## Standard report for Vivax Malaria

### WWARN Vivax Primaquine Study Group

For further information go to https://www.iddo.org/wwarn/vivax-reports

08 May, 2025

#### Introduction

This report has been produced for countries: Afghanistan

The studies included within this report are shown in Table 0.

Table 0: Studies included in this report

Author-year	Country	Recruitment Period	Age range (years)	Follow up (days)	Included treatment arms*	PQ supervison	Patients avail- able
Taylor-2019	Afghanistan	2014 - 2016	0.8 - 70	365	Cq,	Fully supervised	426
					Cq_Pq_7.0_14d_D0,		
					Cq_Pq_7.0_7d_D0		
Awab-2017	Afghanistan	NA	2 - 84	390	Cq,	<50% supervised	544
					Cq_Pq_3.5_14d_D0		

<sup>\*</sup> ACT – artemisinin-based combination treatment; As – artesunate; AL – artemether-lumefantrine; Aq – amodiaquine; Cq – chloroquine; DP – dihydroartemisinin-piperaquine; GI – gastrointestinal; Mf – mefloquine; PQ/Pq – primaquine; SP – sulfadoxine-pyrimethamine;

Treatment code describes (schizontocidal drug)(hypnozoitocidal drug)(total primaquine dose)(duration of primaquine treatment eg 14d = 14 days)(primaquine start day)

#### 1: EFFICACY

#### 1.1: Description

The efficacy study was undertaken to better understand the impact of primaquine dose on the prevention of P. vivax recurrences. Inclusion in the efficacy meta-analysis was restricted to studies with 42 days or more follow up and patients with data on day 0 parasitaemia.

In this report the efficacy study includes 970 patients across 3 study sites, from 2 studies.

#### 1.2: Characteristics of Study Population

Table 1\_eff: Characteristics of the study population for the efficacy study analysis, categorised by total primaquine category

	Primaquine (PQ) Treatment				
	No primaquine (N=351)	Very low dose total primaquine (<2 mg/kg)(N=7)	Low dose total primaquine (2 - <5 mg/kg)(N=297)	High dose total primaquine (>= 5 mg/kg)(N=315)	Total ( <b>N=970</b> )
Age (years)					
Mean (SD)	18 (14)	21 (11)	20 (14)	16 (12)	18 (13)
Age Category					
<5	25 (7%)	0 (0%)	18 (6%)	27 (9%)	70 (7%)
5-<15	156 (44%)	1 (14%)	113 (38%)	156 (50%)	$426 \ (44\%)$
>=15	170 (48%)	6 (86%)	166 (56%)	132 (42%)	474 (49%)
Gender	, ,	, ,	, ,	, ,	, ,
Male	243 (69%)	6 (86%)	196 (66%)	217 (69%)	662 (68%)
Female	108 (31%)	1 (14%)	101 (34%)	98 (31%)	308 (32%)
Weight (kg)					
Mean (SD)	44 (21)	59 (21)	49 (22)	39 (20)	44 (21)
Missing	5 (1.4%)	0 (0%)	4 (1.3%)	0 (0%)	9 (0.9%)
Malnutrition	0 (1.470)	0 (070)	4 (1.570)	0 (070)	3 (0.370)
No	17 (5%)	1 (14%)	15 (5%)	30 (10%)	63 (6%)
Yes	11 (3%)	0 (0%)	12 (4%)	6 (2%)	29 (3%)
	` '	` /	` ,	` '	` /
Missing	$323 \ (92.0\%)$	6 (85.7%)	270 (90.9%)	279 (88.6%)	878 (90.5%)
Fever day 0	OF (FOI)	. (==01)	0 (004)	F0 (0.404)	111 (1007)
No	25 (7%)	4 (57%)	9 (3%)	76 (24%)	114 (12%)
Yes P. vivax baseline parasitaemia	326 (93%)	3 (43%)	288 (97%)	239 (76%)	856 (88%)
Median (IQR)	2200 [1200, 3620]	1481 [833, 2685]	2407 [1400, 4000]	1185 [648, 3463]	2000 [931, 3662
Haemoglobin day 0 (g/dL)					
Mean (SD)	12 (1.7)	14(2.0)	12 (1.6)	13 (1.7)	12(1.7)
Missing PQ daily dose (mg/kg)	3 (0.9%)	0 (0%)	1 (0.3%)	1 (0.3%)	5 (0.5%)
Mean (SD)  Duration of PQ treatment		1.2 (0.41)	3.4 (0.50)	8.0 (2.0)	5.7(2.7)
7 days		4 (57%)	14 (5%)	154 (49%)	172 (28%)
14 days		3 (43%)	283 (95%)	161 (51%)	447 (72%)
Method to calculate PQ dose		, ,	` '	` '	, ,
Per actual dose		7 (100%)	23 (8%)	315 (100%)	345 (56%)
Per dosing protocol Start day of PQ treatment		0 (0%)	274 (92%)	0 (0%)	274 (44%)
Day 0		7 (100%)	294 (99%)	315 (100%)	616 (100%)
Day 1		0 (0%)	2 (1%)	0 (0%)	2 (0%)
=		` ′	` ′	` ′	` '
Day 2		0 (0%)	1 (0%)	0 (0%)	1 (0%)

