A Phase III, Randomized, Non-Inferiority Trial to Assess the Efficacy and Safety of Dihydroartemisinin-Piperaquine in Comparison with Artesunate-Mefloquine in Patients with Uncomplicated Plasmodium falciparum Malaria in Southern Laos

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Abstract. We conducted an open, randomized clinical trial of oral dihydroartemisinin-piperaquine (DP) versus artesunate-mefloquine (AM) in 300 patients in Laos with uncomplicated Plasmodium falciparum malaria as part of a multicentre study in Asia. Survival analysis and adjustment for re-infection showed that the 63-day cure rates (95% confidence interval [CI]) were 100% for AM and 99.5% (96.4–99.8%) for DP. The 63-day cure rates per protocol were 99% (97 of 98) for AM and 99.5% (196 of 197) for DP (P = 0.55). The difference (AM minus DP) in cure rates (95% CI) was −0.5% (−5.1 to 2.0%), which is within the 5% non-inferiority margin. The median fever and parasite clearance times were also similar for AM and DP. The proportion of patients with at least one recorded potential adverse event was significantly higher in the AM group (38 of 87, 44%) than in the DP group (57 of 182, 31%) (relative risk = 0.6, 95% CI = 0.4–0.9; P = 0.04). Dihydroartemisinin-piperaquine is not inferior to AM in the treatment of uncomplicated P. falciparum malaria in Laos and is associated with fewer adverse effects. The results of this study were similar to those of the larger multicentre study.

INTRODUCTION

Malaria remains an important public health challenge in southern Laos with a median incidence of Plasmodium falciparum infection of 4.7–23.5/1,000 population. As in many other tropical countries, antimalarial drug resistance to P. falciparum poses a public health threat. The Lao Government changed national policy for first-line antimalarial drug treatment of uncomplicated P. falciparum malaria to artemether-lumefantrine (AL) in 2005. This artemisinin-combination treatment (ACT) and artesunate-mefloquine both have high efficacies and good tolerability with 42-day failure rates of ≤ 6% in Laos. However, there is uncertainty as to the clinical importance of reduced bioavailability of lumefantrine when taken without fatty food, especially with evidence of resistance to artesinin derivatives in adjacent Cambodia increasing the required contribution to efficacy by the partner drug. Artemether-lumefantrine has to be taken twice a day, increasing the required contribution to efficacy by the partner drug. Dihydroartemisinin-piperaquine (DP) is a potential alternative; it can be taken once a day, does not have food dependence, and can be produced for a lower cost than AL. A trial comparing DP (Artekina®; Holleykin Pharmaceutical Co., Guangzhou, China) with artesunate (Guilin Pharmaceutical Co., Guilin, China) plus mefloquine (Lariam®; Roche, Basel, Switzerland) in southern Laos demonstrated that DP had a 100% cure rate (n = 110) assessed at 42 days-follow up. The DP used in this study was produced in compliance with Chinese Good Manufacturing Practice (GMP) standards, but not with the GMP standards required by drug regulatory authorities in Europe or the United States. A DP formulation produced to standards compliant with International Conference on Harmonization GMP has been developed by Sigma-Tau (Rome, Italy) with support from the Medicines for Malaria Venture (MMV).

We therefore conducted a phase III, randomized, non-inferiority trial of this new formulation to assess the efficacy and safety in comparison with artesunate-mefloquine (AM) in patients with acute, uncomplicated P. falciparum malaria in Laos. This study was part of a multi-center study in Asia that included India, Laos, and Thailand. This paper reports the detailed results from Laos and their implications.

MATERIALS AND METHODS

Study site, patients, clinical procedures, and laboratory investigation. The study was conducted during June–October in 2005 and 2006 at Phalanxay (10 beds) and Xepon (30 beds) District Clinics and during June–October 2006 at Xepon District Clinic, Savannakhet Province. Phalanxay and Xepon districts (Savannakhet Province) are located 605 km and 665 km, respectively, southeast of Vientiane, the capital of Laos. Phalanxay (78 villages, population = 24,730) and Xepon (88 villages, population = 48,000) are inhabited predominantly by rice farmers of the Lao Theung ethnic group. A sample size calculation was performed for the whole multi-center study. For Laos, 300 patients were assigned to be recruited.

