was 0.201 (95% CI 0.106–0.362), 0.312 (95% CI 0.190–0.485), and 0.424 (95% CI 0.274–0.615) for group 7D and 0.100 (95% CI 0.033–0.279), 0.100 (95% CI 0.033–0.279), and 0.138 (95% CI 0.054–0.327) for group 14D, respectively. These results are represented in Figure 1. Analysis showed that the standard simplified regimen 15 mg PQ per day for 14 days and 30 mg/day for 7 days was inadvertently below the recommended 3.5 mg/kg total dose in the majority of patients in both the groups.

### Determinants of recurrent parasitemia

The HR for recurrent parasitemia in the shorter duration treatment group (7D) was significantly higher than the longer treatment duration group (14D) (HR = 3.4; 95% CI 1.1–10.2). Sex, age, and weight were not associated with parasitemia recurrence (Table 2). Adjusting for weight in the final model, only shorter duration treatment remained significantly associated with parasitemia recurrence (HR = 5.2; 95% CI 1.5–18.1).

### Recrudescence analysis

Recrudescent analysis could be performed on 14 of the 19 recurrent infections. In group 7D, 54.5% (6/11) of the recurrent infections were homologous infections when compared to the parasite present at inclusion. In group 14D, 100% (3/3) of the infections were homologous (p=0.231). The longest relapse period observed for the homologous infections was 7 months.

Statistical analysis of the efficacy of the two regimens taking homologous and heterologous infections into account was not possible, due to the limited numbers in each category and the fact that heterologous cases considered as new infections can also be true relapses displaying genetic differences originating from the hypnozoites.

### Discussion

We found a higher incidence of recurrent parasitemia in group 7D (PQ 30 mg/day for 7 days) than in group 14D (15 mg PQ daily for 14 days), p<0.05. There are numerous studies evaluating different treatment regimens. For decades, the generally accepted view has been that the total dose, rather than the duration of treatment, influenced the prevention of recurrent episodes, which was largely based on a preclinical study in 1977 by Schmidt et al and a clinical study by Clyde et al. The latter study showed an effective prevention of recurrences in patients using 60 mg/day over a period of 7 days. The follow-up period was adequate, and 8 of the 11 patients were followed up for at least 7 months, yet the number of patients in that specific study was small (n=11). A more recent study by Krudssoord et al compared a regimen comprising 30 mg bid for 7 days to a regimen comprising 30 mg qd for 11 and 14 days. There was no difference between the two groups, thus supporting the total dose theory. However, their follow-up period was a mere 30 days, excluding late recurrences. On the other hand, a study conducted in Peru reported similar recurrence rates in groups with a 7-day regimen of 0.5 mg/kg/day and with 0.25 mg/kg/day (maximum of 15 mg per day) for 14 days. There was no difference between the two groups, thus supporting the total dose theory. However, their follow-up period was a mere 30 days, excluding late recurrences. On the other hand, a study conducted in Peru reported similar recurrence rates in groups with a 7-day regimen of 0.5 mg/kg/day and with 0.25 mg/kg/day (maximum of 15 mg per day) for 14 days. A study by Alving et al and a more recent study by Leslie et al showed that a regimen of a once-weekly dose of 45 mg/week and 0.8 mg/kg/week of PQ, respectively, for 8 weeks was just as effective for radical cure of *P. vivax* as 0.5 mg/kg/day for 14 days. Our results are in concordance with a study by Rajgor et al from India in 2003, which demonstrates higher recurrence and relapse rates in the group with 7 days double dose as compared to 14 days single dose. A Cochrane database systematic review published in 2013 did not find enough evidence to promote a different regimen than single-dose PQ for 14 days. Our findings add to the data, which fuel the debate about the most efficacious PQ regimen to prevent recurrence.
Our study has some limitations. First of all, the study findings date from 2006 to 2008, but the data are still relevant since PQ is so far the only approved drug for the prevention of *P. vivax* relapses, and several countries, including our neighbor Brazil, still use the short regimen. Second, the number of patients in both groups is small. This study was initiated in the period in which the number of malaria cases started to drastically decline due to novel control strategies. Also, the randomization was weakened, since the local circumstances in the village Kwamala Samutu dictated the use of one treatment within the same “house” in the same period, but fortunately, this did not significantly influence the outcome. Furthermore, the number of recurrent episodes was low, therefore complicating comparisons, besides the inadequacy of current molecular methods to distinguish between a true recrudescence of the initial infection or a relapse from hypnozoites generated from the prior blood stage infection. Last, the therapeutic dose in both the groups was suboptimal if the actual bodyweight was taken into account. However, statistical analysis showed that this did not influence recurrent parasitemia cases.

### Conclusion

In summary, general analysis showed a significantly higher incidence of recurrent parasitemia in group 7D (>0.05), suggesting that the duration of the treatment is, in fact, more important than the total dose. Heavier weight, resulting in possible subtherapeutic doses, was not identified as a confounder. Since the recurrence of parasitemia was higher than 10% even in group 14D and the majority of patients in our study received subtherapeutic doses, replacing the current standard dose of 15 mg per day for adults by an accurately calculated dose of 15 mg/kg of bodyweight or prescribing the 30 mg/day regimen, already implemented in Thailand and other countries in South-East Asia, is recommended.

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### Disclosure

The authors report no conflicts of interest in this work.

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