	No primaquine (N=351)	Very low dose total primaquine (<2 mg/kg)(N=7)	Low dose total primaquine (2 - <5 mg/kg)(N=297)	High dose total primaquine (>= 5 mg/kg)(N=315)	Total (N=970)
Day 4 Day 5 Day 6		0 (0%) 0 (0%) 0 (0%)	0 (0%) 0 (0%) 0 (0%)	0 (0%) 0 (0%) 0 (0%)	0 (0%) 0 (0%) 0 (0%)
Level of PQ supervision Unsupervised Partially supervised Fully supervised Was PQ taken with food?		0 (0%) 0 (0%) 7 (100%)	0 (0%) 274 (92%) 23 (8%)	0 (0%) 0 (0%) 315 (100%)	0 (0%) 274 (44%) 345 (56%)
No Yes Recommended Other treatment given		0 (0%) 7 (100%) 0 (0%)	0 (0%) 23 (8%) 274 (92%)	0 (0%) 315 (100%) 0 (0%)	0 (0%) 345 (56%) 274 (44%)
AL	2 (1%)	0 (0%)	0 (0%)	0 (0%)	2 (0%)
AsAq AsMf Cq DP Transmission intensity of the site	0 (0%) 0 (0%) 349 (99%) 0 (0%)	0 (0%) 0 (0%) 7 (100%) 0 (0%)	0 (0%) 0 (0%) 297 (100%) 0 (0%)	0 (0%) 0 (0%) 315 (100%) 0 (0%)	0 (0%) 0 (0%) 968 (100%) 0 (0%)
Low Moderate High Not available <b>Geographical region</b>	0 (0%) 0 (0%) 351 (100%) 0 (0%)	0 (0%) 0 (0%) 7 (100%) 0 (0%)	0 (0%) 0 (0%) 297 (100%) 0 (0%)	0 (0%) 0 (0%) 315 (100%) 0 (0%)	0 (0%) 0 (0%) 970 (100%) 0 (0%)
Africa Americas Asia-Pacific Relapse Peridocity Low periodicity	0 (0%) 0 (0%) 351 (100%) 351 (100%)	0 (0%) 0 (0%) 7 (100%) 7 (100%)	0 (0%) 0 (0%) 297 (100%) 297 (100%)	0 (0%) 0 (0%) 315 (100%) 315 (100%)	0 (0%) 0 (0%) 970 (100%) 970 (100%)
High periodicity G6PD categories	0 (0%)	0 (0%)	0 (0%)	0 (0%)	0 (0%)
(Qualitative test) <30% >=30% G6PD categories (Quantitative test)	0 (0%) 351 (100%)	0 (0%) 7 (100%)	0 (0%) 297 (100%)	0 (0%) 315 (100%)	0 (0%) 970 (100%)
<30% 30-<70% >=70% Missing	0 (0%) 0 (0%) 0 (0%) 351 (100%)	0 (0%) 0 (0%) 0 (0%) 7 (100%)	0 (0%) 0 (0%) 0 (0%) 297 (100%)	0 (0%) 0 (0%) 0 (0%) 315 (100%)	0 (0%) 0 (0%) 0 (0%) 970 (100%)

#### 1.3: Risk of recurrence

Kaplan-Meier survival analysis was used to calculate risk of recurrence between day 7 and 365. Patients were left censored at day 7 and right censored at the first of: the day last reviewed, the last day prior to a 60-day blood smear gap or the last day of study follow up. Outcomes were stratified by primaquine treatment arm: no primaquine, low total dose primaquine (2 to <5 mg/kg) and high total dose primaquine ( $\ge 5$  mg/kg). Very low total dose primaquine (<2 mg/kg) was not presented due to low numbers of patients treated with this dose.

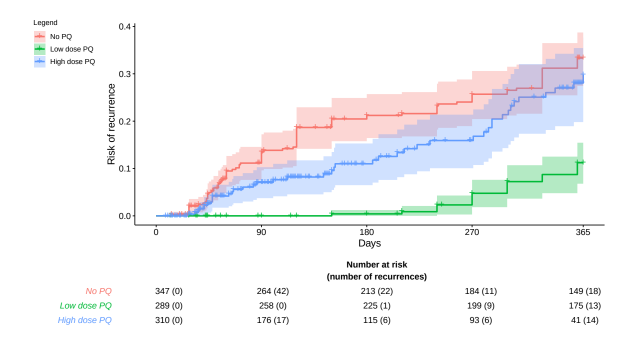


Figure 0\_eff: Kaplan-Meier figure of cumulative risk of recurrence between day 7 and day 365 for primaquine treatment category. Please interpret the results of this figure with caution as there may not always be paired treatment comparisons in the original studies contributing to these pooled results.

Cox regression analysis for the time to first vivax recurrence between day 7 and 180 was performed to determine the effect of primaquine dose. Analysis was restricted to patients treated with daily primaquine or no primaquine. Potential confounders including sex, age and baseline parasitaemia were adjusted for with shared frailty for study site.

Similar but separate multivariable Cox regression analyses were undertaken to investigate primaquine duration, also adjusting for total actual mg/kg dose, in i) patients treated with low total dose primaquine and ii) patients treated with high total dose primaquine.

Care should be taken when interpreting these results, as model assumptions have not been fully assessed in this automated report format.

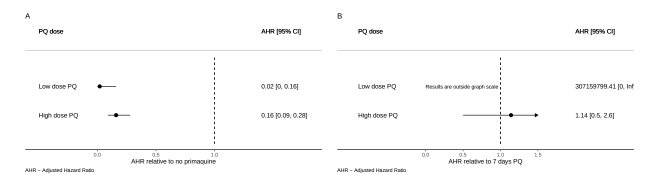


Figure 1\_eff: Hazard ratio between day 7 and day 180 for A: total dose of primaquine and B: 14-day vs 7-day primaquine duration, stratified by total dose of primaquine

#### 2: HAEMATOLOGY

#### 2.1: Description

Haematological safety is a key concern for clinicians and policymakers in the implementation of primaquine radical cure, due to the risk of haemolysis in patients with G6PD deficiency. This individual patient data meta-analysis was conducted to assess the evidence for adverse haematological outcomes related to primaquine dose, with consideration of patients G6PD status.