Patient inclusion and exclusion criteria have been reported. At presentation, venous blood samples were obtained for parasite count, hematologic tests, and biochemical tests; three blood spots were collected on 3MM filter paper (Whatman, Maidstone, United Kingdom) for genotyping by polymerase chain reaction (PCR) in the event of reappearance of P. falciparum during follow-up. A 12-lead electrocardiogram (ECG) was obtained.

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If the study inclusion criteria were met, the patients were randomly assigned to receive either 1) AM: artesunate (Artesunate, 100 oblong Lactab; Mepha, Aesch-Basel, Switzerland), 4 mg/kg/day for 3 days (days 0–2) plus mefloquine (Mephaquin Lactab; Mepha, Aesch-Basel), 15 mg base/kg on day 1 and 10 mg base/kg on day 2 or 2) DP: dihydroartemisinin-piperaquine (Sigma-Tau), 2.1/16.8 mg/kg in a single daily dose for 3 days. Each tablet contained 40 mg of dihydroartemisinin and 320 mg of piperaquine for adults and 20 mg of dihydroartemisinin and 160 mg of piperaquine for children <15 years of age. The use of AM and DP in this study was reviewed and approved by the Food and Drug Department, Ministry of Health, Laos.

The treatment choice was kept in a sealed opaque envelope, which was opened only after the decision to recruit had been made. An unequal randomization, 2:1 (DP:AM), was used to provide more precise estimates of DP cure rates and to provide more patients for the safety database of DP. Axillary temperature was measured every six hours. Study drug administration was observed directly by the study physicians. Study medications administered to older children (>5 years of age) and adults were given as tablets or fractions of tablets orally with a glass of water and those given to young children (<5 years of age) were crushed, mixed with water, and administered as a slurry if the children were unable to swallow. Patients were observed for one hour to ensure that the medications were not vomited or regurgitated.

Patients were reviewed daily until parasite clearance was observed, then weekly for 63 days from the start of treatment, or at other times if he or she was ill. At each visit, finger prick blood was obtained for a malaria blood smear and hematocrit. Three blood spots were collected on filter paper from those with reappearance of asexual parasitemia for genotyping. Twelve-lead ECGs were obtained on days 2 and 7 and on days 28 and 63 if the ECG result was abnormal on day 7; ECGs were also obtained on the day of \( P. falciparum \) recurrence. Venous blood samples were obtained for hematologic and biochemical tests on days 28 and 63 if the results were abnormal on day 28 and were also performed on the day of \( P. falciparum \) recurrence.

Patients who received AM and had recurrent \( P. falciparum \) parasitemia or treatment failure were withdrawn from the study, re-treated with oral artesunate (2 mg/kg/day) plus doxycycline (4 mg/kg/day) (or artesunate alone for children <8 years of age) for 7 days and followed-up. Those persons who received DP and had recurrent \( P. falciparum \) parasitemia or treatment failure were withdrawn from the study, re-treated with AM for three days, and followed-up. Patients in whom severe disease developed or required rescue therapy were re-treated with artesunate (2 mg/kg/day) plus mefloquine after the axillary temperature first decreased below 37.5°C and remained below 37.5°C (48 hours), 7) gametocyte carriage (blood slide positive for gametocytes) after treatment, 8) fractional changes in hematocrit after antimalarial treatment, and 9) adverse events.

Statistical analysis. Data were analyzed by using SPSS version 11.0 (SPSS, Chicago, IL). Comparisons between two groups were made by using the Mann-Whitney U test, the Student’s t-test, the chi-square test, and Fisher’s exact tests, as appropriate. Cure rates were calculated as the proportion of patients with PCR-confirmed recrudescence by using intention-to-treat (ITT) and per-protocol (PP) populations. In the ITT population, all losses to follow-up were treated as failures. In the PP population, losses to follow-up were excluded from the analysis. Patients with new infections were regarded as cures in both analyses. Survival analysis (Kaplan-Meier estimates) was used to calculate cure rates by using StatXact version 9 (StatSoftCorp, College Station, TX) for all randomized patients.

Results. The numbers of patients screened for malaria and enrolled in the trial is shown in Figure 1. The admission features of the two study groups were similar (Table 1). At the time of enrollment, 129 (43%) patients were afebrile but had a history of fever prior to admission. The proportion of patients without documented fever on admission was similar in the AM and DP groups. Most patients (68%, 205 of 300) were children (≤15 years of age) (69 of 98 in AM group and 136 of 202 in the DP group). Two patients in the DP group refused to swallow the study tablets, and three patients had persistent vomiting after taking study drugs, and thus were excluded from
the study and were not followed-up. Therefore, 98 (100%) and 197 (97.5%) patients in AM and DP groups, respectively, completed 63 days of follow-up (Figure 1).