Inclusion in the haematological safety meta-analysis was restricted to studies with 28 days or more follow up, patients with data on day 0 parasitaemia, patients with available data on day 0 haemoglobin levels or haematocrit, patients with an available haemoglobin measurement on at least one more day during the follow-up period and patients with data on daily primaquine dose.

The haematology study included 965 patients across 3 study sites, from 2 studies.

#### 2.2 Characteristics of Study Population

Table 1\_saf: Characteristics of the study population for the safety study analysis, categorised by total primaquine category

		Primaquin	e Treatment			
	No primaquine (N=348)	Low dose daily primaquine (<0.375 mg/kg/day) (N=255)	Intermediate dose daily primaquine (>= $0.375 \& < 0.75 $ mg/kg/day) (N=182)	High dose daily primaquine (>= $0.75$ mg/kg/day) (N= $180$ )	Total ( <b>N=965</b> )	
Age (years) Mean (SD) Age Category	18 (14)	21 (14)	18 (14)	14 (9.9)	18 (13)	
<5 5-<15	25 (7.18%) 154 (44.25%)	14 (5.49%) 86 (33.73%)	12 (6.59%) 86 (47.25%)	19 (10.56%) 97 (53.89%)	70 (7.25%) 423 (43.83%)	
>=15 Gender	169 (48.56%)	155 (60.78%)	84 (46.15%)	64 (35.56%)	472 (48.91%)	
Male Female Weight (kg)	240 (68.97%) 108 (31.03%)	170 (66.67%) 85 (33.33%)	129 (70.88%) 53 (29.12%)	119 (66.11%) 61 (33.89%)	658 (68.19%) 307 (31.81%)	
Mean (SD) Missing Malnutrition	44 (21) 5 (1.4%)	51 (21) 4 (1.6%)	42 (22) 0 (0%)	36 (19) 0 (0%)	44 (21) 9 (0.9%)	
No Yes	17 (4.89%) 11 (3.16%)	14 (5.49%) 8 (3.14%)	13 (7.14%) 5 (2.75%)	19 (10.56%) 5 (2.78%)	63 (6.53%) 29 (3.01%)	
Missing Fever day 0	320 (92.0%)	233 (91.4%)	164 (90.1%)	156 (86.7%)	873 (90.5%)	
No Yes P. vivax baseline parasitaemia	25 (7.18%) 323 (92.82%)	7 (2.75%) 248 (97.25%)	47 (25.82%) 135 (74.18%)	35 (19.44%) 145 (80.56%)	114 (11.81%) 851 (88.19%)	
Median (IQR)	2200 [1200, 3642]	2600 [1450, 4000]	1296 [653, 3028]	1287 [667, 3704]	2000 [926, 3667	
Haemoglobin day 0 (g/dL) Mean (SD) PQ daily dose (mg/kg)	12 (1.7)	12 (1.6)	13 (1.7)	13 (1.8)	12 (1.7)	
Mean (SD) Missing Mean (SD) Missing Method to calculate PQ dose		3.4 (0.48) 0 (0%) 14 (0) 0 (0%)	6.6 (1.9) 0 (0%) 13 (1.9) 0 (0%)	8.1 (2.8) 0 (0%) 7.9 (2.3) 0 (0%)	5.7 (2.7) 0(0%) 12 (3.1)	

	No primaquine (N=348)	Low dose daily primaquine ( $<0.375$ mg/kg/day) ( $N=255$ )	Intermediate dose daily primaquine (>= $0.375 \& < 0.75 $ mg/kg/day) (N=182)	High dose daily primaquine (>= 0.75 mg/kg/day) (N=180)	Total (N=965)
Per dosing protocol		248 (97.25%)	22 (12.09%)	3 (1.67%)	273 (44.25%)
Missing Start day of PQ treatment Day 1 Day 2		0 (0%) 2 (0.78%) 1 (0.39%)	0 (0%) 0 (0.00%) 0 (0.00%)	0 (0%) 0 (0.00%) 0 (0.00%)	0(0%) 2 (0.32%) 1 (0.16%)
Day 3 Day 4 Day 5 Day 6 Missing Level of PQ supervision		0 (0.00%) 0 (0.00%) 0 (0.00%) 0 (0.00%) 0 (0%)	0 (0.00%) 0 (0.00%) 0 (0.00%) 0 (0.00%) 0 (0%)	0 (0.00%) 0 (0.00%) 0 (0.00%) 0 (0.00%) 0 (0%)	0 (0.00%) 0 (0.00%) 0 (0.00%) 0 (0.00%) 0(0%)
Partially supervised Fully supervised Was PQ taken with food? No		248 (97.25%) 7 (2.75%) 0 (0.00%)	22 (12.09%) 160 (87.91%) 0 (0.00%)	3 (1.67%) 177 (98.33%) 0 (0.00%)	273 (44.25%) 344 (55.75%) 0 (0.00%)
Recommended		248 (97.25%)	$22\ (12.09\%)$	3 (1.67%)	273 (44.25%)
Other treatment given $AL$ $AsAq$ $Cq$ $DP$	2 (0.57%) 0 (0.00%) 346 (99.43%) 0 (0.00%)	0 (0.00%) 0 (0.00%) 255 (100.00%) 0 (0.00%)	0 (0.00%) 0 (0.00%) 182 (100.00%) 0 (0.00%)	0 (0.00%) 0 (0.00%) 180 (100.00%) 0 (0.00%)	2 (0.21%) 0 (0.00%) 963 (99.79%) 0 (0.00%)
Transmission intensity of the site Low Moderate High Not available	0 (0.00%) 0 (0.00%) 348 (100.00%) 0 (0.00%)	0 (0.00%) 0 (0.00%) 255 (100.00%) 0 (0.00%)	0 (0.00%) 0 (0.00%) 182 (100.00%) 0 (0.00%)	0 (0.00%) 0 (0.00%) 180 (100.00%) 0 (0.00%)	0 (0.00%) 0 (0.00%) 965 (100.00%) 0 (0.00%)
Geographical region Africa Americas Asia-Pacific Relapse Peridocity	0 (0.00%) 0 (0.00%) 348 (100.00%)	0 (0.00%) 0 (0.00%) 255 (100.00%)	0 (0.00%) 0 (0.00%) 182 (100.00%)	0 (0.00%) 0 (0.00%) 180 (100.00%)	0 (0.00%) 0 (0.00%) 965 (100.00%)
Low periodicity High periodicity G6PD categories (Qualitative test)	348 (100.00%) 0 (0.00%)	255 (100.00%) 0 (0.00%)	182 (100.00%) 0 (0.00%)	180 (100.00%) 0 (0.00%)	965 (100.00%) 0 (0.00%)
<30% >=30%	0 (0.00%) 348 (100.00%)	0 (0.00%) 255 (100.00%)	0 (0.00%) 182 (100.00%)	0 (0.00%) 179 (99.44%)	$\begin{array}{c} 0 & (0.00\%) \\ 964 & (99.90\%) \end{array}$
Unknown G6PD categories (Quantitative test) <30% 30-<70% >=70%	0 (0.00%) 0 (0.00%) 0 (0.00%) 0 (0.00%)	0 (0.00%) 0 (0.00%) 0 (0.00%) 0 (0.00%)	0 (0.00%) 0 (0.00%) 0 (0.00%) 0 (0.00%)	1 (0.56%) 0 (0.00%) 0 (0.00%) 0 (0.00%)	1 (0.10%) 0 (0.00%) 0 (0.00%) 0 (0.00%)
Unknown	348 (100.00%)	255 (100.00%)	182 (100.00%)	180 (100.00%)	965 (100.00%)

#### 2.3 Summary of the haematology outcomes

Table 2 below provides a summary of the outcome experienced within each primaquine treatment arm for participants with G6PD activity  $\geq 30\%$ .

Table 2\_saf: Summary of safety outcomes, categorised by total primaquine category

		Primaquin	e Treatment			
	No primaquine	Low dose daily primaquine $(<0.375 \ \mathrm{mg/kg/day})$	Intermediate dose daily primaquine ( $0.375~\&<0.75$ mg/kg/day)	High dose daily primaquine ( 0.75 mg/kg/day)	Total	
Drop in haemoglobin of >25% AND Hb below 7 g/dL						
No Yes Missing Drop in haemoglobin of >5 g/dL from baseline between days 1-14	339 (97.4 %) 0 (0.0 %) 9 (2.6%)	252 (98.8 %) 0 (0.0 %) 3 (1.2%)	175 (96.2 %) 0 (0.0 %) 7 (3.8%)	168 (93.9 %) 0 (0.0 %) 11 (6.1%)	934 (96.9 %) 0 (0.0 %) 30 (3.1%)	
No Yes Missing Drop in haemoglobin to <5 g/dL between days 1 and 14	338 (97.1 %) 1 (0.3 %) 9 (2.6%)	252 (98.8 %) 0 (0.0 %) 3 (1.2%)	175 (96.2 %) 0 (0.0 %) 7 (3.8%)	168 (93.9 %) 0 (0.0 %) 11 (6.1%)	933 (96.8 %) 1 (0.1 %) 30 (3.1%)	
No	339~(97.4~%)	252 (98.8 %)	175~(96.2~%)	168 (93.9 %)	934 (96.9 %)	
Yes Missing	0 (0.0 %) 9 (2.6%)	0 (0.0 %) 3 (1.2%)	0 (0.0 %) 7 (3.8%)	0 (0.0 %) 11 (6.1%)	0 (0.0 %) 30 (3.1%)	
Anaemia developed at days 2 or 3  Nil (Hb: >=11 g/dL)  Mild (Hb: >=8 g/dL & <11 g/dL)	228 (65.5 %) 6 (1.7 %)	185 (72.5 %) 0 (0.0 %)	119 (65.4 %) 13 (7.1 %)	101 (56.4 %) 10 (5.6 %)	633 (65.7 %) 29 (3.0 %)	
Moderate (Hb: $>=5$ g/dL & $<8$ g/dL) Severe (Hb $<5$ g/dL) Missing Anaemia developed at days 5-7	0 (0.0 %) 0 (0.0 %) 114 (32.8%)	0 (0.0 %) 0 (0.0 %) 70 (27.5%)	0 (0.0 %) 0 (0.0 %) 50 (27.5%)	0 (0.0 %) 0 (0.0 %) 68 (38.0%)	0 (0.0 %) 0 (0.0 %) 302 (31.3%)	
Nil (Hb: >=11 g/dL)	238~(68.4~%)	187 (73.3 %)	118 (64.8 %)	102~(57.0~%)	645 (66.9 %)	
$\label{eq:midd} \begin{array}{ll} \mbox{Mild (Hb: }>=8\ \mbox{g/dL \& }<11\ \mbox{g/dL)} \\ \mbox{Moderate (Hb: }>=5\ \mbox{g/dL \& }<8\ \mbox{g/dL)} \\ \mbox{Severe (Hb }<5\ \mbox{g/dL)} \\ \mbox{Missing} \\ \mbox{Change in haemoglobin on days 2-3} \\ \mbox{from day 0} \end{array}$	5 (1.4 %) 0 (0.0 %) 0 (0.0 %) 105 (30.2%)	1 (0.4 %) 0 (0.0 %) 0 (0.0 %) 67 (26.3%)	4 (2.2 %) 0 (0.0 %) 0 (0.0 %) 60 (33.0%)	9 (5.0 %) 0 (0.0 %) 0 (0.0 %) 68 (38.0%)	19 (2.0 %) 0 (0.0 %) 0 (0.0 %) 300 (31.1%)	
Mean (SD)	-0.151 (0.628)	-0.0588 (0.546)	-0.225 (0.942)	-0.000741 (0.890)	-0.112 (0.720)	
Missing Change in haemoglobin on days 5-7 from day 0	38 (10.9%)	12 (4.7%)	40 (22.0%)	44 (24.6%)	134 (13.9%)	
Mean (SD) Missing	-0.0134 (0.560) 28 (8.0%)	0.0159 (0.679) 9 (3.5%)	0.134 (1.06) 50 (27.5%)	0.219 (1.36) 47 (26.3%)	0.0557 (0.856) 134 (13.9%)	
Relative percentage (%) change in haemoglobin on days 2-3 from day 0 Mean (SD) Missing Relative percentage (%) change in haemoglobin on days 5-7 from day 0	1.07 (4.42) 38 (10.9%)	0.370 (4.84) 12 (4.7%)	1.71 (7.22) 40 (22.0%)	-0.224 (7.24) 44 (24.6%)	0.763 (5.64) 134 (13.9%)	
Mean (SD)	-0.0751 (4.36)	-0.303 (6.02)	-1.19 (8.22)	-2.19 (10.8)	-0.656 (6.90)	
Missing	28 (8.0%)	9 (3.5%)	50 (27.5%)	47 (26.3%)	134 (13.9%)	