Cure rates, fever and parasite clearance, and changes in hematocrit and hemoglobin levels. Severe disease developed in a four-year-old girl who had uncomplicated *P. falciparum* malaria (97,968 parasites/μL). She had one 5-minute convulsion 30 hours after receiving AM and remained unconscious (Glasgow Coma Score $\geq 9/15$) for 70 minutes. She recovered after receiving supportive treatment. This patient was not excluded from the study and completed 63 days of follow-up and showed cure.

Of 8 and 12 patients with subsequent *P. falciparum* reappearance in the AM and DP groups, respectively, PCR analysis indicated that only one patient, in the DP group, had a recrudescence infection. If we considered the four-year-old patient with the convulsion and coma after treatment as an early treatment failure, the 63-day cure rates per protocol, excluding patients who refused to swallow tablets or had persistent vomiting, or experienced re-infection, were 99% (97 of 98) for AM and 97% (196 of 202) for DP ($P = 0.43$). Similarly, 28-day cure rates by intention-to-treat analysis were 99% (97 of 98) for AM and 97% (196 of 202) for DP ($P = 0.43$). Conventionally intention-to-treat analysis showed that the 63-day cure rates adjusted for re-infection were 99% (97 of 98) for AM and 97% (196 of 202) for DP ($P = 0.43$). Similarly, 28-day cure rates by intention-to-treat analysis were 99% (97 of 98) for AM and 97% (196 of 202) for DP ($P = 0.43$). The number of patients with *P. vivax* appearance in the AM and DP groups during follow-up was similar (Table 2). At presentation, 57% of all patients had fever, as defined by a body temperature $\geq 37.5^\circ$C. At day 2, 99% and 97.5% in AM and DP groups, respectively, were afebrile ($P = 0.67$), and all patients (except one in the DP group) had no parasites detected. The median fever and parasite clearance times were similar between the AM and DP groups (Table 2).

At presentation, 57% of all patients had fever, as defined by a body temperature $\geq 37.5^\circ$C. At day 2, 99% and 97.5% in AM and DP groups, respectively, were afebrile ($P = 0.67$), and all patients (except one in the DP group) had no parasites detected. The median fever and parasite clearance times were similar between the AM and DP groups (Table 2).

The mean hematocrit on day 7 was significantly higher in the DP group than in the AM group ($P = 0.02$). Mean leukocyte counts were significantly higher on day 28 than on day 0 in both treatment groups ($P < 0.001$).
### Table 1

Admission demographic, clinical, and laboratory details for patients in a study comparing AM and DP for treatment of *Plasmodium falciparum* malaria in Laos