## 2.4: Change in Haemoglobin (Hb) levels between primaquine treatment groups

The following figure provides the estimated change in haemoglobin from day 0 for different primaquine doses at at day 2/3 and days 5/7, adjusted for baseline haemoglobin, age, sex and day 0 parasitaemia and allowing for clustering by study site, in participants with  $\geq 30\%$  G6PD activity.

Care should be taken when interpreting these results, as model assumptions have not been fully assessed in this automated report format.



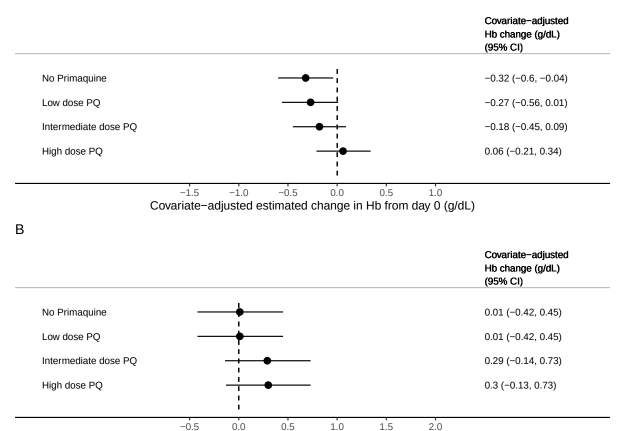


Figure 1\_saf: The covariate-adjusted estimated change in Hb between primaquine daily dose groups on (A) days 2-3 and (B) days 5-7, in patients with  $\geq 30\%$  G6PD activity.

Covariate-adjusted estimated change in Hb from day 0 (g/dL)

1.5

#### 3: TOLERABILITY

#### 3.1: Description

This individual patient data meta-analysis was conducted in order to understand the effect of primaquine dose on the gastrointestinal side effects.

Inclusion in the gastrointestinal tolerability meta-analysis was restricted to studies with 28 days or more followup, data from pre-specified symptom questionnaires (symptom checklist), patients with data on vivax parasite count at baseline, patients starting primaquine by day 2, patients not receiving intermittent primaquine (defined as primaquine administered weekly or monthly, rather than daily) and patients with data on daily primaquine dose.

The tolerability study included 970 patients across 3 study sites, from 2 studies.

#### Characteristics of Study Population

Table 1\_tol: Characteristics of the study population for the tolerabilty study analysis, categorised by total primaquine category

		Primaquine	Treatment		
	No primaquine (N=351)	Low dose daily primaquine $(<0.375$ mg/kg/day) $(N=256)$	Intermediate dose daily primaquine (>=0.375 & $<0.75$ mg/kg/day) (N=182)	High dose daily primaquine (>= 0.75 mg/kg/day) (N=181)	Total ( <b>N=970</b> )
Age (years) Mean (SD) Age Category	18 (14)	21 (14)	18 (14)	14 (9.9)	18 (13)
<5 5-<15	25 (7%) 156 (44%)	14 (5%) 86 (34%)	12 (7%) 86 (47%)	19 (10%) 98 (54%)	70 (7%) 426 (44%)
>=15 <b>Gender</b>	170 (48%)	156 (61%)	84 (46%)	64 (35%)	474 (49%)
Male Female Weight (kg)	243 (69%) 108 (31%)	171 (67%) 85 (33%)	129 (71%) 53 (29%)	119 (66%) 62 (34%)	662 (68%) 308 (32%)
Mean (SD) Missing Malnutrition	44 (21) 5 (1.4%)	51 (21) 4 (1.6%)	42 (22) 0 (0%)	36 (19) 0 (0%)	44 (21) 9 (0.9%)
No Yes	17 (5%) 11 (3%)	14 (5%) 8 (3%)	13 (7%) 5 (3%)	19 (10%) 5 (3%)	63 (6%) 29 (3%)
Missing Fever day 0	323 (92.0%)	234 (91.4%)	164 (90.1%)	157 (86.7%)	878 (90.5%)
No Yes P. vivax baseline parasitaemia	25 (7%) 326 (93%)	7 (3%) 249 (97%)	47 (26%) 135 (74%)	35 (19%) 146 (81%)	114 (12%) 856 (88%)
Median (IQR)	2200 [1200, 3620])	2600 [1475, 4000])	1296 [653, 3028])	1278 [667, 3704])	2000 [931, 3662])
Haemoglobin day 0 (g/dL) Mean (SD) Missing PQ daily dose (mg/kg)	12 (1.7) 3 (0.9%)	12 (1.6) 1 (0.4%)	13 (1.7) 0 (0%)	13 (1.8) 1 (0.6%)	12 (1.7) 5 (0.5%)
Mean (SD) <b>Duration of PQ treatment</b> 7 days 14 days		3.4 (0.48) 0 (0%) 256 (100%)	6.6 (1.9) 14 (8%) 168 (92%)	8.1 (2.8) 158 (87%) 23 (13%)	5.7 (2.7) 172 (28%) 447 (72%)
Method to calculate PQ dose					

	No primaquine (N=351)	Low dose daily primaquine $(<0.375$ mg/kg/day) $(N=256)$	Intermediate dose daily primaquine (>=0.375 & $<0.75$ mg/kg/day) (N=182)	High dose daily primaquine (>= $0.75$ mg/kg/day) (N=181)	Total (N=970)
Per actual dose		7 (3%)	160 (88%)	178 (98%)	345 (56%)
Per dosing protocol		249 (97%)	22 (12%)	3 (2%)	274 (44%)
Start day of PQ treatment		` /	,	, ,	, ,
Day 0		253 (99%)	182 (100%)	181 (100%)	616 (100%)
Day 1		2 (1%)	0 (0%)	0 (0%)	2 (0%)
Day 2		1 (0%)	0 (0%)	0 (0%)	1 (0%)
Day 3		0 (0%)	0 (0%)	0 (0%)	0 (0%)
Day 4		0 (0%)	0 (0%)	0 (0%)	0 (0%)
Day 5		0 (0%)	0 (0%)	0 (0%)	0 (0%)
Day 6		0 (0%)	0 (0%)	0 (0%)	0 (0%)
Level of PQ supervision					
Unsupervised		0 (0%)	0 (0%)	0 (0%)	0 (0%)
Partially supervised		249 (97%)	22~(12%)	3 (2%)	274~(44%)
Fully supervised		7 (3%)	160 (88%)	178 (98%)	345 (56%)
Was PQ taken with food?					
No		0 (0%)	0 (0%)	0 (0%)	0 (0%)
Yes		7 (3%)	160 (88%)	178 (98%)	345 (56%)
Recommended		249 (97%)	22 (12%)	3 (2%)	274 (44%)
Other treatment given					
AL	2 (1%)	0 (0%)	0 (0%)	0 (0%)	2 (0%)
AsAq	0 (0%)	0 (0%)	0 (0%)	0 (0%)	0 (0%)
Cq	349 (99%)	256 (100%)	182 (100%)	181 (100%)	968 (100%)
DP	0 (0%)	0 (0%)	0 (0%)	0 (0%)	0 (0%)
Transmission intensity of					
the site	0 (007)	0 (001)	0 (007)	0 (007)	0 (007)
Low	0 (0%)	0 (0%)	0 (0%)	0 (0%)	0 (0%)
Moderate	0 (0%)	0 (0%)	0 (0%)	0 (0%)	0 (0%)
High	351 (100%)	256 (100%)	182 (100%)	181 (100%)	970 (100%)
Not available	0 (0%)	0 (0%)	0 (0%)	0 (0%)	0 (0%)
Geographical region Africa	0 (0%)	0 (0%)	0 (0%)	0 (0%)	0 (0%)
	` '	` '	` '	` '	
Americas	0 (0%)	0 (0%)	0 (0%)	0 (0%)	0 (0%)
Asia-Pacific	351 (100%)	256 (100%)	182 (100%)	181 (100%)	970 (100%)
Relapse Peridocity Low periodicity	351 (100%)	256 (100%)	182 (100%)	181 (100%)	970 (100%)
High periodicity	0 (0%)	0 (0%)	0 (0%)	0 (0%)	0 (0%)
• •	0 (0,0)	0 (0,0)	0 (0,0)	0 (0,0)	0 (0,0)
G6PD categories					
(Qualitative test) <30%	0 (0%)	0 (0%)	0 (0%)	0 (0%)	0 (0%)
>=30%	351 (100%)	256 (100%)	182 (100%)	181 (100%)	970 (100%)
G6PD categories	(-00/0)	(+00/0)	(100/0)	(10070)	2.2 (20070)
(Quantitative test)					
<30%	0 (0%)	0 (0%)	0 (0%)	0 (0%)	0 (0%)
30-<70%	0 (0%)	0 (0%)	0 (0%)	0 (0%)	0 (0%)
>=70%	0 (0%)	0 (0%)	0 (0%)	0 (0%)	0 (0%)
Missing	351 (100%)	256 (100%)	182 (100%)	181 (100%)	970 (100%)

#### 3.3 Summary of the gastrointestinal tolerability outcomes

The primary endpoint for this analysis was a composite indicator including the presence of vomiting or anorexia or diarrhoea on days 5-7 after enrolment.

Secondary endpoints for this analysis were:

a) the presence of vomiting, nausea, anorexia, abdominal pain, diarrhoea or dizziness assessed separately

on days  $5-7^1$ 

b) the presence of the composite endpoint including vomiting or anorexia or diarrhoea on day 0, days 1-2 and days 5-7, assessed separately

<sup>1</sup>Assessment of nausea, dizziness and abdominal pain was restricted to patients older than 5 years Table 2 provides a summary of the outcome experienced within each Primaquine treatment arm.