| Variable | All (n = 300) | AM (n = 98) | DP (n = 202) |
|----------|--------------|-------------|--------------|
| Sex, Male, no. (%) | 179 (60) | 59 (60) | 120 (59) |
| Age, years | 14.4 (12.8–16.0) | 14.1 (11.2–17.0) | 14.6 (12.7–16.5) |
| Body weight, kg | 29.3 (27.2–31.4) | 28.2 (24.5–31.9) | 29.8 (27.2–32.4) |
| Previous malaria attack, no. (%) of patients† | 99 (33) | 34 (35) | 65 (32) |
| Axillary temperature, °C | 37.9 (37.7–38.0) | 37.9 (37.7–38.2) | 37.9 (37.7–38.0) |
| Patients without fever on admission, no. (%) | 129 (43) | 39 (40) | 90 (44.5) |
| Systolic blood pressure, mm Hg‡ | 103.3 (101.8–104.9) | 104.5 (101.6–107.4) | 102.8 (100.9–104.6) |
| Diastolic blood pressure, mm Hg‡ | 64.5 (63.5–65.7) | 65.0 (62.8–67.3) | 64.2 (62.5–65.7) |
| Pulse, beats/min | 109.6 (107.1–112.0) | 110.2 (105.6–114.9) | 109.2 (106.3–112.2) |
| Splenomegaly, no. (%) of patients | 100 (33) | 37 (38) | 63 (31) |
| Hepatomegaly, no. (%) of patients | 54 (18) | 21 (21) | 33 (16) |
| Parasitemia, geometric mean parasites/μL | 20,564 (17,873–23,659) | 22,851 (18,433–28,333) | 18,505 (15,438–22,182) |
| Erythrocytes/mm³ | 4.91 (4.81–5.02) | 4.79 (4.62–4.96) | 4.97 (4.85–5.10) |
| Lymphocytes, % | 34.5 (33.2–35.8) | 33.4 (31.2–35.7) | 35.1 (33.4–36.7) |
| PMN, % | 61.8 (60.4–63.2) | 63.6 (61.2–65.9) | 60.9 (59.2–62.7) |
| Hemoglobin, g/dL | 11.4 (11.1–11.7) | 11.2 (10.7–11.7) | 11.5 (11.1–11.8) |
| Hematocrit, % | 36.6 (35.8–37.4) | 35.9 (34.6–37.3) | 36.9 (35.9–38.0) |
| Leukocytes/mm³ | 6,670 (6,322–7,035) | 6,754 (6,237–7,270) | 6,642 (6,173–7,112) |
| PMN = polymorphonuclear leukocytes; ALT = alanine aminotransferase; AST = aspartate aminotransferase, GGT = γ-glutamyl transferase. \* Values are presented as mean (95% confidence intervals) unless otherwise indicated. AM = mefloquine plus artesunate for 3 days; DP = dihydroartemisinin plus piperaquine for 3 days; PMN = polymorphonuclear leukocytes; ALT = alanine aminotransferase; AST = aspartate aminotransferase, GGT = γ-glutamyl transferase. † Defined as patient or patient’s guardian reporting that the patient had had a febrile illness with a positive malaria slide. § Data were available from only 201 patients in DP group. ¶ Significant difference from day 28. # Significant difference between AM and DP groups (P < 0.01).

### Comparisons between adults and children. The mean admission temperature was significantly higher in children (38.0°C [37.8–38.1°C]) than adults (37.7°C [37.4–37.9°C]) (P = 0.04). A palpable spleen (47% and 4%; P < 0.001) and liver (25% and 2%, P < 0.001) were significantly more frequent in children than adults. The admission geometric mean parasitemia/μL was significantly higher in children than adults (13,718 parasites/μL [10,578–17,795 parasites/μL] versus 6,471 parasites/μL [4,578–9,147 parasites/μL]) (P = 0.001), as were the proportions of patients with clinical anemia and vomiting at presentation (41 [20%] of 205 versus 1 [1%] of 95; P < 0.001 and 94 [46%] of 205 versus 29 [30.5%] of 95; P < 0.001, respectively). The frequency of the following admission symptoms and signs were significantly lower in children than adults:

### Table 2

Outcome measures for the treatment of patients enrolled in a study comparing AM and DP for treatment of *Plasmodium falciparum* malaria in Laos

| Variable | All (n = 300) | AM (n = 98) | DP (n = 202) |
|----------|--------------|-------------|--------------|
| 63-day cure rate, no. (%) of patients† | – | 97/98 (99) | 196/202 (97) |
| 63-day cure rate per protocol, no. (%) of patients | – | 97/98 (99%) | 196/197 (99.5) |
| 63-day cure rate by survival analysis, % (95% CI) | – | 100 | 99.5 (96.4–99.8) |
| Fever clearance time, median hours (range)‡ | 22 (7–68) | 24 (7–68) | 21 (7–64) |
| Parasite clearance time, median days (range)§ | 2 (1–3) | 2 (1–2) | 2 (1–3) |
| Positive parasitemia at day 1, no. (%) of patients | 199/298 (67) | 70/98 (71) | 129/200 (64.5) |
| Positive parasitemia at day 2, no. (%) of patients | 1/298 (0.3) | 0 | 1/200 (0.5%) |
| Gametocytemia after treatment, no. (%) of patients | 7/300 (2.3) | 1/198 (1%) | 6/202 (3%) |
| P vivax appearance after treatment of *P. falciparum*, no. (%) of patients | 10/295 (3) | 2/298 (6) | 4/197 (2) |
| Day 0 hematocrit, mean % (95% CI) | 36.6 (35.8–37.4)¶ | 35.9 (34.6–37.3)¶ | 36.9 (35.9–38.0)¶ |
| Day 7 hematocrit, mean % (95% CI)¶ | 32.6 (32.0–33.3) | 31.6 (30.5–32.7) | 33.1 (32.4–34.0) |
| Day 28 hematocrit, mean % (95% CI)¶ | 38.6 (38.2–39.1) | 38.2 (37.4–39.0) | 38.8 (38.2–39.5) |