Table 2\_tol: Summary of gastrointestinal outcomes, categorised by total primaquine category

		Primaquir	ne Treatment			
	No primaquine	Low dose daily primaquine $(<0.375$ mg/kg/day)	Intermediate dose daily primaquine (0.375 & <0.75 mg/kg/day)	High dose daily primaquine ( 0.75 mg/kg/day)	Total	
Outcomes include participants of a	all ages					
	(N=351)	(N=256)	(N=182)	(N=181)	(N=970)	
Composite on day 0						
No	232 (66.1 %)	180 (70.3 %)	119 (65.4 %)	124 (68.5 %)	655 (67.5 %)	
Yes	119 (33.9 %)	76 (29.7 %)	63 (34.6 %)	57 (31.5 %)	315 (32.5 %)	
Composite between days 1-2						
No	294 (84.7 %)	206 (80.5 %)	153 (85.5 %)	155 (88.6 %)	808 (84.4 %)	
Yes	53 (15.3 %)	50 (19.5 %)	26 (14.5 %)	20 (11.4 %)	149 (15.6 %)	
Missing	4 (1.1%)	0 (0%)	3 (1.6%)	6 (3.3%)	13 (1.3%)	
Composite between days 5-7						
No	342 (99.1 %)	248 (97.3 %)	173 (97.7 %)	156 (92.9 %)	919 (97.2 %)	
Yes	3 (0.9 %)	7 (2.7 %)	4 (2.3 %)	12 (7.1 %)	26 (2.8 %)	
Missing	6 (1.7%)	1(0.4%)	5 (2.7%)	13 (7.2%)	25 (2.6%)	
Vomiting between days 5-7						
No	344 (99.7 %)	253 (99.2 %)	175 (99.4 %)	165 (98.2 %)	937 (99.3 %)	
Yes	1 (0.3 %)	2 (0.8 %)	1 (0.6 %)	3 (1.8 %)	7 (0.7 %)	
Missing	6 (1.7%)	1(0.4%)	6 (3.3%)	13 (7.2%)	26 (2.7%)	
Anorexia between days 5-7						
No	76 (98.7 %)	7 (100.0 %)	152 (98.1 %)	156 (94.5 %)	391 (96.8 %)	
Yes	1 (1.3 %)	0 (0.0 %)	3 (1.9 %)	9 (5.5 %)	13 (3.2 %)	
Missing	274 (78.1%)	249 (97.3%)	27 (14.8%)	16 (8.8%)	566 (58.4%)	
Diarrhoea between days 5-7						
No	340 (99.7 %)	247 (98.0 %)	175 (98.9 %)	165 (98.2 %)	927 (98.8 %)	
Yes	1 (0.3 %)	5 (2.0 %)	2 (1.1 %)	3 (1.8 %)	11 (1.2 %)	
Missing	10 (2.8%)	4 (1.6%)	5 (2.7%)	13 (7.2%)	32 (3.3%)	
Outcomes restricted to participan	ts >5 years old	. ,				
resultated to participan	(N=322)	(N=234)	(N=164)	(N=157)	(N=877)	
Nausea between days 5-7*	(1022)	(1201)	()	(	(20)	
No	310 (98.4 %)	207 (89.2 %)	158 (98.8 %)	140 (95.2 %)	815 (95.4 %)	
Yes	5 (1.6 %)	25 (10.8 %)	2 (1.2 %)	7 (4.8 %)	39 (4.6 %)	
Missing	7 (2.2%)	2 (0.9%)	4 (2.4%)	10 (6.4%)	23 (2.6%)	
Abdominal pain between days	. ( -/-//	( )	()	- (- /0)	- (, 0)	
5-7*						
No	73 (100.0 %)	7 (100.0 %)	133 (95.0 %)	137 (93.2 %)	350 (95.4 %)	
Yes	0 (0.0 %)	0 (0.0 %)	7 (5.0 %)	10 (6.8 %)	17 (4.6 %)	
Missing	249 (77.3%)	227 (97.0%)	24 (14.6%)	10 (6.4%)	510 (58.2%)	
Dizziness between days 5-7*	- ()	. (,	( -/-/	- (- /0)	- (/0)	
No	72 (98.6 %)	5 (71.4 %)	139 (99.3 %)	145 (98.6 %)	361 (98.4 %)	
Yes	1 (1.4 %)	2 (28.6 %)	1 (0.7 %)	2 (1.4 %)	6 (1.6 %)	
Missing	249 (77.3%)	227 (97.0%)	24 (14.6%)	10 (6.4%)	510 (58.2%)	

Figure 1\_tol: Distribution of primaquine daily dose by primaquine mg/kg daily dose category. Primaquine daily dose categories: Low: <0.375 mg/kg/day, Int (intermediate):  $\ge 0.375$  mg/kg/day and <0.750 mg/kg/day, and High:  $\ge 0.750$  mg/kg/day

#### 3.4: Risk of gastrointestinal intolerance

The risk of gastrointestinal intolerance on days 5-7 was calculated from the number of patients reporting the composite outcome as a proportion of the total number of patients asked about each of the individual components of the composite; i.e. those asked about vomiting or anorexia or diarrhoea on any day between days 5-7. The 95% confidence intervals (CIs) for the risks were calculated as exact binomial CIs. The risks were stratified by primaquine daily dose categories