*AM = mefloquine plus artesunate for 3 days; DP = dihydroartemisinin plus piperaquine for 3 days. † Intention-to-treat analysis. ‡ Data were available from only 81 and 151 patients in the AM and DP groups, respectively. § Significant difference from day 28 (P < 0.01). ¶ Significant difference between AM and DP groups (P < 0.02).
adults: weakness (154 [75%] of 205 versus 85 [89%] of 95; \(P = 0.004\)), muscle pain (32 [15%] of 205 versus 80 [84%] of 95; \(P < 0.001\)), headache (144 [70%] of 205 versus 91 [96%] of 95; \(P < 0.001\)), nausea (76 [37%] of 205 versus 48 [50.0%] of 95; \(P = 0.02\)), pruritis (0% versus 3% \(P < 0.03\)), and tinnitus (3 [1%] of 205 versus 15 [16%] of 95; \(P < 0.001\)).

The median and range fever and parasite clearance times were not significantly different between children (23 [7–64] hours and 2 [1–3] days) and adults (21 [7–68] hours and 2 [1–3] days) \(P = 0.48\) and 0.06, respectively, by Mann-Whitney U test). Mean hematocrit values on admission and on all days \([1–2\) days] \(P = 0.48\) were not significantly different between children (23 [7–64] hours and 2 [1–3] days) and adults (21 [7–68] hours and 2 [1–3] days) \(P = 0.001\) in both treatment groups.

*Plasmodium falciparum* gametocyte carriage. Twenty-three (7.7%) patients had patent gametocytes on admission, 6 (6%) in the AM group and 17 (8%) in the DP group \(P = 0.48\). The time to clearance of gametocytes was longer in the DP group than in the AM group. All patients in the AM group and only 8 (47%) patients in the DP group cleared their gametocytes by day 7 \(P = 0.048\), by Fisher’s exact test). After treatment, gametocytogenesis developed on day 56 in one patient in the AM group who did not have gametocytes on admission at the same time as a new *P. falciparum* infection. Gametocytogenesis also developed after admission (days 1–2), which cleared by day 14, in 6 other patients in the DP group who did not have gametocytes on admission. The proportion of patients in whom gametocytes developed in the AM and DP groups was not significantly different \(P = 0.68\). Person-gametocyte-weeks was estimated as 12 weeks/1,000 patients in the AM group and 22 weeks/1,000 patients in the DP group \(P = 0.069\). The proportion of patients with gametocytes at any time point was higher in children (13%, 27 of 205) than in adults (3%, 3 of 95) \(P = 0.006\), by Fisher’s exact test), and the proportion of patients with gametocytes after treatment was 4% (7 of 185) in children and 0% (0 of 92) in adults \(P = 0.10\), by Fisher’s exact test).

**Adverse events.** Adverse events were reported by 269 (89.7%) patients ≥ 3 years of age who were able to answer questions about these symptoms. A convulsion and coma developed in a four-year-old patient after treatment with AM (see above). Severe malaria or other life-threatening events after treatment did not develop in any other patients. The frequency of patients with symptoms and signs before treatment that may subsequently be confused with drug-related adverse events did not differ significantly between groups \(P > 0.05\). The proportion of patients with at least one recorded potential side effect/ adverse event after treatment was significantly higher in the AM group (38 of 87, 44%) than in the DP group (57 of 182, 31%) \(P = 0.04\). The incidence of post-treatment dizziness, nausea, insomnia, and anorexia were all significantly higher in AM group than in the DP group \(P \leq 0.01\) (Table 3). The frequency of some adverse events was significantly higher in adults than in children (see above).

**Electrocardiograms.** No significant ECG abnormalities were found before or after treatment. The mean ECG-determined heart rate was significantly lower after treatment at days 2 and 7 for both treatment groups \((P < 0.001)\) than at admission (Table 4). The mean PR interval was significantly longer after treatment (on days 2 and 7) than on admission for both groups \((P \leq 0.001)\); mean PR interval (95% CI) difference between baseline and days 2 and 7 were –9.5 (–13.1 to –6.0) and –5.7 (–9.1 to –2.4), respectively, for the AM group and –5.3 (–7.1 to –3.5) and –3.6 (–5.6 to –1.6), respectively, for the DP group. Four patients (two in the AM group and two in the DP group) had PR interval prolongation (> 200 ms) after treatment on day 2 or day 7 but the PR intervals on day 7 or day 28 were normal. The mean QRS interval was also significantly longer on day 2 than on admission \((P = 0.001)\) but was similar to the baseline value by day 7 \((P > 0.05)\) in both treatment groups (Table 4).

Modeling the QT/RR relationship\(^{20–22}\) gave a correction formula of \(Q T C = R R^{-0.5} × P_{. F A L C I P A R U M} M A L A R I A}^\) on day 2, and \(Q T C = R R^{-0.493}\) on day 7. Because this calculation approximates to Bazzet’s formula \((Q T C = Q T/RR^{0.5})\), Bazzet’s correction was used. There was weak negative correlation or no correlation between QTc and RR intervals at baseline or after treatment (Table 4). The mean (95% CI) QTc (ms) was
significantly prolonged on day 2 than at baseline in the DP group (P < 0.001) (Table 4). The mean (95% CI) QTc (ms) prolongation on days 2 and 7 compared with those at baseline were 3.5 (−1.1 to 8.2) and 5.2 (−0.9 to 11.4), respectively, for the AM group and 9.5 (5.9–13.1) and 1.4 (−2.5 to 5.4), respectively, for the DP group. The proportion of the patients with prolonged QTc pre-treatment and on days 2 and 7 did not differ significantly between the AM and DP groups (P > 0.05) (Table 4). The numbers of patients with QTc lengthening (defined as > 30 ms and > 60 ms by the U.S. Food and Drug Administration33,34) were not significantly different between the two treatment groups (P > 0.05) (Table 4).

**DISCUSSION**

This study confirms that the fixed-dose, co-formulated ACT dihydroartemisinin-piperaquine is not inferior to the more expensive, not co-formulated AM in the treatment of uncomplicated *P. falciparum* malaria in southern Laos and showed a 63-day follow-up cure rate of almost 100% for both drugs. Our previous study conducted in the same area, using GMP drugs from China, also demonstrated that the 42-day follow up cure rates were 95% in 22 of the 26 studies. In a meta-analysis of individual patient data from 7 trials in Africa and Southeast Asia that included 1,814 patients, it was found that the proportion of patients in whom gametocytemia developed and the gametocyte carriage rate was higher after DP than after the comparator. However, in Laos, the proportion of patients with gametocytemia at any time point on or after day 7 and the mean (95% CI) person-gametocyte-weeks did not significantly differ between the AM and DP groups, probably because of the smaller sample size.

An important limitation of this trial for public health policy in Laos is that DP was compared with AM and not with AL, which is the current Lao national treatment policy. Artesunate-mefloquine was the comparator because this was required for the multicentre trial. In 2005, AM was the intended national treatment policy but sudden and unexpected unavailability of AM led the Government of Laos to change to AL. Artesunate-mefloquine is national policy in Cambodia, Myanmar (Burma), and Thailand, and DP is national policy in Vietnam and China. There have been six comparisons of DP versus AL, all in children from Africa, and the 28–63-day cure rates were not significantly different between AL (84–96%) and DP (93–100%).

In the multi-center study, overall *P. falciparum* gametocyte prevalence (days 0–70) measured as person-gametocyte-weeks was significantly higher after DP than AM (9.7% versus 4.8%; P = 0.006; modified ITT). In addition, Zwang and others53 found that the proportion of patients in whom gametocytemia developed and the gametocyte carriage rate was higher after DP than after the comparator. However, in Laos, the proportion of patients with gametocytemia at any time point on or after day 7 and the mean (95% CI) person-gametocyte-weeks did not significantly differ between the AM and DP groups, probably because of the smaller sample size.

Of 26 published clinical trials of either DP or DP plus trimethoprim (Table 5) that included 6,010 patients, 8 deaths were reported (6 children and 2 adults). One adult death was probably caused by severe malaria that developed during treatment with DP; the other seven deaths were considered unlikely to be a consequence of DP. No deaths were reported in the larger multi-center trial of DP in Asia of 767 patients, of which this trial was a part.52

Although piperaquine has a chemical structure similar to chloroquine, which can cause lethal cardiologic effects, results of our study are consistent with that of previous studies, which suggested that at therapeutic doses, DP does not have clinically significant effects on the ECG results.