	N		Percentage (95% CI)		N		Percentage (95% CI)
No PQ	345	•	0.87 (0.18, 2.52)	No PQ	345	•	0.29 (0.01, 1.60)
Low dose PQ	255	•	2.75 (1.11, 5.57)	Low dose PQ	255	•	0.78 (0.10, 2.80)
Intermediate dose PQ	177	•-	2.26 (0.62, 5.68)	Intermediate dose PQ	176	•-	0.57 (0.01, 3.12)
High dose PQ	168		7.14 (3.75, 12.14)	High dose PQ	168	<b>←</b>	1.79 (0.37, 5.13)
norexia		0 5 10 15 20 25 30		Diarrhoea		0 5 10 15 20 25 30	
погеліа	N		Percentage (95% CI)	Diamicea	N		Percentage (95% CI)
No PQ	77	•	1.30 (0.03, 7.02)	No PQ	341	•	0.29 (0.01, 1.62)
Low dose PQ	7	•	0.00 (0.00, 40.96)	Low dose PQ	252	<b>-</b>	1.98 (0.65, 4.57)
Intermediate dose PQ	155	•	1.94 (0.40, 5.55)	Intermediate dose PQ	177	•-	1.13 (0.14, 4.02)
High dose PQ	165	-	5.45 (2.52, 10.10)	High dose PQ	168	•	1.79 (0.37, 5.13)
		0 5 10 15 20 25 30				0 5 10 15 20 25 30	
lausea				Abdominal Pain			
	N		Percentage (95% CI)		N		Percentage (95% CI)
No PQ	315	•-	1.59 (0.52, 3.67)	No PQ	73	•—	0.00 (0.00, 4.93)
Low dose PQ	232	-	10.78 (7.10, 15.50)	Low dose PQ	7	•	0.00 (0.00, 40.96
Intermediate dose PQ	160	•-	1.25 (0.15, 4.44)	Intermediate dose PQ	140	<b>-</b>	5.00 (2.03, 10.03
High dose PQ	147	-	4.76 (1.94, 9.57)	High dose PQ	147		6.80 (3.31, 12.15
		0 5 10 15 20 25 30				0 5 10 15 20 25 30 Percentage	
Dizziness							
	N		Percentage (95% CI)				
No PQ	73	•	1.37 (0.03, 7.40)				
Low dose PQ	7	<b>→</b>	28.57 (3.67, 70.96)				
Intermediate dose PQ	140	•	0.71 (0.02, 3.92)				
			1.36 (0.17, 4.83)				

Figure 2\_tol: Risk of gastrointestinal intolerance by symptoms. For each outcome the risk was estimated as the number of individuals experiencing the symptom as a proportion of the number of individuals asked about the symptom on any day between days 5-7. The confidence intervals (CIs) are exact binomial CIs.

# 3.4.1: Adjusted association between primaquine daily dose categories and gastrointestinal intolerance days 5-7

The plots below show the estimated proportion of patients with gastrointestinal symptoms on days 5–7 by primaquine treatment regimen, adjusted for age, sex and baseline parasite density. A logistic model was fit and the adjusted proportions were estimated using mean values for age, sex, and log10 baseline parasite density.

Care should be taken when interpreting these results, as model assumptions have not been fully assessed in this automated report format.

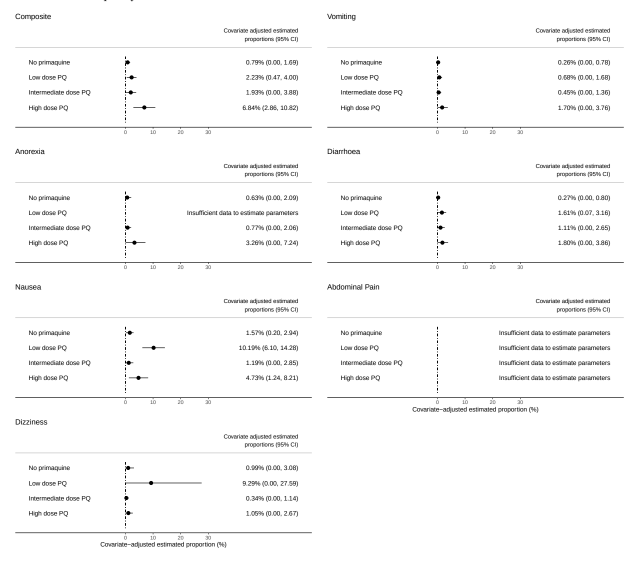


Figure 2\_tol: Covariate-adjusted estimated proportion of patients with gastrointestinal symptoms on days 5-7 by primaquine treatment regimen. A logistic mixed effects model was fit, with study site as the random effect, and the adjusted proportions were estimated using mean values for age, sex, and log10 baseline parasite density. Note: The lower confidence intervals have been limited to 0% and the upper limited to 100%.

#### 3.4.2: Risk of Acute Vomiting on days 0-2 and 3-13

The unadjusted risk of vomiting within an hour of primaquine administration (acute vomiting) was calculated on days 0-2 and days 3-13 for each primaquine dose group.

Table 3\_tol: Risk of acute vomiting on days 0-2 and 3-13 by primaquine daily dose categories

	Risk of acute vomiting		
Primaquine treatment	Days 0-2	Days 3-13	
Low dose daily primaquine ( $<0.375 \text{ mg/kg/day}$ ) Intermediate dose daily primaquine ( $>=0.375 \& <0.75 \text{ mg/kg/day}$ ) High dose daily primaquine ( $>=0.75 \text{ mg/kg/day}$ )	7/254 (2.8%) 1/180 (0.6%) 1/181 (0.6%)	0/7 (0.0%) 0/156 (0.0%) 0/167 (0.0%)	

Days 0–2			Days 3–13					
	N	Percentage (95% CI)		N	Percentage (95% CI)			
Low dose PQ	254	2.76 (1.12, 5.60)	Low dose PQ	7	0.00 (0.00, 40.96)			
Intermediate dose PQ	180 —	0.56 (0.01, 3.06)	Intermediate dose PQ	156 •	0.00 (0.00, 2.34)			
High dose PQ	181 —	0.55 (0.01, 3.04)	High dose PQ	167 •	0.00 (0.00, 2.18)			
	0 1 2 3 4 Percentage			0 1 2 3 4 Percentage				

Figure 3\_tol: Risk of acute vomiting on days 0-2 and 3-13 by primaquine daily dose categories. The confidence intervals (CIs) are exact binomial CIs.