In Laos, AL is provided by the Global Fund at a cost of < 1.4 U.S.$/adult treatment course and is free to patients through...
The public sector. When compared with AL, DP has potential benefits because it is less expensive to manufacture, may have better adherence, and can be used in a once a day regimen rather than a twice a day regimen. Although a study of the unsupervised effectiveness and supervised efficacy of DP and AM treatment in western Myanmar suggested good adherence to both regimens, there have been no studies directly comparing DP and AL adherence. Although DP has been successfully used as rescue therapy for multidrug-resistant \textit{P. falciparum} malaria in pregnancy (PCR-adjusted cure rate at 63 days was 92.3%), further studies are needed to establish its safety in pregnancy.

In conclusion, DP demonstrated non-inferiority to the reference treatment of AM in Laos and in the overall three-country study. The high efficacy and relative safety of DP and its availability as a fixed combination and probable lower cost compared with other candidate ACTs suggests that it could play a significant role in the first-line treatment of uncomplicated \textit{P. falciparum} malaria in Laos.

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The table below shows the review of efficacy or effectiveness of DP in 26 clinical trials (1,004 adults; 2,976 children; 2,030 not identified as children or adults; total = 6,010) with reference to the comparator:

| Reference | Country | Comparator | Day of follow-up | No. patients | Cure rate (%) | No. deaths |
|-----------|---------|------------|------------------|--------------|---------------|------------|
| Asia      |         |            |                  |              |               |            |
| 37        | China   | 0          | 28               | 60           | 96.7          | Unknown    |
| 38        | China   | AL         | 28               | 51           | 100           | Unknown    |
| 39        | Cambodia| 0          | 56               | 30           | 76            | 95.5       |
| 33        | Cambodia| 0          | 28               | 32           | 30            | 100        |
| 40        | Cambodia| A + M      | 64               | 215          | 97.5          | None reported |
| 41        | Vietnam | A + M      | 56               | 76           | 97.4          | None reported |
| 42        | Vietnam | AQ + P     | 28               | 84           | 0             | None reported |
| 43        | Vietnam | A + AQ     | 42               | 34           | 15            | None reported |
| 44        | Thailand| A + M      | 28               | 234          | 0             | None reported |
| 34        | Thailand| A + M      | 63               | 353          | 97            | 1 man at day 28 from gunshot |
| 45        | Thailand| A + M      | 28               | 120          | 0             | None reported |
| 46        | Thailand| A + M      | 63               | 333          | 99.7          | 1 woman at < 24 hours after admission in the clinic (probably from severe malaria) unlikely caused by DP. 1 girl (13 years of age) at day 3 (unlikely caused by DP). |
| Africa    |         |            |                  |              |               |            |
| 47        | Thailand| AN + PPQ, A + M, and AL | 28 | 82 | 98.8 | None reported |
| 35        | Burma   | A + M      | 42               | 327          | 99            | One 11-year-old died aparasitemic at day 21 |
| 11        | Laos    | A + M      | 42               | 36           | 74            | 100        |
| 48        | Indonesia| A + AQ | 42               | 92           | 76            | 96         |
| 49        | Papua New Guinea | CQ + SP, A + SP, and AL | 42 | 482 | 88 | None reported |
| 50        | Vietnam | AN + PPQ   | 28               | 51           | 100           | None reported |
| Africa    |         |            |                  |              |               |            |
| 28        | Uganda  | AL         | 63               | 0            | 351†          | None reported |
| 29        | Burkina Faso, Kenya | AL | 42 | 0 | 1,039 | 86 | 1 child (3 year-old girl) died 24 hours after treatment with DP. The most likely cause was sepsis or severe malaria. |
| 51        | Rwanda  | A/SP + AQ  | 28               | 0            | 252           | 95.2       |
| 25        | Uganda  | AL         | 42               | 0            | 211           | 93         |
| 30        | Kenya   |             | 28               | 0            | 73            | 100        |
| 27        | Uganda  | AL         | 42               | 0            | 215           | 98         |
| 26        | Burkina Faso | AQ + SP and AL | 42 | 0 | 187 | 98 | None reported |
| South America |         |            |                  |              |               |            |
| 52        | Peru    | A + M      | 63               | 168          | 94            | 98.4       |

* DP = dihydroartemisinin plus piperaquine for 3 days; AL = artemether-lumefantrine; A = artesunate; AN = artemisinin; M = mefloquine; AQ = atovaquone; P = proquanil; PPQ = piperaquine; CQ = chloroquine; SP = sulfadoxine-pyrimethamine.
† No. treatments rather than no. patients (longitudinal study).
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